Title
Single cell analysis: the new frontier in 'Omics'

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Single cell analysis: the new frontier in ‘Omics’

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Cellular heterogeneity arising from stochastic expression of genes, proteins, and metabolites is a fundamental principle of cell biology, but single cell analysis has been beyond the capabilities of ‘Omics’ technologies. This is rapidly changing with the recent examples of single cell genomics, transcriptomics, proteomics, and metabolomics. The rate of change is expected to accelerate owing to emerging technologies that range from micro/nanofluidics to microfabricated interfaces for mass spectrometry to third- and fourth-generation automated DNA sequencers. As described in this review, single cell analysis is the new frontier in Omics, and single cell Omics has the potential to transform systems biology through new discoveries derived from cellular heterogeneity.

Single cell analysis: needs and applications

Cellular heterogeneity

Cellular heterogeneity within an isogenic cell population is a widespread event [1, 2]. Stochastic gene and protein expression at the single cell level has been clearly demonstrated in different systems using a variety of techniques [3-5]. Therefore, analyzing cell ensembles individually with high spatiotemporal resolutions will lead to a
more accurate representation of cell-to-cell variations instead of the stochastic average
masked by bulk measurements. Disconnect between single cell and average cell
measurements is exemplified in Figure 1a. Using an integrated microfluidic bioprocessor
for single cell gene expression analysis, Mathies group showed that siRNA knockdown
of GAPDH gene expression led to two distinct groups of individual Jurkat cells-partial
knockdown (~50%) and complete knockdown (~0%). The average result from 50 cells
(~21%) is not representative of any one individual cell [6].

To fully understand the cellular specificity and complexity of tissue
microenvironments under physiological conditions, it is necessary to measure molecular
signatures with single cell resolution. A clear example is provided by the recent work
from Kim and colleagues, who analyzed single cell gene expression profiles using high-
resolution confocal microscopy and correlated them with known cell lineages in
*Caenorhabditis elegans* [7]. The group generated expression profiles of 93 genes in 363
specific cells from L1 stage larvae. Cells were clustered into groups in a two-dimensional
scatter plot according to their correlation in gene expression (Figure 1b). Two features of
the scatter plot stand out: first, cells are diverse, but cluster with known fates such as
muscles and neurons; second, cells from homogeneous tissue (e.g. intestinal cells) cluster
more tightly than those from heterogeneous tissue (e.g. neurons). However, even within
the homogeneous tissue, individual cells show clear heterogeneity. Single cell analysis
(SCA) will therefore be critical for elucidating cellular diversity and heterogeneity.

**Potential applications of single cell analysis**

At the simplest level, SCA reduces biological noise. It provides fundamental
improvements in experimental design and data analysis for applications predicated on single cells. Stem cells, for example, hold great potential for regenerative medicine because they can self-renew and differentiate along different lineages. However, embryonic stem cells, adult stem cells, and induced pluripotent stem (iPS) cells are all heterogeneous populations [2, 8, 9]. SCA can target specific populations and therefore elucidate signaling pathways and networks for self-renewal and for differentiation. Cancer is a heterogeneous disease and dissecting cell-to-cell variations is extremely important in understanding tumor initiation, progression, metastasis, and therapeutic responses. For example, cancer cells have recently been shown to harbor homoplasmic and heteroplasmic mutations in mitochondrial DNA (mtDNA) that are above and beyond the widespread heterogeneity of mtDNA in normal human cells [10]. SCA may functionally differentiate normal and cancer cells and also cancer cells at various development stages. Interestingly, the key therapeutic targets in tumors may be cancer stem cells [11], which represent a small percentage of the total mass of tumors, but may be responsible for tumor repopulation following treatment [12]. SCA has the potential to more accurately identify these cells and their unique susceptibilities. Neurons are the basic unit of the nervous system. Their electronic properties have been well-characterized by electrophysiology, but the molecular complexity of ion channels, neurotransmitters, and neuropeptides at the single cell level and even the subcellular level is only beginning to be understood [13]. SCA can help elucidate neural communication in unprecedented detail, which may yield new strategies to understand and treat neurological disorders.

