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Applications of Single-cell DNA Sequencing

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Abstract

Over the past decade, genomic analyses of single cells—the fundamental units of life—have become possible. Single-cell DNA sequencing has shed light on biological questions that were previously inaccessible across diverse fields of research, including somatic mutagenesis, organismal development, genome function, and microbiology. Single-cell DNA sequencing also promises significant future biomedical and clinical impact spanning oncology, fertility, and beyond. While single-cell approaches that profile RNA and protein have greatly expanded our understanding of cellular diversity, many fundamental questions in biology and important biomedical applications require analysis of the DNA of single cells. Here, we review the applications and biological questions for which single-cell DNA sequencing is uniquely suited or required. We include a discussion of the fields that will be impacted by single-cell DNA sequencing as the technology continues to advance.

Keywords

Single-cell DNA sequencing; single-cell genomics; lineage tracing; germ cells; organismal development; DNA modifications

Introduction

Single-cell DNA sequencing (scDNA-seq) encompasses a suite of technologies and approaches that interrogate DNA at the level of single cells. These technologies contrast with standard DNA sequencing, also known as “bulk” sequencing, that homogenizes the DNA content of usually thousands to millions of cells. Much like the voice of a single individual or a small number of individuals can be “drowned out” in a large crowd, genomic signals (variants, DNA modifications, or structural properties of DNA) that are present in only one or a small number of cells in a sample may be undetectable without interrogating

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single-cell genomes. The development of scDNA-seq methods has at every step been motivated by biological questions that seek to explore this cellular genomic diversity that would otherwise be missed by bulk sequencing.

In this review, we focus on the major applications of scDNA-seq rather than its technological aspects. We begin by introducing the basic concepts of scDNA-seq to provide a framework for understanding why certain applications depend on its unique capabilities and why for some important biological and biomedical questions it is the only suitable technology. Following this, we discuss the major biological fields that scDNA-seq has impacted and the discoveries it has enabled. These include a wide array of fields: somatic mutation and mosaicism, organismal development, germ cell mutation and development, fertility, cancer, epigenetic regulation of the genome, genome organization, and microbiology. This review specifically focuses on single-cell genomics methods, i.e. those that profile single-cell DNA at larger genomic scales or genome-wide. Single-cell genomic approaches have only become feasible over the past decade, whereas profiling of one or small numbers of loci in single cells was previously available. We will also use the term single-cell genomics interchangeably with single-cell DNA sequencing. While single-cell genomics is sometimes used in the literature to refer to single-cell RNA sequencing, to avoid terminological confusion, here, single-cell genomics and single-cell transcriptomics will refer to single-cell DNA and single-cell RNA sequencing, respectively.

A conceptual framework for single-cell DNA sequencing

In this section, we summarize key concepts necessary for understanding the subsequent sections that focus on the applications of scDNA-seq. These concepts are not only important to understand as a terminological and technical reference, but they also help explain the unifying characteristics of the applications for which scDNA-seq is suitable. scDNA-seq unlocks a particular set of capabilities, described below, that are advantageous or unique relative to other technologies; therefore, every application of scDNA-seq is distinguished by its reliance on these capabilities. While this section explains this unifying framework, we will not review in detail specific scDNA-seq technologies, as these have been well reviewed previously (52, 66).

scDNA-seq is characterized by three core capabilities, which we term (a) Fidelity, (b) Co-presence, and (c) Phenotypic association, that together help understand the types of biological questions and applications for which it is best suited (Figure 1A). As we discuss below, different scDNA-seq technologies may possess different subsets of these three capabilities, and moreover, none of these capabilities are unique to scDNA-seq. However, scDNA-seq is distinguished from other approaches by its potential to achieve all three capabilities on a genome-wide scale.

The ‘Fidelity’ capability of scDNA-seq (Figure 1A) is its ability to overcome the limits of bulk DNA sequencing to detect features of DNA (mutations, modifications, or other properties of DNA) that are at a low level of mosaicism; i.e. features present in only a small subset of cells in a sample. Bulk DNA sequencing is limited by sequencing error that is a constant fraction of the total sequencing coverage. While increasing the depth

of bulk sequencing coverage initially increases sensitivity for lower-level mosaic DNA features by virtue of deeper sampling, the fact that sequencing error is a constant fraction of total coverage means that further increases in coverage eventually reach a hard limit: mosaic features with less than ~0.5% mosaicism cannot be detected because they cannot be distinguished from sequencing error (18, 146). In contrast, scDNA-seq is not limited by sequencing error, since the sequencing error rate is much lower than the expected 50% signal level of heterozygous DNA features or even 25% single-stranded DNA features. Nevertheless, errors of single-cell DNA amplification can in some cases rival the signal level of true genetic features, motivating the on-going development of new methods for high-fidelity single-cell genome amplification (52).

The ‘Co-presence’ capability of scDNA-seq (Figure 1B) refers to its ability to ascertain which mosaic DNA features were present together in the same cells. For example, if a tissue sample contains two different low-level mosaic genetic variants, only a single-cell method can determine whether both variants are present in the same cells or in mutually exclusive subsets of cells. Bulk methods homogenize samples prior to sequencing, so the information about which mosaic DNA features were present in the same cell or subsets of cells is lost. In contrast, scDNA-seq preserves this information. While there exist bulk methods that either directly profile large DNA fragments or preserve long-range phasing information that can technically ascertain co-presence for small numbers of DNA features, these methods have limited phasing distance and they cannot ascertain co-presence of DNA features on different chromosomes or genome-wide co-presence of a large number of DNA features. Therefore, co-presence is a key capability of scDNA-seq that enables many of its unique applications.

The ‘Phenotypic association’ capability of scDNA-seq (Figure 1C) is its potential to be combined with simultaneous single-cell phenotyping to identify the cell type(s) or cell state(s) in which specific DNA features are present. This capability derives from the fact that scDNA-seq analyzes single cells, thereby preserving the link between a cell’s DNA features and its phenotype. However, ‘Phenotypic association’ requires technology that combines scDNA-seq with phenotypic profiling, for example single-cell DNA plus RNA, or single-cell DNA plus proteomic profiling. Alternatively, some single-cell DNA features such as methylation themselves provide information about cell phenotype. The ‘Phenotypic association’ capability of scDNA-seq is technically challenging since combining genome-wide scDNA-seq with phenotypic profiling can compromise the resolution of the scDNA-seq component (106). However, new multi-omic scDNA-seq technologies are emerging (31, 105) and are a focus of current technology development given their importance for many of the applications we review below.

The ‘Co-presence’ and ‘Phenotypic association’ capabilities are together particularly important for samples composed of genetically distinct heterogeneous cells, a common situation often central to the applications of scDNA-seq. This includes both applications profiling cells within an organism, such as distinguishing tumor versus normal cells in a tumor sample, and profiling mixtures of cells from distinct species, e.g., soil samples or the gut microbiome.

