Single-Cell Transcriptomics: A High-Resolution Avenue for Plant Functional Genomics

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Plant function is the result of the concerted action of single cells in different tissues. Advances in RNA-seq technologies and tissue processing allow us now to capture transcriptional changes at single-cell resolution. The incredible potential of single-cell RNA-seq lies in the novel ability to study and exploit regulatory processes in complex tissues based on the behaviour of single cells. Importantly, the independence from reporter lines allows the analysis of any given tissue in any plant. While there are challenges associated with the handling and analysis of complex datasets, the opportunities are unique to generate knowledge of tissue functions in unprecedented detail and to facilitate the application of such information by mapping cellular functions and interactions in a plant cell atlas.

Insights Gained from Single-Cell Profiling

The functionality of complex organs and organisms is the result of an orchestrated interplay of different cell types and their specific functions [1–3]. To completely understand and identify most critical cellular processes in complex tissues therefore requires changes to be captured at the cell-type or even single-cell level (Figure 1A). Progress in cell type-specific transcriptomics represented a vital first step in this direction and helped to reveal essential cellular activities involved in plant development and stress adaptation [2–4]. Recent advances in microfluidics-based approaches to single-cell RNA-seq (scRNA-seq) (see Glossary) equip us now with a unique opportunity to study transcriptional changes at cellular resolution in any given organism [5–12]. In animal-based research, scRNA-seq has revolutionised and stimulated cellular studies. In addition to facilitating the discovery of new cell types, it enables the study of stochastic principles in cellular gene network regulation and the trajectories of transcriptomic changes underlying cell fate choices and organ functionality [13–17].

As a new technique that has thus far been substantially applied only in animal/human tissues, the true potential of scRNA-seq for diverse aspects of plant science has just started to be recognised. Here, we highlight the opportunities and challenges associated with performing scRNA-seq in plants. The scope of this review is to focus on droplet-based methods, primarily Drop-seq and 10X platforms, as these are the methods that have been applied to plant studies so far. However, analyses of data from other droplet-based methods are highly similar to what is reviewed below. Importantly, we introduce the technical principles of scRNA-seq approaches for plant-based studies, explain how scRNA-seq systems operate, and outline standard analytical workflows including underlying statistical analyses of scRNA-seq data.

Common Approaches in Analysing Cell-Specific Signalling Networks

Prior to the invention of droplet-based scRNA-seq, single-cell analyses relied on dissecting individual cells in situ using laser microdissection (LMD) or fluorescence-activated cell sorting (FACS). The disadvantage of LMD is that it is low throughput, capturing 4–40 cells, and is technically challenging [18–21]. FACS-based approaches, in turn, have advanced our understanding of gene networks on a cell-type or tissue level by employing fluorescent marker lines to isolate specific cell populations. Using FACS in combination with microarrays, Birnbaum et al. produced the first gene expression map of the arabidopsis root encompassing five cell types and three developmental zones [2]. Together with subsequent higher-resolution studies, it revealed a specific transcriptional identity for each cell type [3]. Similarly, further studies of cell type-specific responses to salt stress, iron deprivation, nitrogen depletion, varied pH levels, and immunity all revealed the activation of distinct stress responses and improved resilience capacities for the development of crops with enhanced stress resistance.

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Highlights

- Plant tissues comprise a diverse set of cell types that can be distinguished by their functions. The concerted interplay of these cell types determines the functionality and plasticity of plant tissues.
- Deciphering the different functions of cell types in a tissue is essential to understand plant development and adaptation to changing environments.
- Single-cell RNA-seq technologies enable us now to capture transcriptional profiles in each cell type to describe the genetic basis of their identity and function. This knowledge of cell-type-defining gene networks is as equally significant for fundamental science as it is for the development of crops with improved resilience capacities against climatic and other environmental stresses.
gene networks in each cell type [4,22–25]. While FACS has provided essential insights into root tissue organisation, its dependence on fluorescent marker lines has limited its use to studies of known cell type lineages in model plants (mostly Arabidopsis).