Advances in SCA have the potential to accelerate not only biological research, but also diagnostics. Preimplantation genetic diagnosis (PGD), for example, is the analysis of
a single cell from a biopsy of an embryo after in vitro fertilization. PGD is used to test for
genetic diseases and chromosome aneuploidies. The most common molecular analysis
techniques are PCR and FISH (fluorescent in situ hybridization) [14], and newer
methods, such as comparative genomic hybridization (CGH) [15], are beginning to be
utilized. As more genetic diseases and predispositions are identified, multiplexed PGD
using single cell Omics technologies will enable early and accurate identification and
possible treatment of genetic abnormalities in human embryos.

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Single cell Omics: current state-of-the-art

Omics for single cell analysis

‘Oomics’– the large-scale studies of genes (genomics and epigenomics), transcripts
(transcriptomics), proteins (proteomics), metabolites (metabolomics), lipids (lipidomics),
and interactions (interactomics) – is both the foundation and driving force for systems
biology [16]. Omics strives to identify, quantify, and characterize all of the components
in cellular systems with spatiotemporal resolution, and thereby dissect the intracellular
pathways and networks. Currently, major technologies for genomics, epigenomics, and
transcriptomics include DNA sequencing and microarrays (planar-, bead-, and fiber optic
arrays); for proteomics, mass spectrometry (MS) and protein arrays; and for
metabolomics, MS and NMR.

The abovementioned major technologies have undergone dramatic advances in
automation and miniaturization. The logical limit to the ever-increasing miniaturization
of Omics technologies is the comprehensive spatiotemporal analysis of genes, transcripts,
proteins, metabolites, and interactions in single cells and their subcellular compartments.
The movement of Omics into SCA represents a significant shift. Previous well-established methods for SCA, such as imaging and flow cytometry, are limited to the examination of a small number of genes, proteins or metabolites. As a result, these methods can only be used to open narrow windows into the complexity and dynamics of intracellular pathways. By contrast, single cell Omics has the potential to enable systems biology at the level of single cells, ultimately representing the unique convergence of technology and biology.

**Nucleic acid analysis**

Nucleic acid analysis has already reached the single cell level [4, 17-21]. Newer technologies, such as ‘lab-on-a-chip’ [22], have further advanced the field. For example, Mathies and colleagues have shown a hybrid (PDMS/glass) microfabricated bioprocessor for single cell gene expression analysis [6] (as discussed above). In addition, Quake and colleagues have used PDMS structures and on-chip PCR to realize large-scale integration for applications in single cell mRNA isolation and analysis [23]. Quake’s technology has been commercialized by Fluidigm Corporation, demonstrating the promise of miniaturization for single cell Omics. The Fluidigm Dynamic Array™ enables gene expression measurements in individual cells, and represents the first dedicated commercial product for SCA. This technology, however, needs greater multiplexing capabilities for global transcriptomic analysis. It currently can test up to 96 individual cells against 96 genes. The number of genes needs to at least increase by two orders of magnitude for routine transcriptome profiling.

Further improvement in the breadth of single cell mRNA analysis has been
achieved recently using either mRNA sequencing (mRNA-Seq) [24] or quantitative PCR (qPCR) [25]. In the first study, the Surani and Lao groups together developed an assay with mRNA-Seq [26] for single cell whole-transcriptome analysis; in this method, single cell cDNA amplification [27] was improved from a previous version by increasing the cDNA length up to 3 kb. Briefly, the mRNAs from a single cell were reverse-transcribed into cDNAs, which were subsequently amplified by PCR. Finally, amplified cDNAs were digitally counted with an ultrahigh-throughput DNA sequencer. The researchers first verified reproducibility of the method, and then demonstrated its utility in dissecting functional consequences of Dicer1 and Ago2 knockdowns, two of the critical genes for miRNA synthesis and mRNA regulation, by comparing single mouse Dicer1−/− and Ago2−/− oocytes versus wild-types (Figure 2a). The described method has the added advantage over probe-based methods (such as qPCR) in that it is effective at locating splicing variants, including de novo variants, but may face challenges in accurate quantitation due to the required amplification process.