None of the above three capabilities are unique to scDNA-seq. For example, fluorescence in situ hybridization can ascertain co-presence of a small number of genetic variants as well as their phenotypic association with specific cell types. Another example is single-cell cloning (176), where a single cell is expanded *in vitro* to create a bulk sample deriving entirely from one cell, thereby providing both fidelity and co-presence information for mosaic DNA variants that were present in the original single-cell. The above are only a few examples of existing non-single-cell DNA sequencing methods that can achieve one or more of the three key capabilities of scDNA-seq. However, scDNA-seq is distinguished by the fact that: (a) it can potentially achieve all three capabilities simultaneously; (b) it can determine co-presence at a genome-wide scale; and, (c) it can be applied directly to primary cells and is not limited to specific cell types. The last feature distinguishes it from single-cell cloning approaches that entail cloning and culture bottlenecks that may bias results and which are not feasible for some cell types (for example, mature adult neurons and unculturable bacteria). Every application of scDNA-seq can be traced back to one or more of these three core capabilities, and the most advanced applications leverage all three. It is therefore a useful exercise for any proposed scDNA-seq application or research project to critically evaluate which of these three capabilities is needed—Fidelity, Co-presence, and Phenotypic Association—and consequently, whether scDNA-seq is required as opposed to employing an alternative technology. Conversely, consideration of these three capabilities can help identify new biological questions that can be uniquely addressed by scDNA-seq.

It is worth noting also that the fidelity capability of scDNA-seq is only relative to the fidelity of current DNA sequencers. If the fidelity of DNA sequencers were to advance by approximately four orders of magnitude, that would leave only Co-presence and Phenotypic association as distinguishing characteristics of scDNA-seq.

A major limitation of scDNA-seq derives from another limitation of current DNA sequencers: only a small fraction of the DNA that is input into a sequencer is captured and sequenced. Therefore, scDNA-seq requires amplification of DNA prior to sequencing. And because there is no perfect amplification method, these can introduce errors that confound analyses (52). Currently, there are several different single-cell DNA amplification methods, each with a different error profile and performance characteristics, which have been reviewed previously (52, 66). The best single-cell DNA amplification method depends on the application, and ideally, the error mode of the amplification method will be orthogonal to the DNA feature(s) of interest.

Finally, throughput—the number of single cells that can be profiled in one experiment—is an important parameter of different scDNA-seq methods. Some applications can suffice with tens or hundreds of single cells, while other applications and most future applications will require thousands of cells. New methods utilizing droplet encapsulation (117), combinatorial indexing (173), and nanowell devices (81) are increasing the throughput of scDNA-seq to thousands of cells and expanding the range and depth of questions that can be addressed. However, current scDNA-seq methods and sequencing costs impose a trade-off: methods with higher throughput have lower genome-wide coverage or profile fewer DNA loci per cell. The applications discussed in this review differ widely in their throughput and

coverage requirements (Figure 1D), motivating careful matching between an application's requirements and the capabilities of the chosen scDNA-seq method.

Somatic cells: mutational processes and mosaicism

Somatic mutations—mutations that occur post-zygotically in somatic cells—are ubiquitous in multi-cellular organisms due to exogenous and endogenous mutagens, DNA replication errors, chromosomal damage, imperfect DNA repair, and other mutagenic processes (155, 181). Indeed, it can be safely assumed that no two cells in an organism have an identical genome sequence, in particular, organisms whose genomes are significantly larger than the inverse of their per-cell division mutation rate. While most somatic mutations are benign, they are the cause of cancer and many non-cancer genetic diseases, and they are speculated to contribute to physiologic aging (38, 110, 131, 155, 159, 181). Quantifying the rates and spectra of somatic mutations is therefore of fundamental importance. However, within an individual organism, most somatic mutations are either present at very low mosaicism or unique to one cell (159), placing them below the detection limits of current DNA sequencers. The fidelity capability of scDNA-seq allows detection of somatic mutations regardless of mosaicism level and has provided some of the first direct, genome-wide measurements of somatic mutations rates. Here, we will review the application of scDNA-seq in quantifying somatic mutational processes and mosaicism. scDNA-seq of germ cell mutations will be reviewed in a later section. We also recommend an accompanying review in this issue for more in-depth discussion of somatic mosaicism and single-cell sequencing ([Reference review by Christopher Walsh in this same issue]).

Somatic mutation rates vary across tissues, developmental timepoints, cell types, and mutation types, and scDNA-seq studies have begun to systematically quantify somatic mutations across these dimensions. The first scDNA-seq studies quantifying genome-wide somatic mutation in normal (non-cancer) cells profiled LINE-1 (L1) retrotransposon insertions and copy number variants (CNV) in human brain (40, 112). Both studies were motivated by long-standing hypotheses that somatic variation may be prevalent and adaptive in the brain (113). These studies successfully identified somatic L1 insertions and CNVs in human cortical neurons, but at lower rates than prior indirect measurements had anticipated: < 0.6 somatic L1 insertions per neuron, < 2% aneuploid neurons, and 41% of neurons each harboring few (mostly 1-2) megabase-scale CNVs (40, 112). Several subsequent scDNA-seq studies were concordant with these findings (21, 39, 42, 77), suggesting that these types of somatic mutations are largely tolerated physiologically while creating a low-level of neuronal genomic diversity and in rare cases causing neurologic disease (27, 131).

In contrast to these studies of CNVs in mature cells, remarkably high rates of aneuploidy and CNVs have been found in early pre-implantation human embryos by single-cell microarray profiling and in macaque embryos by scDNA-seq (30, 157). scDNA-seq of pre-implantation macaque embryos found that 74% had at least one blastomere with a whole or partial chromosome copy number abnormality, and remarkably, detected some of the missing chromosomal material in cellular fragments arising from aberrant multipolar divisions at the 1- or 2-cell stage (30).

The possibility of pathogenic mosaic CNVs in human pluripotent stem cells (hPSC) used for cellular therapies has motivated scDNA-seq studies of these cells. One scDNA-seq study found that 23% of hPSCs had at least one large megabase-scale CNV, one third of which had more complex karyotypes. Broader application of scDNA-seq for routine safety assessment and screening of cellular and gene-modifying/editing therapies can be anticipated.

In addition to measuring somatic L1, CNV, and aneuploidy rates, scDNA-seq studies have quantified single-nucleotide variant (SNV) somatic mutation rates. scDNA-seq SNV studies face a greater analytic challenge separating single-cell amplification artifacts from bona fide somatic variants, and have recently greatly benefited from computational methods utilizing local spatial variation and read-based phasing of candidate variants to germline variants in order to filter artifacts (16, 104). The first genome-wide scDNA-seq SNV study of normal (non-cancer) cells examined human cortical neurons and found much higher rates of SNV mutation relative to other mutation types and a mutational strand bias suggesting some originate from transcription-coupled damage (96). A subsequent study refined these estimates across a larger set of neurons and age groups to quantify 300-900 somatic SNVs per neuron at birth and ~20 new somatic SNVs per year of life, leading to more than triple the number of SNVs (~2-3,000) by the time an individual is >80 years-old (95). Interestingly, these aging mutation rates were higher in hippocampal neurons and in individuals with DNA-repair syndromes (95).