Drop-seq was the first single-cell technology to use microfluidics combined with a barcoding system to enable parallel, high-throughput sequencing of single-cell transcriptomes [26]. This groundbreaking new approach has transformed single-cell analysis in mammalian studies and is beginning to be implemented in plants [8–12].

Figure 1. Generation of Single-Cell Transcriptomic Data Using Microfluidic Technology.

(A) Complex tissue is dissociated into individual cells. Root cross-section was adapted from the Plant Illustrations repository (E. Sparks, 2017; https://doi.org/10.6084/m9.figshare.4688371.v1). (B) Microparticle beads covered with DNA probes are used to capture mRNA molecules and introduce barcodes for cells into the cDNA. Probes comprise a PCR handle, a cell barcode common to all probes on a bead, unique molecular identifiers (UMIs) enabling computational removal of PCR duplicates, and poly-dT sequences. (C) Individual cells are encapsulated in droplets together with a barcoded bead and lysis buffer. (D) Cells are lysed within droplets. (E) Reverse transcription yields cDNA molecules hybridised to probes on beads. (F) Sequencing of cDNA yields a library of transcriptomes of thousands of individual cells. (G) Software is used to count unique reads per gene and per cell yielding a digital gene expression (DGE) matrix. Created with BioRender.com.

Glossary

Barcoded beads: microparticles that carry DNA oligonucleotide probes for capturing and barcoding of mRNA transcripts.

Digital gene expression (DGE) matrix: matrix of unique read counts per gene per cell.

Drop-seq: a microfluidics-based method to quantify mRNA transcript levels in thousands of single cells by encapsulating individual cells in droplets containing barcoded beads.

Fluorescence-activated cell sorting (FACS): technique to isolate single cells based on their cell type-specific marker gene-associated fluorescence.

Index of Cell Identity (ICI): adapted algorithm used to assign known cell-type identities to single cells without relying on unique marker genes.

Laser microdissection (LMD): laser-based isolation of single cells or cell clusters from tissue for subsequent analyses.

Principal component analysis (PCA): dimensionality reduction method based on linear algebra.

Pseudotime: algorithm used to assign a cell’s position along a trajectory of transcriptomic changes such as differentiation.

Single-cell RNA sequencing (scRNA-seq): set of methods for the sequencing of transcriptomes of single cells.

Single-cell transcriptomes attached to microparticles (STAMPs): cDNA molecules derived from a single cell hybridised to probes on the surface of barcoded beads.

t-Distributed Stochastic Neighbourhood Embedding (tSNE): machine learning algorithm to compute positions in a low-dimensional (normally 2D) space for a set of high-dimensional data points such that the proximity of close data points is preserved.

Uniform Manifold Approximation and Projection (UMAP): a method for dimensionality reduction designed to preserve more of the structure of distance relationships than tSNE.

Unique molecular identifier (UMI): short DNA sequence synthesised randomly which is used to uniquely label DNA molecules.
For plant studies, processing tissue with cell walls has been a challenge for the application of single-cell technologies, often resulting in low capture rates (discussed in the following section). Currently, droplet-based single-cell methods produce relatively low resolution of transcriptomes (~10,000 reads per cell [21]) and thus the information gained from single-cell analytics can be limited to the most highly expressed genes. However, the latest single-cell technologies (e.g., the 10X platform), in combination with efficient tissue lysis protocols, have improved capture efficiency and gene detection thus increasing the resolution of single-cell analyses irrespective of tissue properties. There are a range of droplet-based scRNA-seq based methods, including inDrop, Drop-seq, and 10X. All of these methods utilise the same principles, whereby microfluidics is used to encapsulate single cells and barcodes into subnanolitre droplets, where cell lysis and barcoding occur. The main features of these various platforms are described in Table 1. The different technologies vary in terms of cell capture, efficiency, doublet rate, and cost and the ideal platform is experiment dependent (reviewed in [21]). Broadly speaking, inDrop and 10X have a higher capture rate, which is ideal for experiments with less available tissue. Furthermore, 10X has the highest sensitivity, making it the better choice.

