Best practices for the execution, analysis, and data storage of plant single-cell/nucleus transcriptomics

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Short title: Best practices in plant single-cell/nucleus-omics

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Abstract Single-cell and single-nucleus RNA-sequencing (scRNA-seq and snRNA-seq) technologies capture the expression of plant genes at an unprecedented resolution. Therefore, these technologies are gaining traction in plant molecular and developmental biology for elucidating the transcriptional changes across cell types in a specific tissue or organ, upon treatments, in response to biotic and abiotic stresses, or between genotypes. Despite the rapidly accelerating use of these technologies, collective and standardized experimental and analytical procedures to support the acquisition of high-quality datasets are still missing. In this commentary, we discuss common challenges associated with the use of single-cell transcriptomics in plants and propose general guidelines to improve reproducibility, quality, comparability, and interpretation, and to make the data readily available to the community in this fast-developing field of research.
Introduction: Plant-specific challenges for single-cell approaches

Plant molecular and developmental biologists are fully embracing single-cell applications. Specifically, single-cell RNA-sequencing (scRNA-seq) and single nuclei RNA-sequencing (snRNA-seq) are gaining a lot of traction while spatial transcriptomics is emerging as a promising complementary technology (Figure 1). Despite an increase in the use and publication of plant single-cell experimentation (Figure 1A), it is fair to say that the plant field has, so far, not settled on common strategies, protocols, or analysis methods. Given the high complexity of the different technologies and sample types (Figure 1B-C), we feel it is important to provide a best-practice workflow and guidelines that will help in establishing a collectively accepted quality cut-off. These guidelines will aid in the evaluation of experimental approaches and computational analyses of single-cell transcriptome data, while also offering solutions to commonly observed challenges, thereby improving the reproducibility and comparability of experiments in the broader field of plant research. The present co-authors collectively accept these guidelines and commit to applying them to their research. We also highlight examples where consensus has not yet been achieved between co-authors, which will need to be resolved when both the technologies and the field develop further. As one example, single-cell multi-omics and spatial transcriptomics are, in our opinion, not established enough in the plant field to propose any sort of definitive rules at this moment in time.

To date, we have identified the following eight main challenges in the field of plant single-cell/nucleus (sc/sn) transcriptomics: 1) deciding on the best single-cell methods to answer a specific biological question; 2) understanding experimental variability; 3) biases in protocols and platforms; 4) deciding on a sequencing strategy; 5) generating expression matrices and defining high-quality cells; 6) constructing cell clusters and mapping them to cell types; 7) trajectory inference methods and applications; and (8) documenting and publishing datasets (Figure 2). Each of these challenges is discussed in detail in the following sections.

Challenge 1: Selecting the best approach to answer a specific biological question

Before considering the best experimental approach to obtain single-cell transcriptome data, it is important to evaluate the potential benefits of accessing single-cell resolution over bulk RNA-sequencing (bulk RNA-seq). This assessment depends on the biological system considered and the biological questions to answer. Sc/snRNA-seq provides a snapshot of the transcriptome of each cell within an organism, offering a high spatiotemporal resolution of the dynamic gene regulation involved in plant development, cell differentiation, or responses to biotic and abiotic stresses. Single-cell transcriptomics can also offer the required resolution to study cell type
specific responses during cellular evolution and adaptation mechanisms among plant species (Guillotin et al., 2023). We thus advocate using sc/sn transcriptomic technologies over bulk RNA-seq when working with a cellularly complex sample or to capture dynamic transcriptomic responses to stimuli. In other cases, the question at hand might be more easily addressed using bulk RNA-seq or targeted gene expression experiments.