Additional scDNA-seq studies have measured somatic SNV rates of ~900 per fibroblast from a human toddler, ~460 per B-lymphocyte in newborns increasing to ~3,000 in centenarians, and ~1,000 per hepatocyte in newborns increasing to ~4-5,000 in elderly individuals (17, 34, 180). Notably, all of the above studies have found that most SNVs are unique to one cell in the sample and can be computationally assigned to an “aging clock-like” mutational process (whose mechanism is enigmatic) previously identified in cancer sequencing studies (COSMIC signatures 1/5) (3). Altogether, these studies reveal a relatively similar burden of somatic SNVs per cell at birth across cell types and a significant increase with age, albeit at different rates in different cells types. The possible functional impact of accumulating somatic SNVs with age throughout the body is an important open question.

Over the long-term, a full view of somatic mutation rates will necessitate scDNA-seq studies spanning numerous developmental timepoints, cell types, tissues, and mutation types. Most scDNA-seq studies have to date also only focused on one or a few cell types among hundreds of cell types in the body. Bulk methods for measuring somatic mutation rates have begun to fill this gap, such as clonal expansion of primary single cells followed by bulk sequencing (5), deep sequencing of tissues where clones remain spatially restricted (20), and RNA-seq (174), with largely consistent findings to scDNA-seq. However, clonal expansion of primary cells is susceptible to culture bottlenecks and cannot be applied to many cell types, deep sequencing of spatial clones is not applicable to many tissues and cannot resolve different cell types, and bulk RNA-seq cannot easily measure cell type-specific mutation rates. Therefore, going forward, the unique advantage of scDNA-seq relative to other methods for somatic mutation quantification will be via multi-omic scDNA-seq methods

(i.e. utilizing scDNA-seq's phenotypic association capability) to quantify mutation burdens of specific cell types from any tissue.

Organismal development and lineage tracing

One of the long-standing goals of biology has been to understand how a single cell (the zygote) transforms into the billions or trillions of cells of a complex multi-cellular organism (1, 115). The development of an organism can be represented by a lineage tree—a diagram of the cell divisions from the zygote to each of its cells, ideally annotated with spatial data, cell phenotypes, and the molecular determinants of lineage decisions (74). Lineage trees are the blueprints of development, depicting the relationships between progenitor cells and cell types throughout development, and providing a framework for understanding stem-cell and cell-type hierarchies, aberrant lineages in disease, and tissue homeostasis. Lineage tracing—the collection of methods used to determine lineage trees—are therefore among the most fundamental tools of biology (79). In this section, we review the application of scDNA-seq for lineage tracing with a focus on organismal development. scDNA-seq lineage tracing of cancer is reviewed in the 'Cancer' section.

Lineage tracing methods are either prospective or retrospective, and scDNA-seq can be used for either approach (7, 169). In prospective lineage tracing, cells are experimentally tagged by a heritable marker (e.g., a genetically-encoded barcode), and their progeny are profiled at a later timepoint. Because prospective lineage tracing requires invasive manipulation of the organism, it is only applicable to animal models. In retrospective lineage tracing, lineage trees are reconstructed from somatic DNA mutations that occurred spontaneously during an organism's lifetime, so it is non-invasive and feasible for studying human development. Regardless of whether DNA mutations are experimentally induced (prospective approach) or occur spontaneously (retrospective approach), scDNA-seq can detect these mutations to construct a matrix showing which mutations were present in which single cells (note, this is another way of describing scDNA-seq's 'co-presence' capability). This mutation-by-cell matrix is then transformed into a lineage tree through phylogenetic reconstruction algorithms (115).

Although scDNA-seq can be used for both prospective and retrospective lineage tracing, its greatest long-term potential lies in retrospective lineage tracing. Recently, prospective methods of lineage tracing have been developed in which DNA mutations are created using CRISPR-Cas9 specifically in sites that are transcribed into RNA, such as 3' untranslated regions (2, 26, 47, 133, 147, 167). This allows DNA mutations to be detected by scRNA-seq, in conjunction with phenotypic profiles, without requiring scDNA-seq. These methods, which have been reviewed elsewhere (7, 74, 115, 160), have enabled high-throughput phenotypic lineage tracing for the first time and have removed the requirement for scDNA-seq in prospective lineage tracing of animal models. In contrast, scDNA-seq is essential for retrospective lineage tracing—the only method that can be applied to humans. This is because most spontaneous somatic mutations, which retrospective lineage tracing relies on, are in the large majority of the genome that is not transcribed and cannot be captured by scRNA-seq. We therefore focus this section on the use of scDNA-seq for retrospective lineage tracing. scRNA-seq-based prospective lineage tracing methods are also currently

limited by their inability to induce mutations continuously across broad development timepoints (115). scDNA-seq retrospective lineage tracing, however, leverages somatic mutations that occur in every cell division that can theoretically reconstruct the lineage of any tissue and any developmental timepoint at the maximum possible resolution (1, 48). Once scDNA-seq retrospective lineage tracing scales to the throughput of the above scRNA-seq prospective lineage tracing methods and is combined with phenotypic (e.g., RNA or protein) profiling—though this may take many years to realize—one can expect it will become the primary lineage tracing approach for both humans and animal models.

The resolution of retrospective lineage tracing scales with both the number of loci that are genotyped and their somatic mutation rate (14). Microsatellites are abundant genomic elements (>1 million loci in the human genome) consisting of repeated 1-6 base pair motifs that have the highest mutation rates of any type of genomic element (48). This makes them an attractive target for retrospective lineage tracing (48, 141). In fact, it has been estimated that every cell division may be tagged by at least one microsatellite mutation, theoretically allowing reconstruction of an organism's complete lineage tree (48). Early pioneering scDNA-seq studies of microsatellite mutations utilized only a small set of < 100 loci but were able to confirm some basic lineage information such as clustering liver cells separately from other organs' cell types (140) and closer lineage relationships of muscle satellite cells that are physically proximal (163). Another study of ~120 microsatellite loci found evidence for periodic replacement of all the cells in a colon crypt with progeny of a single stem cell, a process termed monoclonal conversion (136).

The number of microsatellite loci profiled per cell has subsequently increased to achieve greater lineage resolution, with ~1-2,000 loci per cell in two different studies (14, 166) and ~12,000 loci in a third study (153). One of these studies was able to simultaneously profile methylation in the same cells and recovered a known cultured lineage tree with 88% accuracy (166). However, there remain significant challenges associated with artifactual microsatellite mutations that occur during single-cell genome amplification and library preparation (14). These will need to be overcome before the full potential of microsatellites for scDNA-seq lineage tracing can be realized.

Several scDNA-seq studies have used other types of somatic mutations for retrospective lineage tracing, such as SNVs, which are also estimated to occur in every cell division (1), and L1 retrotransposon insertions. Nearly all these studies have so far focused on the human brain. One study traced the spatial distribution of two somatic L1 retrotransposon insertions identified by scDNA-seq in the brain of an individual, finding both a focally restricted lineage in the frontal lobe and a widely distributed lineage consistent with spatial mixing of progenitor cells during early development (41). The L1 insertion of the focal lineage also harbored a poly-A microsatellite that itself mutated somatically at a high rate to mark different subclades in each brain region, consistent with tangential migration and intermingling of radial clones of progenitors in the cortex (41). A notable study profiled 18 somatic mutations, including 15 SNVs, to reconstruct a lineage tree of 136 single neurons with 4 clades and additional sub-clades (96). Remarkably, the 4 root clades in this individual were dispersed across the cortex at low mosaicism, again indicating significant spatial mixing of early brain progenitors, whereas later sublineages had more focal distributions

(96). A more recent study performed simultaneous scDNA-seq and RNA profiling of > 1,000 cortical neurons (65). It found that SNVs of later sub-lineages were progressively restricted to either excitatory neurons or to inhibitory neurons, consistent with these cell types' different developmental origins (65). Upper layer neurons also comprised a greater fraction of later lineage branches, consistent with the inside-out formation of cortical layers (65). These studies have confirmed processes of brain development previously seen only in animal models using prospective (invasive) approaches. The theme of these studies is that as their throughput has been increasing, more detailed features of brain development have emerged. Further such studies may eventually reveal human-specific developmental processes.