To improve mRNA quantitation from single cells, the Kambara group performed direct qPCR from a cDNA pool without pre-amplification [25]. Gene expression for four house-keeping genes (TBP, SDHA, B2M, and EEF1G) was quantitatively analyzed in 14 single cells, showing that the absolute amounts of each of the four genes differed from cell to cell (Figure 2b). They further showed that the average number of cDNA molecules obtained from single cells was comparable to the per-cell number measured from the cell pools (10-1000 cells); however, much larger standard deviation was observed for the single cells, as expected from the cellular heterogeneity. The research group is currently developing higher-throughput methodology to analyze more individual
cells (personal communication). A larger sample size will enable better statistical analysis of cell-to-cell variation. The method would also benefit from increased multiplexing of the capture beads for the immobilized cDNA library to increase transcriptome coverage. Single cell genomics has been achieved for microorganisms. Church and colleagues have sequenced the genome of single cells of *Escherichia coli* and *Prochlorococcus* through so-called polymerase cloning (“ploning”), an optimized version of multiple displacement amplification (MDA) using random primers [28]. They obtained ~65% sequence coverage at an average depth of 4-fold for a single *Prochlorococcus* cell through whole-genome shotgun sequencing. However, the method may suffer background contamination during the amplification of single cell DNA and difficulty in assembling the genomes. Compared to microorganisms, which typically have a genome size of several million base pairs containing a few thousand genes, mammalian genomes are much more complex. For example, the human genome has 3 billion base pairs containing estimated 20,000-25,000 genes. To the best of our knowledge, there has been no report yet on single cell genomics (i.e., global analysis of chromosomal DNA) or epigenomics (i.e., global analysis of DNA methylation etc) for mammalian cells. However, this may change soon with the recent introduction of several next-generation sequencing technologies (discussed in the next section), particularly those centered on single molecule sequencing.

**Protein and metabolite analysis**

Proof-of-principle SCA has been performed to characterize peptides using matrix-assisted laser desorption ionization (MALDI) MS [29-31], or to characterize small
molecules using electrospray ionization (ESI) MS [32-34]. The Sweedler group previously profiled peptides directly from single neurons and individual organelles using MALDI-MS [35, 36]; however, the number of peptides detected was limited. Recently, the group was able to couple capillary electrophoresis with ESI-MS (CE-ESI-MS) and achieved reproducible metabolomic analysis of an isolated single R2 neuron and its different subcellular regions [33]. As shown in Figure 3a, different subcellular regions of a single R2 neuron (soma vs. neurite) have different relative amounts of several metabolites with the extracted ion of 146 m/z. However, this method requires offline manual isolation and lysis of single cells in a vial before the front-end CE separation followed by MS, thereby posting a significant challenge for in situ metabolomic analysis.

In addition, CE sample injection is dictated by capillary inlet and outlet of the sampling interfaces; currently only 0.1% of the total content of a single metacerebral cell, corresponding to 6 nL of the cell lysate, was injected (although more than 100 compounds were detected). Increasing the injected amount of cellular contents may improve the metabolome coverage from a single cell.

More recently, laser ablation electrospray ionization mass spectrometry (LAESI-MS) was demonstrated for in situ metabolomic profiling of single cells [34] (Figure 3b). Single cell ablation was achieved by delivering mid-IR laser pulses through a GeO\textsubscript{2}-based glass fiber (15 μm radius at the tip). When comparing two physically adjacent, individual Allium cepa cultivar cells with differences in purple pigmentation, LAESI-MS revealed similar essential metabolites for both variants, with colorless cells containing anthocyanin and pigmented cells containing anthocyanidins, other flavonoids, and their glucosides. One of the key advantages of this method is that single cells can be analyzed
in situ, although, as mentioned by the authors, further improvement in optics and ablation geometry is needed for cell-by-cell analysis of biological tissue. Another challenge for the LAESI-MS is the lack of front-end separation of cellular contents using either LC or CE, because laser ablation is performed directly on whole cells. This hinders further improvement in metabolome coverage and identification.