**scRNA-seq versus snRNA-seq**

A thorough understanding of the different strategies and types of protocols (Figure 1C) is essential before one can make an educated decision on which technology will best answer a specific biological question. A first example of an important choice is whether to profile the transcriptome from isolated nuclei or cells. When doing scRNA-seq, the most popular choice to obtain single-cells from a plant organ requires the enzymatic digestion of cell walls and the generation of so-called protoplasts. There are a number of disadvantages of using protoplasts such as: some tissues (e.g. sclerenchyma) and species (e.g. Sorghum; (Guillotin et al., 2023)) are recalcitrant to cell wall digestion; enzymatic digestion affects the transcriptional status of the plant cells and could bias the outcome of experiments (Birnbaum et al., 2005); and the large size of protoplasts reduces their capture efficiency with most of the currently available commercialized single-cell platforms. Nuclei isolation followed by snRNA-seq gained traction in plant single-cell transcriptomics as well. However, the recovered data content per nucleus (e.g. UMI or genes) is up to ten (for UMIs) and three times (for genes) lower compared to scRNA-seq (Farmer et al., 2021; Guillotin et al., 2023). Furthermore, even though the transcriptome coverage is similarly efficient between scRNA-seq and snRNA-seq (e.g. 89% of all Arabidopsis expressed genes were captured in snRNA-seq data (Farmer et al., 2021; Guillotin et al., 2023)), a nuclear and a cellular transcriptome are not equivalent (Lee and Bailey-Serres, 2019; Reynoso et al., 2019). For example, differences in abundance and composition between transcripts obtained from nuclear versus polyA RNA under hypoxia, point towards nuclear transcript retention or enrichment as part of the stress response (Lee and Bailey-Serres, 2019; Reynoso et al., 2019). Furthermore, the half-life of the transcripts (estimations range between 12 min to more than 24 hours in Arabidopsis cells (Narsai et al., 2007)) suggests that the cellular transcriptome is the result of the accumulation of the transcript synthesis over time, while the nuclear transcriptome is considered to accommodate faster to changes in gene activity. These differences are important to consider when selecting and later interpreting a single-cellular versus a nuclear-based transcriptome and should be determined by the experimental system or biological question. Therefore, when studying e.g. early stress responses of plant cells, a snRNA-seq could achieve higher resolution of rapid transcriptome changes, while scRNA-seq might be more informative when understanding the biology of a cell type or when studying cells that are enucleated at some
stages of development (e.g. sieve element cells in the phloem cell lineage (Miyashima et al., 2019)).

**Biological replicates in single-cell transcriptomics**

As for all scientific observations, generating robust sc/sn data sets requires performance evaluation across multiple, independent biological replicates. We hereby note that a biological replicate relies on the independent growth, harvesting and processing of various plant samples. Any separation after protoplast or nuclei isolation cannot be classified as biological replicates and can only be reported as technical replicates. No standardized metrics are available within the community to evaluate reproducibility between replicates. We propose that a correlation coefficient of the average gene expressions among all cells would be an informative assay. Alternatively, one could compare the frequency of cell types or cell clusters across replicates. As such, we advise analyzing cell cluster-specific differentially expressed genes and annotating each replicate separately, before merging the replicates and applying batch effect correction. Other parameters, e.g. Average Silhouette Width (ASW) and Adjusted Rand Index (ARI), have been used to quantify cell type purity assessments after batch effect correction and can also be informative to evaluate replicate robustness (Tran et al., 2020).

From a statistical point of view, independent biological replicates are unconditionally advised to increase the significance of biological datasets (Heumos et al., 2023). However, in many cases, replicates in sc/snRNA-seq experiments are currently performed to increase the total number of cells or nuclei analyzed, while the replicate information and a comparison between replicates is not necessarily incorporated in the actual statistical analysis. A statistical comparison among biological replicates is thus strongly advised to ensure high data quality and to prevent cluster formation based on replicate-specific artefacts. As such, biological replicates are imperative to add certainty on the reproducibility of the experiment. However, merely adding biological replicates does not remove transcriptional artefacts introduced during sample preparation in each of the replicates. One example is the effect of the enzymatic digestion needed to generate protoplasts or the procedures to extract nuclei on the transcriptome. Therefore, performing replicates by themselves does not provide sufficient confidence in the data to draw biological conclusions. To achieve confident biological interpretation, extensive downstream experimental validation is always required in the form of e.g. reporter lines, *in situ* hybridization or spatial transcriptomics.
Challenge 2: Experimental variability during sample preparation