Because somatic mutations accumulate in every cell division, scDNA-seq retrospective lineage tracing can also be used to estimate for any cell or tissue its approximate number of cell divisions since the zygote, also known as the lineage depth. This can help understand organogenesis and tissue turnover. In mismatch-repair deficient mice with higher microsatellite mutation rates, one scDNA-seq study estimated a depth of 29 for oocytes, 24-40 for various types of adult stem cells consistent with their relative quiescence, and a linear increase in depth of about 1 cell division per day for B-cells (164). This method could be used to investigate stem cell activity in other organs such as the brain and in disease states.

An important recent advance has been scDNA-seq retrospective lineage tracing that uses mitochondrial DNA (mtDNA) instead of nuclear DNA mutations (83, 99, 171). The small size of the mitochondrial genome and the presence of hundreds or more copies per cell enable this approach to scale to thousands of cells via droplet-based methods (83). mtDNA is also readily captured in standard scDNA-seq chromatin accessibility assays, such that mtDNA mutations and cell phenotype information are captured simultaneously from the same cells (83, 99, 171). The throughput of this approach is illustrated by one experiment that traced the lineage relationship of >16,000 CD34⁺ hematopoietic stem cells (HSCs) sampled at one point and peripheral blood mononuclear cells sampled three months later from the same individual (83). They identified >250 clonal groups, indicating that a large pool of HSCs is active in healthy blood production, but they also observed variability in the contribution of individual HSC clones to blood cell production (83). This method has also been used to lineage trace blood and colorectal cancers (83, 99, 171).

Overall, the main advantages of single-cell mtDNA lineage tracing are its significant throughput in terms of cell number and the fact that it does not require *a priori* knowledge of sub-clonal somatic mutations. This contrasts with nuclear scDNA-seq lineage tracing where sequencing the entire genome across thousands of cells is not feasible. However, the advantages of mtDNA lineage tracing are balanced by the limited number of mtDNA mutations that occur in the small mitochondrial genome, even accounting for mtDNA's higher mutation rate (1). In the long-run, this will limit the resolution of mtDNA lineage trees relative to nuclear genome lineage trees. Additionally, mtDNA is not a completely faithful lineage marker, because mutations in only a small subset of a cell's mitochondria (heteroplasmy) may not always segregate to both daughter cells. Nevertheless, the biological insight provided by scDNA-seq mtDNA is promising due to its relative ease of integration

into established single-cell workflows, and it provides a glimpse into the future of what high-resolution scDNA-seq lineage tracing may achieve. By virtue of its ability to ascertain somatic DNA mutations genome-wide, scDNA-seq—combined with cell phenotyping—has the potential to provide the first systematic and truly general approach for high-resolution lineage tracing.

Germ cells: meiotic recombination and germline mutation

Meiosis is a specialized cell division in which haploid gametes are generated from diploid cells through one round of genome replication followed by two rounds of cell division (108). During this process, mutation generates new alleles and recombination generates new combinations of alleles via exchange of genetic material between chromosomes known as crossovers (108). Together, they create heritable genetic variation and the substrate for evolution via selection. Recombination is mechanistically essential to success in meiosis: it is initiated by programmed DNA double-strand breaks, of which only a fraction are repaired with a crossover (69). Insufficient crossovers or their improper localization leads to aneuploidy, which affects 20-30% of human eggs and 1-8% of human sperm, leading to miscarriage in over 25% of pregnancies and severe developmental defects in 0.3% of live births (22, 56, 57). Errors in recombination also lead to numerous genetic disorders (23). In this section, we discuss the application of scDNA-seq for understanding meiotic recombination and germ cell mutation.

The most high-resolution approaches to-date for studying recombination have been mapping of crossovers in pedigrees (10, 55, 78, 89) or admixed populations (58, 165) and inference of historical recombination via breakdown of linkage disequilibrium (so-called LD-based maps) (37, 46). However, these approaches have significant limitations—(i) Individual-versus population-level maps: as a germline process with strong impact on fitness, it is important to measure recombination at the level of gametes or in individuals who are unable to reproduce, because pedigree-based and LD-based methods only probe viable offspring. Separately, the fine-scale landscape of recombination in humans and many vertebrates is determined predominantly by the protein PRDM9, whose DNA-binding properties may vary across individuals in the population (120, 126). Interpretation of recombination maps aggregated across individuals is therefore complicated by *PRDM9* allelic diversity in the population. (ii) Sample availability: identification of crossovers at high-resolution in pedigrees requires dense genotyping or whole-genome sequencing in large numbers of related individuals (55, 78, 92, 142). Such resources are challenging and expensive to accumulate in humans and model organisms and this approach is not typically feasible for species in the wild. (iii) Gender differences: recombination-related phenotypes show strong sexual dimorphism in human and many other species (19, 78, 102). However, understanding of female recombination has lagged behind male recombination, in part due to the challenges of working with limited amounts of tissue that are difficult to obtain. As discussed below, scDNA-seq of germ cells and gametes has overcome many of these limitations.

scDNA-seq studies of sperm demonstrate that human sperm have high rates of genome instability and aneuploidy (12, 98, 161), in striking contrast with mouse (59). Analysis

of tens of thousands of single sperm from multiple human males revealed significant inter-individual differences in aneuploidy rates ranging from 1% to 5%, with an apparent excess of whole-chromosome losses over gains (12). These data revealed that whole-chromosome gains may result from improper segregation at either of the two meiotic divisions with a two-fold higher error rate in the second meiotic division for the autosomes. The number and localization of crossovers covary significantly between cells (12), and a higher number of crossovers within a cell appears to be protective against chromosome mis-segregation (12, 98). Further insight into male meiotic segregation was provided by scDNA-seq of spermatocytes (precursors of sperm) which showed an unexpected class of meiotic segregation error in an azoospermic mouse (173). Single-sperm sequencing in mouse has also revealed novel insights into the dynamic processes underlying the pairing of homologous chromosomes and the determination of which DNA breaks become crossovers: a high-resolution map of crossovers in mouse sperm showed that DNA breaks that repair more quickly are also more likely to become crossovers (59). scDNA-seq may also enable individualized quantification of sperm mutation rates and the risk of transmitting de novo mutations to offspring (161), however further technological improvements in throughput, coverage, and accuracy will be required.