Previously, flow cytometry (i.e. fluorescence-activated cell sorting, or FACS) has been used to profile phosphoprotein networks in single cancer cells [37] and monitor global protein levels in single Saccharomyces cerevisiae [38]. In addition, fluorescence imaging has been used to measure the levels and locations of nearly 1,000 different endogenously tagged proteins in individual living cancer cells at high temporal resolution [39]. Current technologies based on fluorescence require prior in vivo or in vitro labeling, and are generally limited to 10-20 simultaneous measurements due to the spectral overlap. For in vitro labeling, background arising from cross-reactivity and/or nonspecific binding remains a challenge. For in vitro labeling, tagging with GFP/YFP and their derivatives may alter folding or other characteristics (e.g. interactions with other proteins) of the endogenous proteins. Furthermore, photobleaching of fluorophores and autofluorescence of cells may interfere with kinetic measurements. A potential solution is MS-based flow cytometry (‘mass cytometry’), which has been described for a single cell multi-target immunoassay [40]. However, the method still requires labeling of antibodies with elemental tags (e.g. lanthanide) prior to antigen binding.

Mass spectrometry-based single cell proteomics has not been achieved yet despite tremendous interest in the field. Recently, an LC-MS/MS-based method was described for quantitative proteomic analysis of single pancreatic islets containing 2,000-4,000 cells
[41]. A combination of improved chromatographic methods, direct replicate measurements, and high accuracy (1-2 ppm) and sensitivity (attomole to femtomole) provided by a LTQ-Orbitrap mass spectrometer, allowed for detection of more than 6,000 proteins from nanogram quantities of protein mixtures. This ultrasensitive method represented an important step toward single cell proteomics. However, due to the extreme complexity and huge dynamic range of the proteome, new breakthroughs in sample manipulation and detection are needed to achieve in situ single cell proteomics with deep proteome coverage (discussed in the next section).

**Single cell Omics: future prospects**

Technologies for analyzing nucleic acids, proteins, and metabolites are evolving rapidly. This section includes notable examples that have the potential to make major contributions to single cell Omics.

*New technologies for nucleic acid analysis*

Driven by personal genomics and the NIH-launched "$1000 Genome" project, the technologies for nucleic acid analysis, particularly the next-generation sequencing technologies, have undergone dramatic advances [42-47]. Figure 4 showcases some of the technologies that may contribute to SCA.

Geiss and colleagues from NanoString Technologies, for example, have developed the nCounter gene expression system (Figure 4a) for direct multiplexed measurement of gene expression with color-coded probe pairs without amplification [42]. The technology utilizes molecular barcodes and single-molecule imaging to detect and
count hundreds of unique transcripts directly in a single reaction. The company
demonstrated that the nCounter system is more sensitive than microarrays and similar in
sensitivity to real-time PCR. The technology has the potential for single cell
transcriptomics.

In addition, multiple breakthroughs in single molecule sequencing technologies
could pave the way for single cell genomics, epigenomics, and transcriptomics. Turner
and Korlach together with colleagues from Pacific Biosciences have developed a
technology for single molecule and real-time DNA sequencing by a single DNA
polymerase (Figure 4b) [44]. They detected the enzymatic incorporation of
fluorescently-labeled deoxyribonucleoside triphosphates into the growing DNA strand
with zero-mode waveguide nanostructure arrays. They demonstrated a median accuracy
of 99.3%, with no systematic error beyond the fluorophore-dependent error rates, with
the consensus sequences generated from the single-molecule reads at 15-fold coverage.
Optimism for this technology is high: the company has raised approximately $266
million to date through a combination of grants and venture capital according to the
company’s website.