Table 1. An Overview of Available Microfluidics-Based Single-Cell Transcriptomics Devices

<table>
<thead>
<tr>
<th>Supplier</th>
<th>Drop-seq</th>
<th>inDrop</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bead type</td>
<td>Hydrogel</td>
<td>Solid</td>
</tr>
<tr>
<td>Main features</td>
<td>Gel bead in emulsion (GEM) droplets; reverse transcription within droplets</td>
<td>Commercial pairing with Illumina reagents for an end-to-end solution</td>
</tr>
<tr>
<td>System name</td>
<td>Chromium</td>
<td>ddSeq</td>
</tr>
<tr>
<td>Cells captured per sample</td>
<td>80,000+</td>
<td>300–375 per well</td>
</tr>
<tr>
<td>Samples per run</td>
<td>Up to 8</td>
<td>Up to 4</td>
</tr>
<tr>
<td>Run time</td>
<td>&lt;9 min</td>
<td>&lt;5 min</td>
</tr>
<tr>
<td>Capture efficiency</td>
<td>Up to 65%</td>
<td>3%</td>
</tr>
<tr>
<td>Doublet rate</td>
<td>~0.9% for 1000 captured cells, ~8% for 10,000 captured cells</td>
<td>~6% for 1384 cells</td>
</tr>
<tr>
<td>Cost per cell</td>
<td>&lt;$0.35 (incl. sequencing)</td>
<td>&lt;$0.35 (incl. sequencing)</td>
</tr>
<tr>
<td>Analysis pipeline</td>
<td>Cell Ranger</td>
<td>BaseSpace app</td>
</tr>
<tr>
<td>Ease of use</td>
<td>Plug and play</td>
<td>Plug and play</td>
</tr>
<tr>
<td>Refs</td>
<td>[21,70]</td>
<td>[71,72]</td>
</tr>
</tbody>
</table>

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for the detection of lowly expressed transcripts. Drop-seq is the most cost-efficient protocol but has a lower capture rate and sensitivity. However, all of the methods produce high-quality data for single-cell profiling [21], and key considerations for the usage of scRNA-seq are summarised in Box 1.

In addition to droplet-based approaches, there are also well-based approaches to scRNA-seq such as SMART-seq2 [27]. The SMART-seq2 protocol requires cells to be sorted (often using FACS) into individual wells on a plate, whereby each cell is processed ready for sequencing separately. The advantages of SMART-seq2 are a significantly higher capture rate and the sequencing of full-length transcripts and lower technical noise at the cost of capturing fewer cells (a limit imposed by the number of plates that the experimenter is able to process). SMART-seq2 also has the highest sensitivity of all scRNA-seq methods (exceeding 10X), making it the best choice for the detection of lowly expressed transcripts. Furthermore, rare tissues can be investigated by combining SMART-seq2 with FACS, as reported by Efroni et al. (2016) [28] for their study of root meristem regeneration. As with droplet-based approaches, well-based scRNA-seq has not been used widely in the plant community, but it is particularly suited for experiments on small organs or rare tissues where the tissue yield is low.

Applying scRNA-seq to Plants
For plants, single-cell experiments require removal of the cell wall by enzymatic digestion (protoplasting) before the collection of a representative (unbiased) pool of cells. Differences in cell wall composition (due to degree of lignification or suberisation) and the position of cell layers within a tissue can result in an incomplete dissociation of cell files, resulting in biased cell harvesting unless suitably accounted for in the digestion process. In addition, resuspension of protoplasts for input to the single-cell systems must consider the properties of the resuspension buffer: (i) viscosity and likelihood of crystallisation; (ii) compatibility (e.g., Ca²⁺-containing cell suspension buffers cause precipitation of Drop-seq lysis buffer); and (iii) osmolarity to maintain the viability of the cells without

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**Box 1. Balancing Costs and Capacity: Considerations for scRNA-seq**

**How Many Cells Are Enough?**
Typical numbers of single-cell transcriptomes identified for a single sample range from about 1000 to about 8000 cells. The input tissue required to achieve this will depend on the chosen platform. One should consider the expected frequency of the rarest cell type of interest to the study when choosing the number of cells, but also consider that many genes are not detected in single cells and this depth can be improved by capturing a larger number of cells for each cell type. If it is anticipated that the population of cells to be captured will be much smaller than 1000, one should consider using a well-based approach such as SMART-seq2 [27].