Acquisition of oocytes involves a surgical procedure and only small numbers of oocytes (< 10) are typically harvested per individual in human and other mammals (61, 127). Limited tissue availability, together with the fact that all the products of a single meiosis can be harvested together (oocyte and polar bodies), makes scDNA-seq of oocytes a powerful avenue for studying female meiosis. Single-cell data has confirmed that crossovers exhibit interference (they are further away from each other than expected by chance) in human oocytes (61, 127) as they are in sperm (98, 161). Human oocytes have very high aneuploidy rates (18% - 70%) (61, 127), and in contrast with males, have significantly higher error rates in the first meiotic division. Genome comparisons of the oocyte with the first and second polar bodies led to identification of a new mode of “reverse” segregation of chromatids, which was observed to be the dominant signature of aneuploidy (127). The authors of that study also showed that crossing over is protective against aneuploidy, with further meiotic drive against non-recombinant chromatids (127). Understanding the mechanisms of aneuploidy further will require phasing of missegregated chromosomes (127), which remains a challenge with the relatively low genome-coverage obtained per cell. A major focus of recent single-cell research has been to maximize throughput and aggregate small amounts of information per cell across large numbers of cells (12, 173). Generating insights into female recombination, however, will require a parallel emphasis on maximizing the amount of information obtained per cell, e.g., by increasing the genome-coverage per cell (59) and long-range haplotyping of individual molecules (150).

Fertility

A major emerging clinical application of scDNA-seq is in prenatal screening and genetic testing in natural conceptions and in vitro fertilization (IVF). During assisted reproduction, multiple embryos are typically created by IVF and screened for aneuploidy and highly deleterious mutations, followed by implantation of selected embryos. Single-cell genomic approaches have improved upon conventional methods of analyzing embryo biopsies by

enabling simultaneous detection of aneuploidy and mutations genome-wide (80, 156, 168). Sequencing of polar bodies can also be used to detect maternally inherited aberrations in oocytes (61). These approaches are adding insight into the nature of aneuploidy and mosaicism in embryos, which may be mitotic or meiotic in origin (114, 145). In natural conceptions, invasive testing carries the risk of procedure-induced miscarriage and non-invasive sequencing of cell-free fetal DNA in maternal plasma has been rapidly adopted in clinical practice (71, 158). However, the reliability of this method is variable and recent developments in extraction of genetic information from fetal cells circulating in maternal blood may lead to future single-cell genomic tools for this application (64, 143). However, remaining challenges are reliable isolation of rare fetal cells from maternal blood and concern that the predominant and more readily isolated fetal trophoblast cells are not always representative of the genome of the fetus (25).

Cancer

Cancer is a disease of the genome in which an aberrant lineage of cells accumulates somatic mutations along an evolutionary trajectory of uncontrolled growth. scDNA-seq has revolutionized our ability to study the dynamic evolutionary processes by which tumor lineages evolve and interact with selective pressures during oncogenesis, tumor growth, and treatment (8, 124, 137). This section reviews the most important applications of scDNA-seq in cancer research: intra-tumoral heterogeneity, clonal evolution, invasion and metastasis, circulating tumor cells, and therapeutic response. We will also highlight emerging approaches and clinical applications.

Tumors evolve genetically heterogeneous lineages over time, and the resulting intra-tumoral heterogeneity (ITH) plays a significant role in tumor growth, metastasis, and treatment response (29). A tumor's ITH, defined by the subclonal structure of its lineages and driver mutations, is its *dramatis personae* and of prime importance for understanding its biology. The first study to examine ITH with genome-wide scDNA-seq utilized copy number aberrations (CNAs) to distinguish subclonal lineages within breast tumors (123). Since then, numerous scDNA-seq studies of CNAs and point mutations have catalogued ITH across diverse hematologic (6, 51, 63, 67), breast (9, 50, 162), ovarian (116), colorectal (13, 86, 94), renal (172), bladder (90), lung (43), liver (62), and brain (45) cancers. These studies have found that most tumors contain more than one major subclonal lineage, in addition to many tumors that harbor a larger number of lower frequency sublineages. In some cases, the number of major subclonal lineages (subclonal diversity) identified by scDNA-seq is associated with the subtype of tumor. For example, there is greater subclonal diversity in estrogen receptor-negative than -positive breast cancers (9) and greater subclonal diversity in acute myeloid leukemias (AML) harboring *FLT3* mutations than non-*FLT3* AMLs (117). In one study, higher subclonal diversity (4 or more subclones) correlated with a worse prognosis (118), suggesting that subclonal diversity may be a useful prognostic biomarker.

While ITH can be inferred from bulk DNA sequencing data (178), scDNA-seq provides a more comprehensive and higher-resolution view of ITH and clonal evolution (8). Subclonal structures derived from bulk DNA and scDNA-seq are largely concordant, but the latter often reveals subclones not found in bulk data (9, 67, 162). scDNA-seq can also profile

ITH when sample size is limiting, such as fine needle aspirate biopsies (81) or when tumor cells are infrequent in the sample (i.e. low tumor purity). Nevertheless, bulk and single-cell methods are usually both implemented in integrated analyses due to the lower cost and currently higher genome-wide quality of bulk sequencing, especially for broader multi-region and multi-timepoint profiling.

Phylogenetic analyses of ITH profiles obtained by scDNA-seq reveal the order in which specific driver mutations and types of mutations occur, which in turn provides important information about the earliest events and mutational processes that initiate cancer. For example, several scDNA-seq studies have shown that in some breast cancers, colorectal cancers, and acute lymphoblastic leukemias, large numbers of structural variants (CNAs and aneuploidy) occur over short time periods early during tumor evolution but infrequently later (termed ‘punctuated clonal evolution’), while point mutations accumulate continuously along with a small number of later-occurring focal CNAs involving specific driver genes (50, 51, 94, 123, 137, 162). This suggests that genome-wide, high-impact mutational events underlie the origins of some cases of these cancer types rather than a gradual accumulation of mutations. Indeed, the mechanism of one such mutational process, chromothripsis, has been studied *in vitro* using scDNA-seq (177). In a large scDNA-seq study of AML, mutations in *DNMT3A* and *IDH1/2*, epigenetic modifier genes, were identified as the most common initiating events, while mutations in signaling genes tended to occur later (117). scDNA-seq has also quantified statistically significant co-occurrence of driver mutations at the single-cell level, for example, *NPM1* and *FLT3* mutations in AML (118), as well as marked copy number heterogeneity of focal amplifications within individual tumors: cells within one brain tumor varied from < 20 to > 100 copies of *EGFR* (45).

Interestingly, some of the above studies identified within the tumor sample near-normal cells that are unrelated to the tumor lineage but which nevertheless contain a small number of CNAs or only one driver mutation (e.g., *APC*-only mutant cells) (9, 50, 86). This likely reflects baseline copy number and driver gene abnormalities at low levels in normal cells (9, 50, 86). The relationship of such cells to the origins of cancer is presently unknown and is an important topic for further research. scDNA-seq may eventually allow identification of the cells of origin of individual tumors and help resolve long-standing debates regarding the existence and role of stem-cell hierarchies in tumors (8, 124). Other mechanisms of tumor evolution may be revealed in tumor types that have not yet been studied using scDNA-seq. Systematic application of scDNA-seq to all cancer types—a single-cell DNA Cancer Genome Atlas—in addition to other single-cell omics programs (139), will be needed to achieve a full understanding of cancer evolution.