Traz and colleagues from Helicos Biosciences have developed a high-throughput
and amplification-free method for transcriptome quantification, named single-molecule
sequencing digital gene expression (smsDGE) (Figure 4c) [45]. The technology utilizes a
reverse-transcription and polyA-tailing sample preparation procedure followed by single-
molecule sequencing that generates a single read per transcript as published earlier [43].
They confirmed accurate quantification using spiked-in RNAs, and demonstrated
sequencing of the yeast transcriptome in a single run, yielding an average of 12 million
aligned reads per channel. While they have not used this technology for SCA, they recently published a ChIP-Seq study in collaboration with the Bernstein group using a small number of cells and only 50 pg of DNA [48].

Bayley and colleagues from Oxford Nanopore Technologies have developed a single molecule sequencing technology using nanopores (Figure 4d) [46]. The nanopores were constructed from hemolysin mutants that were covalently attached to an adapter molecule to detect the bases cleaved from the DNA by an exonuclease enzyme. The researchers demonstrated the continuous identification of unlabelled nucleoside 5’-monophosphate molecules with an average 99.8% accuracy. Methylated cytosine was also distinguished from the four standard bases. At the present time, the company analyzes multiple copies of nucleic acids, but is theoretically capable of single-copy detection, although longer sequences would be required because of diffusion (personal communication). Moreover, the platform could be adapted for analysis of proteins, other polymers, and small molecules, potentially accelerating multiple phases of SCA.

Table 1 summarizes the key technologies and other attributes of five major third-generation DNA sequencing companies, including three mentioned above (Helicos, Pacific Bio, and Oxford Nanopore) and two others (Complete Genomics and Ion Torrent). In addition, Life Technologies is actively developing a single molecule sequencing technology using quantum dots, and Cipriany et al. recently demonstrated a method for single molecule epigenetic analysis using nanofluidics and multicolor fluorescence microscopy to detect DNA methylation [49]. (For more information on next-generation sequencing technologies, please refer to a recent review by Michael Metzker [50]) Further improvements in detection sensitivity, sample manipulation and
recovery, and sequence assembly will enable the abovementioned technologies to become suitable for direct single cell Omics applications.

**New technologies for protein and metabolite analysis**

The “holy grail” of proteomics and metabolomics is to decode the proteome and metabolome at the single cell level. This is challenging due to the fact that the amount of proteins and metabolites is extremely limited in a single cell (i.e. average $1 \times 10^5$ molecules for proteins). Furthermore, there are no amplification methods for proteins and metabolites comparable to those for nucleic acids. We believe the most prevalent bottleneck for SCA of proteins and metabolites *in situ* is the marriage between extremely efficient sample manipulation and highly sensitive detection. Along this line, we believe micro/nanofluidics interfaced with mass spectrometry will be hotly pursued for single-cell proteomics and metabolomics. **Figure 5** showcases some promising technologies that may contribute to this endeavor.

To achieve single cell Omics, it is critical to efficiently select and manipulate single cells. Laser capture microdissection has been used routinely to isolate a small number of cells from tissue [51]. In addition, micropipette-based micromanipulators have been used to manipulate single cells. However, these technologies are difficult to implement in a micro/nanofluidic device, particularly for living single cells. Newer technologies based on droplet encapsulation may circumvent this challenge. For example, Chiu and coworkers have developed femtoliter- and picoliter-sized droplets as nanolabs for manipulating single cells and subcellular compartments [52]. **Figure 5a** shows one of their PDMS microfluidic devices for thermoelectric manipulation of single cells [53]. In
this device, the T-channel was used for droplet generation and freezing (above where a thermo-electric cooler (TEC) was placed). The same group earlier combined optical trapping, microfluidic encapsulation, and rapid laser photolysis, and demonstrated selective encapsulation of single cells and analysis of subcellular organelles through fluorescence spectroscopy [54]. More recently, high-throughput hydrodynamic encapsulation of single cells into picoliter-volume droplets on PDMS chips has been developed by others [55, 56]. Furthermore, online analysis of the contents in single droplets by MS has been demonstrated [57], suggesting promising potential of single cell encapsulation-MS analysis.