**How Does the Number of Cells Recovered Affect Downstream Analyses?**
The number of cells impacts the resolution of downstream analyses; that is, insufficient cells would mean that the impact of drop-out would be higher in all populations and a marker gene/differentially expressed (DE) genes may be hard to detect. Rare populations will also be missed. Careful choice of single-cell platform to match the experimental requirements can mitigate this problem.

**How Does the Sequencing Depth Affect the Sensitivity of Downstream Analyses?**
The sequencing depth required for single-cell analysis depends on the platform and experimental design. 10X recommends sequencing to a minimum 20,000 read pairs per cell. The sequencing saturation can be used to determine whether a dataset has been sequenced enough, which in turn will depend on the experiment. If the aim is to detect lowly expressed transcripts, sequencing saturation of >90% is required. If the aim is to delineate cell types, a lower sequencing saturation is acceptable.

**What Is the Trade-Off between Cell Number and Sequencing Depth Per Cell?**
The experimental design and question should drive the decision of whether to prioritise cell numbers or sequencing depth, based on the considerations above.
inducing transcriptomic changes – for instance, lactose has osmotic properties similar to sucrose, while the latter is sensed by plant cells and regulates the transcription of energy-related signalling processes [29]. Once the cells of interest have been captured in suspension, these are loaded on the droplet-based scRNA-seq device for tagging using barcoded beads (Figure 1B).

The Drop-seq protocol encapsulates individual cells in droplets and utilises a unique primer-barcode system attached to microparticle beads to identify both the cell of origin and a unique molecular identifier (UMI) for each transcript [26] (Figure 1B). The use of UMIs ensures individual tagging of each mRNA molecule and allows the identification of PCR duplicates. Bead-primer-mRNA complexes are referred to as single-cell transcriptomes attached to microparticles (STAMPS) [26]. In the microfluidic device, aqueous flows contain cells in suspension. The barcoded beads are contained in a separate flow of a lysis buffer. The two flows come together and pass through an oil channel, which leads to the formation of a droplet (Figure 1C). Once encapsulated in a droplet, cells are immediately lysed, releasing RNA that binds to primers on the bead surface where it is reverse transcribed into cDNA to generate comprehensive libraries comprising the barcoded cDNA of thousands of single cells (Figures 1D–F). Following sequencing, these libraries are quantified as a digital gene expression (DGE) matrix of read counts per gene and per cell barcode (Figures 1G and 2A).

In general, the number of droplets is much higher than the number of cells in the sample, such that the majority of barcodes will come from droplets without a cell (referred to as ‘empty barcodes’) or one cell. However, incomplete tissue protoplasting can produce droplets with two or more cells (‘doublets’). Identifying empty barcodes and doublets are key challenges in the data analysis process. The critical first step is to define the (approximate) cut-off point between empty and nonempty barcodes. To make this estimate, there are two recommended diagnostic plots: the cumulative distribution plot and the barcode rank plot. Both plots aim to identify the cut-off point based on the distribution of read counts. First, the cumulative distribution plot of the cumulative number of reads per cell barcode in descending order should reveal a ‘knee’ in the distribution indicative of a transition from STAMPS to background noise. For barcode rank plots, cell barcodes are also ranked by descending read count and the barcode ranks are plotted against the read counts using a log scale on both axes. This renders an inverse S-curve, where the drop indicates the separation of STAMPS and background. While these plots can support the decision about which cells to include in the downstream analysis, their quality and interpretability are dependent on the device used, the sequencing depths, and the tissue type. Therefore, an unclear barcode rank or cumulative distribution plot does not necessarily indicate a failed run. In boundary cases it is recommended to look at further characteristics of the data (e.g., gene counts, UMI counts) to get a deeper insight into the data quality. Representative examples for a barcode rank plot for 10X data can be found in [30] and for a cumulative distribution plot in [31].