Much of the mortality of cancer is due to metastatic spread beyond the primary tumor site. scDNA-seq has been used to study the lineage relationships between metastases and the primary tumor in the hopes of finding ways to intercept this lethal process. One scDNA-seq study of ovarian cancer found that most intraperitoneal sites of tumor spread are either monoclonal or derive from a single tumor clade (116). This indicates that once an intraperitoneal metastatic site is established, it is usually not reseeded by new cancer cells. The authors also found one patient with two subclones that were present together in multiple intraperitoneal sites with correlated frequencies, possibly due to co-migration of

the two subclones to multiple locations (116). The mostly monoclonal spread seen in the prior study contrasts with a scDNA-seq study of invasive ductal breast carcinoma that found that most tumor subclones evolve within the breast duct, followed by multiclonal invasion into the surrounding tissue (24). In another study of colorectal cancer, one patient had monoclonal metastatic seeding of the liver, while another patient had polyclonal seeding (86). These divergent findings in different tumor types make clear that the mechanisms and bottlenecks of clonal spread delineated by scDNA-seq vary across tumor types and anatomic environments.

scDNA-seq of circulating tumor cells (CTCs) is a promising approach to non-invasively sample tumors (8). CTCs are especially appealing in relation to more widely used cell-free DNA approaches, because they allow examination of complete tumor cell genomes. CTCs have been successfully isolated and sequenced from diverse cancers, including prostate cancer where they captured early clonal mutations shared by primary and metastatic sites (97), and in lung cancer where mutations were identified in CTCs that could inform treatment (125). CTCs have also been used to non-invasively monitor treatment response (28). However, CTCs are very rare in blood and require both specific tumor cell surface markers, which are not currently available for most tumor types, and a sensitive and specific isolation procedure. In a study of 51 blood samples from 36 prostate cancer patients, 27% of samples had no detectable CTCs, and only a median of 7 CTCs were found in the remaining samples (97). This poses a challenge for CTC profiling to capture the full clonal heterogeneity of a tumor to reliably inform clinical decision-making. Progress will depend on new methods to purify CTCs from large blood volumes and validation of new markers for CTC isolation. These investments will be worthwhile, as they may enable pre-symptomatic detection of tumors, non-invasive treatment monitoring, and early detection of recurrence.

scDNA-seq studies have examined how ITH changes during clinical treatment and how tumors evolve treatment resistance. One study showed that treatment-resistant clones in breast cancer were detectable prior to treatment (75). Another study used multi-omic scDNA-seq plus immune profiling to observe the phenotypic changes that mutant blast cells underwent following treatment with a drug that induces differentiation to a mostly erythroid phenotype (31). This study highlights the potential of scDNA-seq to elucidate how specific lineages with different genotypes can respond differently to the same treatment. If applied broadly, this approach will provide a critical reference for translating subclonal genotypes to predictions of treatment response. However, two limitations of scDNA-seq for treatment monitoring are noteworthy. First, tumor sampling during treatment is usually only possible in discrete, infrequent timepoints (though CTC profiling may mitigate this). Second, genetics is not the sole driver of ITH or treatment resistance; epigenetic heterogeneity also plays an important role in tumor cell biology (see section on 'DNA modifications') (75, 122).

While scDNA-seq studies of cancer have already been impactful, new scDNA-seq technologies are emerging with multi-omic, spatial, and high-throughput capabilities that will transform cancer research. Multi-omic scDNA-seq can build genotype-phenotype correlations to understand how specific subclonal genotypes associate with cellular phenotypes, invasiveness, treatment response, and more. One such method, scTrio-seq,

that simultaneously profiles CNA, methylation, and transcriptomes, found that subclonal lineages defined by CNAs and methylation are concordant, and it further correlated methylation patterns with gene expression (13). Another recent method combined scDNA-seq with oligonucleotide-tagged antibodies to simultaneously profile surface protein expression to correlate genotypic and phenotypic ITH in AML (31, 117). Mitochondrial mutations have also been combined with ATAC-seq for phenotypic lineage tracing of leukemia and colorectal cancer (83). Many new multi-omic methods continue to be developed (122). scDNA-seq is not only being combined with other molecular modalities, but also being applied in ways that preserve spatial information. One study combined laser microdissection with scDNA-seq to show how ITH varies between pre-invasive and invasive regions in breast cancer (24). Finally, the throughput of scDNA-seq is rapidly increasing, an essential step towards achieving high-resolution views of ITH and tumor evolution and for cost-effective profiling of large cohorts. Nanowell- and droplet-based methods have scaled scDNA-seq to thousands of cells per sample with a concomitant increase in ITH resolution (81, 117). This scale finally brings scDNA-seq to the throughput regime of scRNA-seq and shifts the bottleneck to sequencing costs that currently limit these methods to low genome coverage or targeted loci sequencing.

Translation of scDNA-seq to the clinic is a prime goal for delivering on the promise of cancer genomics to improve patient outcomes (8, 124). Three important steps will help make this a reality. First, clinical trials will need to rigorously measure the potential benefits of scDNA-seq profiles of ITH and clonal evolution for assigning patients to personalized or targeted treatment regimens and for assessing treatment response. Second, the current costs and technical challenges associated with scDNA-seq will need to be reduced and simplified to deliver high-throughput and user-friendly workflows that can be implemented reproducibly in a clinical laboratory. Third, while scDNA-seq CNA profiling of formalin-fixed paraffin-embedded (FFPE) samples has been achieved (109), methods for profiling single-cell genomes at high resolution from FFPE samples are needed as FFPE is the main way that clinical samples are processed. These and additional challenges remain, but the potentially life-saving benefits of scDNA-seq for cancer patients make these worthy goals.

DNA modifications

A sizable fraction of the human genome contains regulatory information that dictates the spatial and temporal expression of genes (36). DNA modifications provide an additional means to precisely control gene expression by modulating the biochemical interaction between genomic DNA and transcriptional machinery (100). The most common mammalian DNA modification is 5'-methylcytosine (5mC) and its oxidized forms 5'-hydroxymethylcytosine, 5'-formylcytosine, and 5'-carboxycytosine. Although earlier literature focused on studying 5mC dynamics in CpG dinucleotide islands (CGI) (gene-regulatory regions with increased density of 5mC-marked CpG nucleotides), the development of whole-genome bisulfite sequencing (WGBS) and its application to diverse human tissues has revealed pervasive 5mC dynamics in genomic regions outside CGIs (144, 184). For example, local 5mC depletion is a reliable signature for enhancers and other types of regulatory elements, and WGBS has uncovered other types of 5mC features such

as partially methylated domains (PMDs), DNA methylation valleys/canyons, and abundant 5mC in non-CpG contexts (54).

Cell-type-specific patterns of 5mC have historically been achieved by isolating bulk populations of specific cell types using fluorescence-activated sorting. However, this approach is limited by the availability of genetically-encoded cell type reporters or antibodies used for purification in animal models and human tissues, respectively. These challenges motivated the development of techniques for single-cell profiling of DNA modifications. Currently, there are diverse single-cell cytosine methylome methods that have been well reviewed previously (73). A series of multi-omic approaches have also been developed that combine single-cell transcriptome, chromatin accessibility, or chromatin conformation together with 5mC profiling (182).