Micro/nanofluidic devices have been fabricated to directly analyze proteins from single cells or extremely small amounts of biomaterials. Zare and colleagues have demonstrated chemical cytometry as well as protein counting in a single cell using PDMS microfluidic devices [58, 59]. The analysis chips (Figure 5b) can manipulate, lyse, label, separate, and quantify the protein contents of a single cell using single molecule fluorescence counting [59]. Heath and colleagues have developed an integrated PDMS-based barcode chip (Figure 5c) for rapid multiplexed analysis of proteins in a microliter of blood [60]. The device achieved on-chip blood separation and rapid detection of a panel of plasma proteins using fluorescence-based surface immunoassays. Although no SCA has been demonstrated yet using the device, its efficient separation and sensitive detection of a very small sample amount may enable its application in SCA. Han and colleagues have developed a Si-based microfabricated nanofluidic device, an anisotropic nanofilter array (ANA), for continuous-flow separation of DNA and proteins [61]. The device may serve as a key component for sample preparation and separation in an
integrated bioanalytical system. However, all three of these methods (Zare, Heath, and Han) require labeling of proteins using fluorescent antibody binding, which limits analysis to a small number of previously characterized proteins.

Mass spectrometry, by contrast, can identify and quantify thousands of proteins and their posttranslational modifications (e.g. phosphorylation and acetylation, etc) per experiment. Wang and colleagues have developed Si/SiO$_2$-based microfabricated monolithic multinozzle emitters for mass spectrometry ($M^3$ emitters) ([Figure 5d] [62]). Si/SiO$_2$-based materials were selected because hydrophobic polymers, such as PDMS, have inherently undesirable properties for the ESI-MS application, including strong affinity to proteins and incompatibility with certain organic solvents. In their design, each $M^3$ emitter consists of a parallel silica nozzle array protruding out from a hollow silicon sliver. Once integrated with a mass spectrometer, $M^3$ emitters achieve sensitivity and stability in peptide and protein detection comparable to those of commercial silica-based capillary nanoelectrospray. The additional advantages of these emitters are ease of fabrication and potential to be integrated with other micro/nanofluidic structures. These $M^3$ emitters may serve as a critical component in a fully integrated silicon/silica-based lab-on-a-chip for single cell proteomics and metabolomics.

Key challenges still remain to implement high-quality interfaces between micro/nanofluidic chips and mass spectrometers [63], particularly for SCA. This requires further reduction in the sampling size (picoliter to femtoliter, i.e. for single cells or subcellular components), sample manipulation (i.e. efficient recovery of proteins and peptides), integration of multiple processes (i.e. lysis, separation, and detection), and dramatic improvement of the mass spectrometers (sensitivity, accuracy, resolution, and
dynamic range). If coupled with abovementioned on-chip droplet encapsulation of single cells, MS-chip technologies may open new doors to high-throughput multiplexed SCA. Therefore, breakthroughs in micro/nanofabrication methods and mass spectrometry technologies are clearly needed to develop an integrated micro/nanofluidics device for single cell proteomics and metabolomics.

**Challenges and opportunities**

Single cell Omics presents unprecedented challenges and opportunities, both technologically and biologically. There is still a great need for further improvement in miniaturization, integration, and detection sensitivity. There is also a great need for automation, throughput, and bioinformatics to study multiple individual cells to achieve statistical significance. New software packages for statistics and bioinformatics [64], for example, may have to be developed for the large amount of data generated from multiple single cells. All of these technical challenges will add costs to existing Omics methods in the short term, but miniaturization and automation are expected to enable the development of new affordable high-throughput methods. Moreover, most current Omics-based SCA methods require cell lysis, which means that genes, proteins, and metabolites are studied at specific timepoints outside of their physiological environments. This can be mitigated through live-cell imaging and other *in situ* techniques that can be performed on single cells once the key markers and pathways are discovered through SCA [65]. Furthermore, studying individual cells is inherently reductionist, which means that tissue-level properties such as interactions between cells and their extracellular environment need to be incorporated into experimental designs and data analyses [7].
While these challenges are significant, opportunities abound. At the simplest level, reducing cellular heterogeneity reduces noise, which is a goal of any analytical system. At a higher level, though, the heterogeneity itself becomes informative. As shown in Figure 1a, identifying the cells with complete GAPDH knockdown would significantly reduce the noise in analyses of GAPDH functions. Furthermore, isolating cells with both complete and partial knockdowns allows for an examination of RNA interference mechanisms and the causes of the disparate, but apparently quantized, responses. Detailed analysis of single cells using Omics technologies are expected to elucidate the underlying cellular pathways and networks.