The next step is to remove further sources of technical and biological variation. These include empty barcodes (not captured by initial filtering), doublets, and broken cells, henceforth referred to as low-quality cells. The distributions of the numbers of genes and UMIs and the percentage of mitochondrial and plastid RNA across the cells can be visualised as distribution plots to determine appropriate thresholds for filtering (Figure 2B). The outliers in these plots can represent either broken cells or doublets. Barcodes with both low UMI and gene count (compared with the overall distribution of barcode counts in the dataset, which is dependent on platform and sequencing depth) that also have a high percentage (>10%) mitochondrial reads indicate loss of cytoplasmic RNA from perforated cells [32,33]. The cells displaying these characteristics are likely to be broken and should be removed from downstream analyses.

While empty barcodes and broken cells will have less genetic material, doublets will have more genetic material than real cells (high numbers of detected genes and UMIs). Commonly, a high count threshold is used to filter outliers based on the distribution of the data. Additionally, there are a number of recently published doublet detection tools that offer more complex approaches for the identification of doublets (e.g., Scrublet [34], Doublet Finder [35]).
Current best practice is to initially apply permissive quality control thresholds (Figure 2B), which can be revisited if contamination becomes apparent in downstream analyses; for example, if one of these quality control metrics is driving differences in clustering [33].

**Cell-Type Identification Using Dimensionality Reduction and Clustering**

Having extracted robust and high-quality cells, the biological features of the dataset can be analysed. Currently, all available scRNA-seq datasets produced from plants were performed on roots using the
10X Genomics or Drop-seq platforms [8–12]. The complex structure comprising many different cell types make roots the ideal plant tissue for single-cell transcriptome studies. All of the current root scRNA-seq studies used similar analytical pipelines, such as Seurat and Monocle for data processing and stepwise analysis [36,37]. Briefly, the top most highly variable genes (HVGs) are used to perform principal component analysis (PCA) and t-distributed Stochastic Neighbourhood Embedding (tSNE) or Uniform Manifold Approximation and Projection (UMAP) to visualise the data structure (Figure 2C,D). Visualisation techniques such as tSNE and UMAP have been developed to dissect and visualise complex high-dimensional data such as scRNA-seq read counts [38,39] (Figure 2E,F).

Analysis pipelines for scRNA-seq utilise only the most variable genes based on average gene expression and dispersion to prioritise cell type-specific expression patterns (Figure 2C). HVGs are characterised as being highly expressed in some cells and lowly expressed in others, making them likely drivers of variance between cell subsets. Typically, the optimum number of HVGs is between 1000 and 5000, depending on the complexity of the dataset. Implementations of HVG ranking programs (as in Seurat and Scanpy) include visualisation tools to help in guiding users towards a correct threshold. If important genes are known, Luecken and Theis (2019) suggest that the threshold could be varied to ensure that all of these are captured within the HVGs [33]. PCA is used to further reduce the complexity of the dataset into fewer PCA dimensions prior to employing tSNE or UMAP (Figure 2F shows an example of a tSNE visualisation) for visualisation and clustering algorithms to identify cell subsets with similar transcriptional profiles. tSNE or UMAP take a high-dimensional representation of a dataset and compute a 2D representation that preserves the underlying structure, and in tSNE or UMAP plots, transcriptomically similar cells are usually close neighbours. Clustering algorithms such as k-means or graph-based approaches are applied to the data to identify biologically significant groups, often corresponding to cell types. Clustering algorithms and their application to single-cell data are reviewed in [40,41]. Showing the clusters on the tSNE or UMAP visualisation by colouring cells can be an indicator of the quality of the clustering; if the clusters are well defined and the colour identity matches the spatial arrangement, the clustering is likely to reflect the underlying biology. As with the previous steps, tSNE and UMAP can be sensitive to their parameters (particularly the perplexity parameter for tSNE), which need to be optimised for each dataset. These dimensionally reduced and clustered data (e.g., using hierarchical clustering or graph-based clustering) [40] can provide a better understanding of the data structure on which biological expression patterns can be investigated (Figure 2G). Moreover, it can help to identify groups of similar cells that can often be identified as cell types. Along the analytic pathway, a number of parameter settings need to be made. Optimal parameter settings depend on individual datasets and changes to one parameter can affect the optimal choice of another (Figure 3). This frequently leads to repeated analyses where the different parameter settings are adjusted until the best choices have been identified.