Most single-cell methylome methods produce sparse data that randomly cover 5%-40% of the haploid genome. Notably, an important feature of methylome profiling is that coverage dropout can be unequivocally determined and is independent of the methylation state. The analysis of single-cell methylomes typically takes advantage of the local correlation of methylation patterns: the methylation level of CpG sites within a CGI region, or in non-CpG contexts in different parts of a gene body, are often correlated. This allows extraction of cell-type specific and dynamic methylation information from sparse genome-wide 5mC measurements and enables unbiased classification of cell types or transient cell populations. Typically, the quantification of 5mC in single cells over discrete features such as enhancers requires the reconstruction of pseudo-bulk profiles derived from pooling the data from a larger number of single cells of each type. Several computational approaches have also been developed to impute missing values in single-cell methylomes and to perform clustering analysis to identify cell populations (128).

Several fields in which single-cell methylome profiling has been applied include embryonic development, brain development, hematopoiesis (68), and cancer—processes involving major 5mC dynamics. A series of studies has investigated the single-cell methylation dynamics of germ cell development (88), pre- and post-implantation embryos of mouse and human (183), and mouse gastrulation (4). The advantage of single-cell methods was demonstrated by the classification of single cells into distinct lineages, enabling the quantification of 5mC and chromatin accessibility at lineage specific enhancers (4).

The human brain contains a large diversity of functional regions and cell types that are established during brain development. Single-cell methylome profiling has provided a means to simultaneously categorize brain cell types and to identify cell-type specific regulatory elements (101). Two methods, snmC-seq and sci-MET, have been applied to mammalian brains (101, 103, 119). As part of the BRAIN Initiative Cell Type Census Network (BICCN), single-cell methylomes are being systematically generated across the entire mouse brain, and the first report analyzed 100,000 cells from 45 regions of the anterior mouse brain (93). These and other studies of the human brain have demonstrated similar resolution of cell-type classification using single-cell methylomes as achieved using single-cell transcriptomes (103). Interestingly, a single-cell methylome dataset generated

from finely dissected mouse brain revealed nuanced 5mC dynamics within cell types that correlates with the spatial organization of the brain (93).

Many cancer types are associated with genomic alterations and aberrant 5mC patterns (70, 122), providing an exciting opportunity to associate epigenomic heterogeneity with tumor lineages. In addition, the possibility of reconstructing a history of epigenetic alterations (or epimutations) has motivated single-cell methylome studies comparing primary versus metastatic tumors (13) and circulating tumor cells (53). Single-cell methylome profiling of colorectal cancer has found consistent 5mC patterns in both primary and metastatic tumors within each tumor sublineage, but different 5mC profiles across lineages (13). Combined single-cell methylation and transcriptome profiling of chronic lymphocytic leukemia cells found a consistently increased rate of 5mC alteration (epimutations) compared to normal B cells, and the tumor cell lineage reconstructed using 5mC changes accurately represented somatic mutation patterns (49). Note that in this section we have focused on single-cell profiling of DNA modifications; applications of single-cell chromatin accessibility have been well reviewed previously (73, 76).

Genome organization and chromatin conformation

The folding of the two-meter long human diploid genome into a nucleus measuring less than ten micrometers in diameter is a complex hierarchical process. The resulting three-dimensional organization of the genome, also known as chromatin conformation, plays an important role in genome function. In interphase nuclei of eukaryotes, chromosomes occupy distinct spatial domains called chromosome territories. The development of proximity ligation-based chromatin conformation capture (3C) and the derived Hi-C methods can associate chromatin structures with DNA sequences to obtain a view of genome organization (91). These methods perform restriction digestion of crosslinked nuclei followed by ligation to capture DNA ends that are proximal in three-dimensional space (135). Hi-C methods have revealed a series of hierarchical chromatin domains such as topologically associating domains (TADs) that are hundreds of kilobases to megabases in length and generally conserved between cell types (33). TADs often contain finer-scale structures such as sub-TADs, domains, or individual loops, which in contrast to the relative invariance of TADs, can significantly differ between cell types and across development, suggesting that chromatin conformation dynamics is a critical component of gene regulation in normal and diseased tissues (32).

Single-cell profiling of chromatin conformation has generated novel insights into the process of chromatin organization (149, 152). Single-cell studies of chromatin conformation dynamics during the cell cycle and the oocyte-to-zygote transition have revealed that TADs are highly heterogeneous among individual cells and that TADs previously observed in bulk samples arise via chromatin contacts that are constrained but nevertheless stochastic at the single-cell level (44, 121). Another exciting application of single-cell chromatin conformation profiling is to connect distal enhancers to their regulatory target genes in specific cell types. Genome-wide association studies (GWAS) have found a strong enrichment of disease-associated genetic variants in enhancer elements (111). Cell-type-specific chromatin loop information therefore provides a valuable resource to

mechanistically dissect disease risk loci by connecting non-coding variants to disease risk genes. Single-cell 3C profiling of mouse and human brain cells has demonstrated robust cell-type identification using chromatin conformation (85, 151), supporting the feasibility of reconstructing high-resolution chromatin interaction maps for constituent cell types of mammalian tissues that contain heterogeneous cell populations. Single-cell chromatin conformation profiling will become a powerful approach for fine mapping and functional studies of disease-associated variants in the post-GWAS era.

Different strategies for single-cell chromatin conformation profiling continue to be refined and developed. While single-cell Hi-C methods, which use biotin pulldown of contact junctions, enrich for sequencing reads informative of long-range interactions (121, 149), 3C-based methods that perform ligation immediately after restriction digestion (hence avoiding biotin pulldown) provide greater sensitivity and detect substantially more chromatin contacts in individual cells (44, 152). The trade-off is that 3C-based libraries contain a lower fraction of reads containing a ligation junction and thus increase the assay cost. The throughput of single-cell chromatin conformation studies may also be further increased using a combinatorial indexing approach (134). Furthermore, recently, multi-omic methods have been developed for joint chromatin conformation and methylome profiling of single nuclei (85, 87). Genomic architecture in single cells can also be reconstructed using a method named Genome Architecture Mapping that slices nuclei into sub-micrometer thin sections followed by sequencing of each section (11).

Microbiology

The diversity of microorganisms (bacteria, archaea, and unicellular eukaryotes) is vast but remains largely unknown because most (>99%) have not been successfully cultured (148). Bulk DNA metagenomics and ribosomal DNA sequence surveys have advanced taxonomies of microbial diversity, but their reliance on small genomic fragments limits their ability to identify many organisms and to associate genetic pathways with specific microbes (148, 170). scDNA-seq, however, is a culture-independent approach to resolve microbial diversity even in complex microbial communities. The most important applications of scDNA-seq to microbiology have been to expand the census of microorganisms (especially those that are rare and cannot be cultivated), resolve uncertainties in microbial phylogenies, study the physical associations between microorganisms, associate specific genetic pathways to taxa, and to discover genetic pathways that may be biomedically useful (15, 84, 148, 170).

scDNA-seq was initially used to sequence small numbers of culturable bacteria as a proof of principle (132, 179), followed by uncultivated microbes from oral and soil samples (107, 130). scDNA-seq has since been used to discover numerous new bacterial clades and even superphyla by profiling diverse environmental samples, such as tropical oceans around the world, hydrothermal sites, freshwater bodies, and underground mines (72, 129, 138). In the largest scDNA-seq microorganism study to date, more than 12,000 cells from tropical and subtropical oceans revealed many that could not be assigned to known taxonomies (1.2%, 2.3%, and 11% had no known class, order, or family) (129). One study identified the first example of lateral gene transfer between eukaryotes and archaea (138), a discovery which could not have been made with bulk metagenomics. scDNA-seq of uncultured protists has

even been able to identify genetic sequences of bacteria and viruses that infected the cells (175). The remarkable diversity of microorganisms in almost every environmental sample (170), motivated the creation of a droplet-based method capable of profiling > 50,000 cells, which was then used to characterize the taxonomic distribution of antibiotic resistance genes within a complex seawater sample (82). While challenges remain, such as varying lysis efficiency and uneven genome amplification of some microbes (15), continued application and development of high-throughput scDNA-seq methods promises to greatly advance our knowledge of microbial diversity.