While the potential of sc/snRNA-seq for plant research is evident, its applicability depends largely on establishing reliable cell and nucleus isolation protocols. These protocols must support the generation of high-quality, high-yield nuclear and viable cellular suspensions within a short amount of time, and must be compatible with downstream procedures (e.g., limited usage of PCR inhibitors like CaCl₂). The efficiency of protoplast generation from tomato roots for example was increased by optimizing the pH of the enzyme-containing buffer, and in part also by using hand sections instead of intact tissues (Omary et al., 2022). Preincubation in L-cysteine and sorbitol for roots of Maize, Sorghum and Setaria improves enzymatic cell wall digestion and protoplast generation (Ortiz-Ramírez et al., 2018, 2021), while L-Arginine positively influenced survival rate of Maize meristem protoplasts (Satterlee et al., 2020). In contrast, nuclei can be isolated from fresh (Farmer et al., 2021; Picard et al., 2021; Cervantes-Pérez et al., 2022; Conde et al., 2022; Liu et al., 2022b; Sun et al., 2022), frozen (Sunaga-Franze et al., 2021; Neumann et al., 2022; Abramson et al., 2022; Li et al., 2023) or fixed (Kao et al., 2021) starting material, offering flexibility in terms of sample handling and preparation, while simultaneously securing dynamic transcriptional changes upon their rapid fixation. While nuclei isolation seems more straightforward to conduct than protoplast isolation, the assessment of nuclei quality prior to snRNA-seq library construction remains a difficult task. The leaking and clumping of isolated nuclei should be seen as a sign of breakage of the nuclear membrane leading to RNA leakage and the generation of low-quality libraries.

Overall, careful workflow optimization should include the following:
1) visual assessment of tissue digestion or nuclei release through e.g. the observation of protoplasts/nuclei produced from all desired cell types, via cell wall digestion of fluorescently-tagged cells of a particular cell type (if available) as a proxy, or via gene expression quantification of cell type markers in a pilot experiment.
2) rapid and non-destructive sample clean-up strategies including washing steps (e.g., centrifugation, filtration), fluorescence (and image-based) activated cell/nucleus sorting (FACS/FANS), or microfluidic cell-enrichment devices can increase the population of viable cells and the purity of cellular/nuclear suspensions. A careful analysis of nuclei shape will help to identify problems with RNA leakage.
3) careful analysis of cell sizes for protoplasts since most commercial platforms have a cell size restriction which might introduce a bias in cell capturing and a preference for incorporating smaller over larger cells (see Challenge 3).
4) quantification of cell viability by manual cell counting (upon staining with trypan blue or fluorescein diacetate) or with the help of automated cell counters.
The procedure of cell wall digestion itself (Birnbaum et al., 2005) - as well as external factors introduced during sample collection and generation (e.g. growth and harvesting conditions, enzyme concentration and activity, temperature and timing) - affect cell viability, cell wall digestion efficiency, cell type representation, and the transcriptional profiles of cells. One of the most promising developments for reducing experimental biases is the inclusion of a fixation step. Until now, scRNA-seq compatible cell fixation protocols have mainly been described in mammalian research (Attar et al., 2018; Phan et al., 2021; Wohnhaas et al., 2019; Wang et al., 2021a), but its application could drastically boost the plant single-cell field by massively reducing the effects of external factors during sample processing, including the generation of protoplasts. Indeed, protoplast isolation efficiency increased when plant tissues were fixed and digested at optimal enzyme activity temperature (Marchant et al., 2022). However, concerns about tissue fixation on protoplast shape (Marchant et al., 2022) and the sequencing results have been reported, motivated in particular by reduction of cDNA yields and biases towards 3’ end-enrichment (Wang et al., 2021a). Despite these limitations, the potential gains for the field could be major, warranting dedicated investments in tissue fixation approaches.

Challenge 3: Biases and specificities of commercial platforms for plant single-cell transcriptomic samples