Currently, two methods are most widely applied to assign cell-type identity to clusters: examining the expression of known cell-type markers (supervised) and unsupervised identification of the genes specifically expressed in each cluster. Denyer et al. (2019) and Shulse et al. (2018) both compared the expression marker genes from Brady et al. (2007) using the Index of Cell Identity (ICI) method to identify cell types [3,8,9]. The ICI method computes an ICI score for each cell representing the relative contribution of each known tissue type to the cell identity. This supervised classification approach is based on comprehensive libraries of possible cell and tissue expression profiles that originate from experiments from known cell types. Instead of relying on variable genes and similar expression patterns, the set of informative markers used by the ICI method is chosen based on an adapted approach from information theory and is not required to be uniquely expressed in a single cell type [19]. In turn, Jean-Baptiste et al. (2019) used Spearman’s rank correlations as a supervised approach to identify different cell types by comparing the gene expression within their clusters to existing markers [2,3,11]. Denyer et al. (2019) further used unsupervised cell clustering to identify potential novel cell-type markers and validated these in silico by comparison with the expression in the existing plant expression atlas and then in vivo with transcriptional reporter fusions [9].
Applying scRNA-seq to Arabidopsis roots has revealed that cell types can be readily identified from the data. Specifically, clusters corresponding to pericycle cells, phloem sieve elements, and different epidermis subpopulations were identified [8–11]. Even very small cell populations such as quiescent centre (QC) cells have been discovered. For instance, Denyer et al. (2019) used the expression patterns of 15 proposed QC genes to identify 36 putative QC cells, while Ryu et al. (2019) used 52 proposed QC markers to identify two likely QC cells [9,10,19,42].

This indicates that plant/root scRNA-seq analyses can be used as a springboard to study gene network organisation in rare cell types, helping to further develop our understanding of fundamental aspects of plant life such as the processes regulating stem cell niche function in cell fate decisions, root patterning, or root longevity [28,43–49]. However, given that these captured populations are so small, great care must be taken to ensure sufficient statistical rigour.

Pseudotime Reveals Developmental Structures

One of the attractions of studying the gene expression patterns of roots is the regular patterning of cell types and the existence of a clear developmental axis from stem cell niche and young undifferentiated cells in the root meristem and newly differentiated cells in the elongation zone to fully differentiated cells in the root maturation zone [1]. The concept of ‘pseudotime’ can be used to investigate the development of undifferentiated meristematic cells into mature tissues. In this differentiation process, the different developmental stages correspond to distinct changes in the gene expression profile of the cells. However, cells may show transitions at different speeds. This asynchrony means that gene expression changes should not be evaluated in dependence of time but instead in dependence of progress through developmental processes. Pseudotime is an abstract measure capturing this progress. Pseudotime methods (e.g., Monocle [37], TSCAN [50]) use machine learning to order cells in ‘time’ along a trajectory, which delivers insights into developmental stages and transition.
Denyer et al. (2019) first used pseudotime to show large-scale differentiation from root meristematic tissue to mature tissue. Interestingly, they defined some clusters as ‘meristematic’ tissue, indicating that despite cell fate being defined after a single division, the transcriptional identity of these early cells seems to be more strongly defined by the developmental stage than by cell type. Furthermore, pseudotime can be used to decipher gene regulatory networks involved in cell differentiation, as demonstrated for atrichoblast, trichoblast (root hair cell), and cortex differentiation [9–11], and revealed the involvement of 3000 genes in trichoblast differentiation [9].