In summary, single cell analysis is the new frontier in ‘Omics’. Single cell Omics represents the unique convergence of technology and biology and will transform systems biology by converting cellular heterogeneity from a source of noise to a source of new discoveries.
Acknowledgements

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References


27. Kurimoto, K., et al. (2007) Global single-cell cDNA amplification to provide a template for representative high-density oligonucleotide microarray analysis. *Nat Protoc* 2, 739-752


Figure Legends

Figure 1. Observation of cellular heterogeneity. (a) mRNA expression of GAPDH from individual Jurkat cells after siRNA knockdown. The levels fall into roughly two categories, 50% and 100% knockdowns (i.e. 50% and 0% expression remained). Note that the average GAPDH expression obtained from the measurement of 50 cells (21±4%) was not representative of any individual cell. Adapted with permission from Ref. [6]. (b) Clustering diagram of 363 cells in *C. elegans* according to the similarity of their gene expression profiles. Distance between cells in the x-y plane indicates levels of similarity. Colors indicate different tissue types. Key: b.w.m., body wall muscle; b.neu., body neurons; re.epi., rectal epithelial cells; b.c., other body cells; int., intestine cells; hyp., hypodermal cells; blast, blast cells; ph.m., pharyngeal muscle; ph.neu., pharyngeal neurons; ph.epi., pharyngeal epithelial cells; ph.c., other pharyngeal cells. Adapted with permission from Ref. [7].

Figure 2. Single cell gene expression analysis. (a) Correlation plots of the quantile-normalized mRNA sequencing reads for mouse oocytes, showing (i) one wild-type oocyte versus another wild-type oocyte; (ii) one *Dicer1*<sup>-/-</sup> oocyte versus another *Dicer1*<sup>-/-</sup> oocyte; (iii) one wild-type oocyte versus one *Dicer1*<sup>-/-</sup> oocyte; and (iv) one wild-type oocyte versus one *Ago2*<sup>-/-</sup> oocyte. All of the reads with changes of greater than fourfold are plotted in red. Adapted with permission from Ref. [24]. (b) Number of cDNA molecules measured by qPCR in single human colon carcinoma HCT116 cells and cell pools, showing (i) cell-to-cell variation in expression of “housekeeping” genes among 14 single cells, and (ii) average gene expression for single cells as well as cell pools with
error bars (single cell: mean ± s.d., n=14; 10–1,000 cells: mean ± s.d., n=5). Adapted
with permission from Ref. [25].

**Figure 3.** Single cell metabolome analysis. (a) Metabolomic profiling of a single *Aplysia*
R2 neuron using CE-ESI-MS, showing the extracted ion electropherogram (XIE)
obtained for 146 m/z from different subcellular regions, i.e. soma versus neurite. Adapted
with permission from Ref. [33]. (b) Metabolomic profiling using LAESI-MS, showing (i)
etched optical fiber for laser ablation relative to the target single cell (scale bar =
100 μm), and (ii) optical image of neighboring individual colorless and pigmented
epidermal cells of the purple *A. cepa cultivar* (scale bar = 50 μm). Also shown (ii-a and
ii-b) are the corresponding LAESI-MS spectra. Selected similar and different peaks (m/z)
are indicated by arrows (ii-b). Adapted with permission from Ref. [34].