The most popular commercial platforms and scRNA-seq protocols used for plant samples rely on microfluidic droplet-based cell compartmentalization or nanowell-based cell separation (Figure 1C). Techniques that can be performed by manual handling, such as combinatorial barcoding (Cao et al., 2017; Rosenberg et al., 2018), are rapidly expanding in the animal field, but have yet to be shown in use for plant samples. The choice of the sample processing method or platform must be taken carefully to allow uniform cell size capture rate, resolvability and, if necessary, a sample multiplexing option or flexibility towards cell capturing and lysis steps. Droplet-based platforms allow fast cell/nucleus processing but offer limited flexibility regarding the cell preparation workflow. Also, the level of pressure imposed on the sample when creating the emulsion could cause the bursting of cells into the droplet-based platform. Well-based methods like SMART-Seq2 (Lopez-Anido et al., 2021) and platforms used with plant samples, such as BD Rhapsody (Zong et al., 2022) or iCELL8 (Sunaga-Franze et al., 2021), require longer cell processing protocols, but offer more flexibility during the sample processing. However, the compatibility of commercial platforms to handle the size and fragility of plant protoplasts is not necessarily evaluated. Plant cell sizes typically lie in the range of 10-80 µm, with even larger values observed for endoreduplicated cells, which is far above the recommended cell size maxima from current technology providers (~40 µm). Furthermore, cell size heterogeneity can create cell capture biases, because droplet-based techniques favor smaller cells, while the well
sizes in nanowell techniques must be fine-tuned to reduce the possibilities of doublets from smaller cells while still allowing capture and processing of larger cells. The consequences could be high multiplet rates and/or imbalanced cell type/stage representation. Careful optimization of the maximum cell loading capacity, loading speed, cell compartmentalization time and the number of washing steps is necessary depending on the platform of choice. Identification of cell type or stage capture rates, however, requires in vivo experimental validation by quantification of cell types or tracing of developmental cell stages. This validation has been done by comparing cell numbers per cell type between scRNA-seq data and cell counting via imaging (Wendrich et al., 2020), but it could also be achieved by spiking-in a fixed ratio of cell types using transgenic marker lines.

Furthermore, a detailed plant-specific benchmark study comparing the commercially available platforms and kits is urgently needed to evaluate benefits and pitfalls when applied to plant samples. Similar benchmark studies using human and mouse cell lines allowed practicality and financial comparisons of common methods, while also comparing cell capture rates and technical bias across cells with distinct cell properties (Mereu et al., 2020). Ideally, a benchmark study should cover numerous species and tissue combinations to allow the establishment of quality standards independent of the species or tissue used. A plant-specific benchmark study might in addition also focus on comparing the ability to recreate developmental or spatial cell states, because most plant single-cell samples offer a chance to also capture and compare developmental cell trajectories. Given the sensitive nature of protoplasts, it would also be important to address the effect of total sample processing durations, as well as the abundance of ambient RNA due to for example protoplast bursting.

**Challenge 4: Establishing an efficient sequencing strategy**

**Full-length versus 3’- or 5’-end transcript sequencing**

Two approaches for library preparation are currently used in single-cell methodology, namely full-length transcript coverage and 3’ or 5’-end transcript coverage. While most reported plant single-cell transcriptome datasets today (Figure 1C) were generated using 3’end transcript coverage, full-length transcript sequencing (used for example in Lopez-Anido et al., 2021) bares huge application potential in plant research, as it can help in improving transcriptome and epitranscriptome resources (reviewed in (Shen et al., 2023)) and in providing cell type-selective isoform usage (shown for example during neuronal maturation in mouse embryos (Lebrigand et al., 2020)).
Cell number and sequencing depth

Although cDNA amount and profile after library preparation is used as a proxy for the overall quality of the sc/snRNA-seq library, sequencing followed by analysis currently remains the only way to fully estimate its quality and biological value. Two major issues that require careful consideration during the experimental planning and sample optimization are the number of cells/nuclei needed for optimal coverage of the cell type(s) of interest and the aimed sequencing depth per cell/nucleus. The number of cells/nucleus in published cell atlases is moving from thousands to hundreds of thousands. Increasing numbers is indeed beneficial for predicting novel marker genes, because it allows better coverage of rare populations and lowly expressed genes. This higher coverage in turn helps to outweigh the bias of differential expression analysis tools towards highly expressed genes (Squair et al., 2021) and dataset-specific noise (Fischer and Gillis, 2021). However, at what point does a cell atlas fully capture the cellular diversity of its samples? The meristematic region of an Arabidopsis root has about 3,000-4,000 cells, meaning that a dataset of 100,000 cells sufficiently covers each cell about 20-30 times, assuming that all cells are equally represented in the dataset. Alternatively, when a specific cell type is isolated from a tissue by upstream cell sorting, an atlas of 2,000 cells could already be saturating. As a rule of thumb, when the cell type composition of the tissue of interest is known, the minimum number of cells that need to be analyzed can be estimated from the probability to robustly capture the rarest cell type(s) (e.g. https://satijalab.org/howmanycells/). For example, to obtain at least 10 quiescent center cells (estimated to represent 0.1% of all cells within the Arabidopsis root meristem according to (Cartwright et al., 2009; Shahan et al., 2022)) with 95% confidence, one would need to profile 15K-20K cells.