Potential of scRNA-seq in Plant Science

Fundamentally, a tissue’s function is defined by the specific functions of its constituent cell types. In animal systems, scRNA-seq not only has enabled the examination of individual cell types in the context of complex tissues but also has resulted in the identification of new cell types and cell-type states [51–53]. This has given an unprecedented view of the gene expression dynamics in complex tissues and how these change under differing physiological conditions (e.g., healthy vs diseased/cancerous tissue) [54]. Alongside the improvements in single-cell technologies, novel bioinformatics methods have arisen to study more complex gene regulatory principles such as transcriptional noise and RNA velocity [55–57]. This enthusiasm for scRNA-seq studies has paved the way for the development of a human cell atlas that aims to create an ‘ID card for each cell type’ and a ‘3D map of how cell types form tissues’ [15].

As mentioned above, recent studies have demonstrated the feasibility of applying scRNA-seq to plant systems and conducting projects on a similar scale as reported for animals. Defining gene networks from scRNA-seq data by combining cluster analyses with pseudotime methods allowed the identification of cell type-specific gene expression profiles involved in the differentiation of root cell lineages and, thus, the description of developmental trajectories in a whole tissue [8–11]. In addition to resolving developmental timelines, cell types might further affect the status of each other to define the root body plan. scRNA-seq studies with Arabidopsis mutants lacking defined cell lineages, such as endodermis, trichoblast, or atrichoblast, showed altered single-cell transcriptomes as indicated by an overall change in the patterning of cell-type clusters [9,10]. The ability to detect such transcriptional variation further highlights scRNA-seq as a new resource to explain phenotypes in mutants, plant germplasms, or natural plant populations.

In addition to broad studies of large tissues, such as whole roots, high-throughput scRNA-seq could be utilised for the examination of specific tissue parts or developmental zones, such as the root meristem (with stem cell niche) at root tips. Unlike in mature roots, FACS approaches are less suited to root-tip studies due to the limited availability of appropriate fluorescent markers. It is also difficult to capture cell states within a cell-type file and to process cell types that form only a very small proportion in a tissue, such as the QC in the stem cell niche. The difficulty of identifying rare cell populations is compounded in droplet-based scRNA-seq by low gene detection and ‘drop-out’; that is, if a marker is slowly expressed, the chance of it being detected in an individual cell is low. However, current methods do allow the capture and identification of QC cells [9,10], highlighting the sensitivity of scRNA-seq in resolving distinct cell states whose differences might be mild across a cell-type file but that are essential to steer the incremental transition of stem cells (e.g., epidermis initials) to meristem and mature cells (e.g., trichoblasts).