**Figure 4.** New technologies for nucleic acid analysis. (a) nCounter gene expression
system from NanoString Technologies, showing (i) schematic representation of the
hybridized complex (not to scale); (ii) schematic representation of, from left to right,
binding, electrophoresis, and immobilization; and (iii) false-color image of immobilized
reporter probes. Adapted with permission from Ref. [42]. (b) Single-molecule, real-time
DNA sequencing system from Pacific Biosciences, showing (i, left) schematic
representation of the experimental geometry with a single molecule of DNA template-
bound DNA polymerase immobilized at the bottom of a zero-mode waveguide (ZMW),
(i, right) schematic event sequence of the phospholinked dNTP incorporation cycle, with
a corresponding expected time trace of detected fluorescence intensity from the ZMW;
(ii, top) total intensity output of all four dye-weighted channels with pulses colored corresponding to the least-squares fitting decisions of the algorithm, and (ii, bottom) the entire read that proceeds through all 150 bases of the linear templates. Adapted with permission from Ref. [44]. (c) Single-molecule sequencing digital gene expression system from Helicos Biosciences, illustrating sample preparation and sequencing workflow: (1) preparation of the first-strand cDNA from mRNA, (2) addition of 3’ tail of dATP followed by dideoxy-TTP (ddT) blocking, (3) hybridization of tailed sample to poly-dT oligonucleotide covalently attached to the flow-cell channel surface, (4) sequencing of a single base by adding a Cy5-labeled nucleotide, (5) cleaving off the Cy5 dye label, and (6) adding and imaging of next nucleotide. Adapted with permission from Ref. [45]. (d) Single-molecule nanopore sequencing system from Oxford Nanopore Technologies, showing (i, left) nanopore structure of the WT-(M113R/N139Q)₆(M113R/N139Q/L135C)₁ mutant with the cyclodextrin covalently attached at position 135 (space-filling model), and (i, right) close-up of the β barrel with the arginines at position 113 and the location of the cysteines in the mutants tested in the study; and (ii) single-channel recording from the nanopore that indicates discrimination of dGMP, dTMP, dAMP, and dCMP, with colored bands representative of the residual current distribution for each nucleotide. Adapted with permission from Ref. [46].

**Figure 5.** New technologies for protein and metabolite analysis. (a) A single-cell encapsulator, showing (i) the channel systems, and (ii) a string of droplets generated. Adapted with permission from Ref. [53]. (b) A single-cell analysis chip, showing the cell-manipulation section on the left and the molecule-counting section on the right.
Adapted with permission from Ref. [59]. (c) An integrated blood barcode chip, comprising channels that harness the Zweifach-Fung effect for plasma separation from a finger prick of blood as well as multiple DNA-encoded antibody DEAL barcode arrays patterned on the surface of the plasma-skimming channel. Adapted with permission from Ref. [60]. (d) Microfabricated monolithic multinozzle emitters (M³ emitters), showing (i) a schematic view of a nanoelectrospray emitter with two protruding nozzles, and (ii) SEM images and corresponding magnified views of the M³ emitters with different nozzle numbers (1-10) and dimensions. Adapted with permission from Ref. [62].
Table 1. A comparison of representative third-generation DNA sequencing companies

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b: Company websites
Figure 1

(a) Normalized GAPDH expression with siRNA

Average of 50 cells is not representative of any individual cells

(b)
Figure 2
Figure 3
Figure 4

(a) nCounter (NanoString)

(i) Capture probe

(ii) Reporter probe

(iii) Labeled RNA segments

(b) Single molecule, real-time sequencing (Pac Bio)

(i) A

(ii) B

(c) Single molecule sequencing (Helicos)

1. mRNA
2. cDNA
3. CTG
4. CTG
5. React C

(d) Single molecule sequencing (Nanopore)

(i) MT138, T115C, T17PC, G199C (2.6 Å), N121C (2.9 Å), N123C (10.3 Å), T125C (13.5 Å), L135C (9.1 Å), N139Q (17.0 Å)

(ii) Residual pore current vs. time
Figure 5