Additionally, the required sequencing depth must be adapted depending on the biological question, the tissue complexity, and the sample quality. However, it is recommended that optimal coverage is given with one read per cell per gene (Zhang et al., 2020). Alternatively, sequencing can be staged by first initiating a shallow sequencing of the library (e.g. up to 10,000 reads/cell) before deeper sequencing (e.g. aiming for 50,000 reads/cell). Such shallow sequencing allows evaluating the performance of cell cluster analysis and annotation and is sufficient to capture the entire cell type heterogeneity of the sample (Zhang et al., 2020). Another common suggestion for a preliminary sample quality control is to analyze the expression of a gene subset related to a biological question (Zhang et al., 2020). Sequencing even less for testing the quality of the library is possible but will affect the retrieval of cell types with a lower number of transcripts, which might be lost within the background of empty droplets if the sequencing is too shallow. The desired final read depth will depend on the goal of the experiment but should
ensure sequencing enough cells at a sequencing depth that captures individual events robustly (e.g. 50% sequencing saturation (Table 1)).

When assessing the most cost-effective sc/snRNA-seq technologies, should one profile many cells/nuclei but have shallow sequencing or should one profile fewer cells/nuclei but with deeper sequencing? In many cases, the ideal scenario will be something in-between assuming that the researchers are working on high quality cells/nuclei. Nevertheless, a choice towards either a higher number of cells/nuclei or higher sequencing depth can be made depending on the biological question, the quality of the biological entities used for the analysis, and the relative abundance of each cell type composing the organ. If the aim is to generate an atlas potentially uncovering rare cell types, a better strategy would be to profile many cells/nuclei with a lower sequencing saturation. However, a minimal depth in sequencing (Table 1) upon maximizing cell/nucleus quality is still necessary to ensure that low-abundance transcripts that define rare cell types are captured and to saturate the transcriptome of the sample. Validation of high-throughput technologies in the plant field that enable to access the transcriptomes of hundreds of thousands and even up to one million cells or nuclei, combined with the on-going expansion of sequencing capabilities and the decrease in sequencing costs, could help to overcome this dilemma. If the goal is to do functional gene discovery and generate gene regulatory networks (GRN) for example (Ferrari et al., 2022), a high sequencing saturation per cell/nucleus favors discovery of low abundance gene transcripts. To achieve this goal, one can select a subset of genes related to the biological question and adjust the sequencing depth until at least one read per cell for each of those genes is reached (Zhang et al., 2020). In all cases, we recommend optimizing cell and nucleus isolation methods to ensure the capture of the largest number of transcripts from each biological entity.

**Challenge 5: Generating an expression matrix from high-quality cells/nuclei**

Once mRNA sequence reads are obtained, reads are mapped to the genome and ultimately to genes and cells of origin using a reference genome and UMI and cellular barcode information. Standard data analysis workflows further include quality control filtering, quantification of gene expression in each cell, clustering and visualization of cells based on transcriptome similarity (Figure 3). While the recommendations below are given with the intention to standardize sample quality parameters within the field (see recommendations in Table 1), we want to highlight that a sn/scRNA-seq experiment (with a read depth of 30K reads/nucleus or cell) should typically result in the capture of 1000 expressed genes/nucleus or 3000 expressed genes/cell, respectively. Furthermore, utilizing universal preprocessing pipelines for single-cell genomics data (e.g.,
Booeshaghi et al., 2023) can help in streamlining cell quality filtering and enhance data reproducibility in the future.

**Read mapping**

Plant genomes in general and crop genomes in particular are poorly annotated compared to e.g. human or mouse genomes. Moreover, due to frequent whole genome duplications (Fox et al., 2020), many plants are polyploid and thus contain multiple similar copies of each gene. Structural annotations of genes are especially important for most droplet-based technologies using a 3’ capturing strategy to properly map the sequencing reads and to quantify transcript abundance. The distribution of the mapped reads on the genome can give an indication about the quality of the annotations. Poor mapping efficiency consequently causes gene loss, which can be dramatic especially for popular 3’-based single-cell technologies. Even for Arabidopsis scRNA-seq datasets, mapping rates vary, but should be e.g. > 85% (Table 1). A high percentage of reads mapping to intergenic regions (e.g. >20%) can be an indication that either not all genes are annotated or that the annotated 3’ UTR regions should be longer. As a note, it is important to mention that the sequencing reads generated upon conducting a snRNA-seq experiment should be mapped against the exonic and intronic sequences of the annotated transcripts, a reflection of the capture of spliced and unspliced transcripts.