In this respect, it is important to note that integrating data across experiments is a challenge as biases are introduced between laboratories, due to varied cell dissociation protocols, library preparation, and sequencing platforms [58]. Various computational methods, such as mutual nearest neighbours (MNN), k-nearest batch-effect test (kBET), and canonical correlation analysis (CCA), are being developed and applied to enable the integration of multiple datasets [59–61]. Furthermore, Seurat v3 has the capacity to integrate multimodal data through the identification of ‘anchor’ cells across datasets or through the use of a reference dataset [36].
scRNA-seq might further help in our understanding of the functional significance of ploidy. As root cells develop and mature, these undergo a process almost unique to plants: root cells change their ploidy [62,63]. This process is driven by endoreduplication (cell cycle without mitosis) and cell ploidy increases with cell development [63,64]. As sessile organisms, plants are exposed to extreme and fluctuating environmental conditions and ploidy is thought to support stress resilience by increasing cell and genome stability [63,65]. In line with this, cells are known to change their ability to respond to stress with cell age. Interestingly, cells lose their ability to regenerate a root stem cell niche once these start to differentiate [28]. Cell type-specific microarray and RNA-seq studies further show that cell identity plays a strong role in response to environmental stress [4,22–25]. To investigate the extent to which stress responses vary between cell types of different age, single-cell studies will be particularly useful. However, Jean-Baptiste et al. showed that when a strong treatment such as heat shock is applied, it is challenging to assign clusters to specific cell identities due to the overwhelming transcriptional impact of the stress and the downregulation of canonical marker genes. This masking of more lowly expressed cell-type markers by more strongly expressed stress gene networks makes it difficult to assign clusters to specific cell identities and to capture the transcriptional impact of the stress and the downregulation of canonical marker genes. This method of single-cell RNA-seq data with imaging techniques is enabling the spatial and temporal reconstruction of cell types and cell states in the tissue. The prospect of the technology, especially in combination with cell epigenetics (e.g., based on scATAC-seq), multimomics approaches, high-resolution imaging, and spatiotemporally resolved cell engineering, is to describe the synergy of ‘microprocesses’ in the development and behaviour of complex tissues under changing environments (see Outstanding Questions). Bundling of those efforts will be essential in developing unprecedented infrastructures such as the creation of a plant cell atlas [66] to underpin new insights into how the interactions between cells influence function, how cells act in a complex system to create an organ, and how cells react as a network to pathogens or environmental changes [67,68]. In addition to advancing fundamental research, a comprehensive plant cell atlas will form an essential scaffold on which to base studies aimed at sustainable improvement of the productivity and value of agricultural ecosystems [69].

Concluding Remarks and Future Perspectives

By enhancing spatiotemporal resolution, scRNA-seq enables us now to virtually dissect and scrutinise whole organisms. To obtain more detailed insights into the organisation of gene regulatory networks in single cells, however, requires further improvements, for instance, in single-cell tagging, labelling, and capturing of mRNA as well as in the development of bioinformatic analyses, to reduce interfering technical noise. Further methodological developments are increasing the range and resolution of single-cell measurements. Application of an assay for transposase-accessible chromatin using sequencing (ATAC-seq) on the single-cell level provides access to the DNA regions harbouring the regulatory codes underlying transcriptomic patterns observed in scRNA-seq data. In addition, the combination of scRNA-seq data with imaging techniques is enabling the spatial and temporal reconstruction of cell types and cell states in a tissue. The prospect of the technology, especially in combination with cell epigenetics (e.g., based on scATAC-seq), multimomics approaches, high-resolution imaging, and spatiotemporally resolved cell engineering, is to describe the synergy of ‘microprocesses’ in the development and behaviour of complex tissues under changing environments (see Outstanding Questions). Bundling of those efforts will be essential in developing unprecedented infrastructures such as the creation of a plant cell atlas [66] to underpin new insights into how the interactions between cells influence function, how cells act in a complex system to create an organ, and how cells react as a network to pathogens or environmental changes [67,68]. In addition to advancing fundamental research, a comprehensive plant cell atlas will form an essential scaffold on which to base studies aimed at sustainable improvement of the productivity and value of agricultural ecosystems [69].

Outstanding Questions

How close are we to virtually reassembling complex plant tissues based on single-cell transcriptomes? Which scRNA-seq parameters represent limiting factors in obtaining deep transcriptome data? Can we apply scRNA-seq technology to any plant tissue and any crop? How do stem cell niche and individual cell types define the plant/root body plan? Which gene networks define and maintain the quiescent centre and stem cells and their interaction? How do stem cell initials define the identity of cell types? What determines the status of individual cell types and how much is it influenced by neighbouring cells? Which signals and gene networks regulate the developmental transition of cell types? What defines individual cell types and to what extent does their status change during development or environmental stress? How do cell types and cell-type states contribute to plant development and environmental stress adaptation? Which regulatory principles beyond transcription and translation (e.g., transcriptional noise) are prevalent in cell types during different developmental stages and under stress?

References