Conclusions and future applications

The applications of scDNA-seq reviewed here are only early forays heralding a rapid growth of future applications that will provide deep biological insights and impactful biomedical uses. The continued advancement of single-cell genome coverage, throughput, and multi-omic approaches will provide opportunities to not only enhance the resolution of developmental lineage trees, germ cell mutation patterns, cancer evolution, genome function, and microbial communities, but also to explore entirely new questions. While unexpected applications will certainly emerge, here, we briefly speculate regarding future and emerging applications of scDNA-seq that were not already addressed in prior sections.

Forensic analysis of low-input DNA samples can be complicated by incomplete genotypes and mixtures of DNA from different individuals. Recent work has shown that scDNA-seq may be useful for separating contributing DNA from mixed semen swab samples (154), and we anticipate increasing use of scDNA-seq in forensics. Outside the setting of forensics, scDNA-seq poses genomic privacy concerns, since near complete or complete genomes of individuals may feasibly be recovered from even the smallest numbers of cells constantly shed into the environment.

Future environmental uses of scDNA-seq include surveillance of pathogens, ecological changes, and antibiotic resistance of microbial species (for example in sewage and diverse ecosystems). In plant biology, scDNA-seq of plant gametes has enabled the generation of crossover maps without an extensive breeding program (35), which may enable more rapid methods of plant breeding. Immunology will benefit from application of scDNA-seq to characterize immunologic lineage expansions and selection in the context of infection and auto-immunity. And further studies of basic genome function such as recently developed single-cell genome replication timing (60) are emerging.

The full potential of scDNA-seq for biomedical research is still constrained by the technical limitations of current methods. Many of the technical challenges of scDNA-seq derive from the fact that it is not only a single-cell method, but also a single-molecule method: with the exception of mitochondrial DNA, each genomic allele is present in one copy per cell. We predict that the field of scDNA-seq will continue to focus heavily on technology development, but that technologies that are robust and scalable will soon be commercialized and adopted by the wider research community, as has occurred for single-cell transcriptomics. With a larger user base, including clinical adoption, we anticipate the

applications of single-cell DNA sequencing will broaden and will continue to expose new dimensions of biology at its most basic unit of life.

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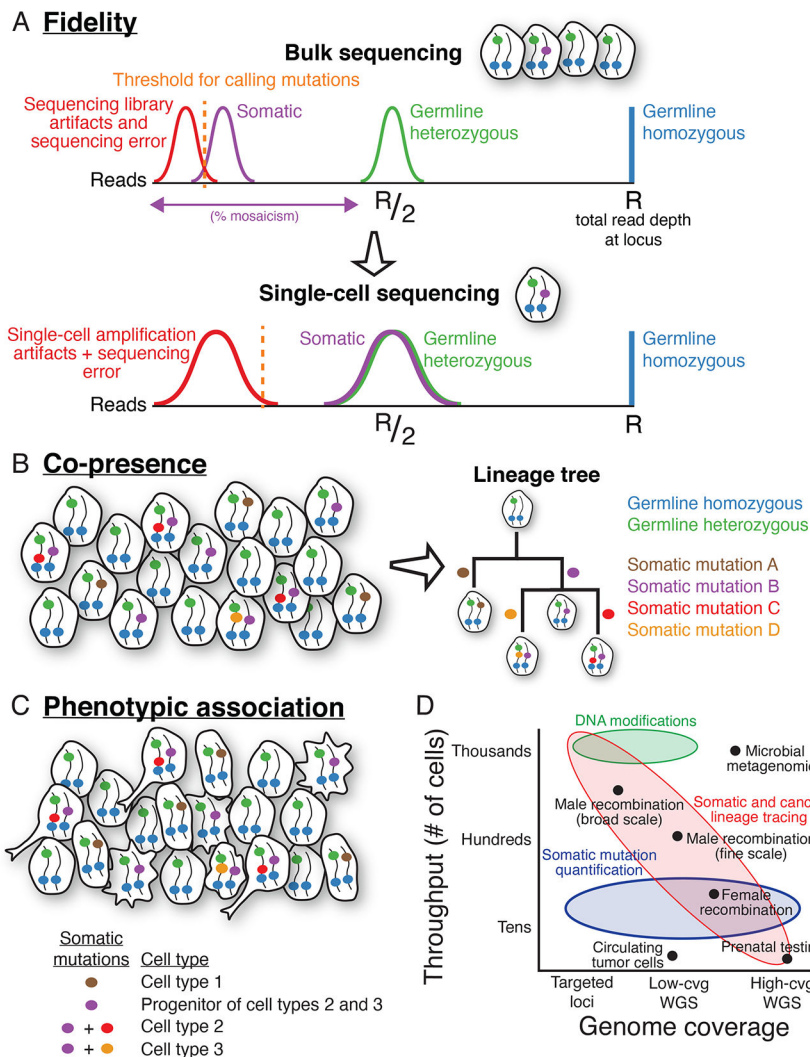


Figure 1. The three core capabilities of single-cell DNA sequencing and application requirements of throughput versus genome coverage.

A-C: The three core capabilities of single-cell DNA sequencing (Adapted from ref. (42)). A, Fidelity: In bulk sequencing, very low-level mosaic mutations cannot be distinguished from sequencing library and sequencing error artifacts. In contrast, in single-cell sequencing, the distribution of read depths is the same for somatic mutations and germline (inherited) heterozygous mutations. This enables detection of somatic mutations regardless of mosaicism, albeit it requires the same total sequencing costs as a hypothetical bulk sequencing method with perfect fidelity due to the need to sequence many single cells. Additionally, single-cell sequencing generally has an increased level of artifacts due to the single-cell amplification process. B, Co-presence: Single-cell sequencing preserves information regarding which somatic mutations are present together in the same cells. This enables reconstruction of lineage trees. C, Single-cell sequencing, when combined with simultaneous cell phenotyping (multi-omic scDNA-seq), preserves information regarding which somatic mutations are present in which cell types. This enables deconvolution of cell types or species in heterogenous samples and annotation of lineage trees with cell

phenotypes. D, Schematic of approximate application-specific requirements and current technological capabilities in terms of throughput (number of single cells per experiment or project) versus genome coverage. Low/High-cvg WGS, Low/high-coverage whole genome sequencing.

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