**Removal of low-quality cells/nuclei**

After read mapping, low-quality cells/nuclei (e.g. cells/nuclei with low number of UMIs and genes) need to be filtered out (see numerical recommendations in Table 1). Impairments in applying low-quality cell filtering may increase the noise in the dataset and reduce the accuracy in downstream analysis, including cell clustering or erroneous identification of cell types (Figure 4A-B). General filter parameters will depend on the application and sample type (i.e., cells vs. nuclei). For plants it is good practice to exclude cells with high mitochondrial (e.g. >10%) and chloroplast (e.g. > 5-10%) reads (Table1). Such cases may indicate cells under stress because of perturbations during the sample preparation. These values might need to be adapted when studying highly or lowly metabolically active cells, or cells undergoing e.g. programmed cell death. If no mitochondria or chloroplast genome is available, plotting the number of genes vs. the total UMI count can be used instead to show cells with low data content. The barcode-rank-plot (“knee plot”) is a commonly used tool to determine sample quality by ranking all barcodes according to their UMI content (Figure 4A). A sample of low quality can be identified if there is no clearly defined boundary between barcodes with high UMI content and barcodes with significantly lower UMI content (Figure 4C). Similarly, poor cluster separation in a sample that
fails to differentiate cell types indicates an insufficient amount of transcript content per cell/nuclei.

Ambient RNA and the presence of empty droplets and doublets – both associated with droplet and nanowell-based technologies – can also lead to noise in the expression matrices, inaccurate cell clusters, and falsely differentially expressed genes. Therefore, it is important to ensure low amount of ambient RNA introduced during sample preparation (e.g., by mixing cells/nuclei from multiple species) and to optimize cell loading concentration (Figure 4C). Bioinformatic tools (e.g. SoupX (Young and Behjati, 2020) or CellBender (Fleming et al., 2023)) can be used to computationally remove transcriptional noise introduced by ambient RNA. Experimentally, even though not yet shown suitable for plant single-cell transcriptomics, doublets can be detected using antibody (cell hashing, (Stoeckius et al., 2018)) or lipid-tagged indices (MULTI-Seq; (McGinnis et al., 2019)) sample multiplexing strategies. Here, doublets are identified if the cell-specific barcode is connected to multiple antibody or lipid-tagged indices, respectively. Mixing cells/nuclei from multiple species in one sample offers another experimental set-up to identify doublets ((Shulse et al., 2019), Figure 4C). In this case the doublet is identified due to the cell-specific barcode being linked to multiple species. While an experimental set-up that allows doublet identification would be the best practice for identifying doublets, it will not help in removing doublets from already existing data sets. Even though significant advances have been made in identifying doublets computationally (see benchmark study (Xi and Li, 2021)), it remains a major challenge in general and even more in plant sc/sn transcriptome analysis due to the presence of endoreduplication. Such polyploid cells will appear as outliers when plotting the gene content or UMI/cell, but additional expression quantification of ploidy marker genes (if available) allows to distinguish endoreduplicating cells (Wendrich et al., 2020).

Identification of protoplast-induced genes

In order to exclude that the invasive enzymatic treatments needed to generate protoplasts might influence the observed transcriptional status of certain cells or cell populations, the overall transcriptome responses induced during these procedures should be determined using bulk RNA-seq in each experimental set-up (Birnbaum et al., 2005; Brady et al., 2007). Indeed, protoplast isolation adds a definite stressing factor to each cell type (Denyer et al., 2019; Wang et al., 2021b; Xu et al., 2021). Although the absence or presence of protoplast induced genes did not alter cell clustering or annotation in Arabidopsis root scRNA-seq data (Denyer et al., 2019), these genes should at least be flagged in the dataset to avoid misinterpretation.
Challenge 6: Cell cluster identification and annotation

Data normalization, dimensionality reduction and cell cluster visualization

Considering the limited number of transcripts per cell, single-cell transcriptomics fail to detect transcripts for most genes in a given cell. This sparsity is further enhanced by intrinsic noise from stochastic transcript fluctuations, cell-cycle state, and cell heterogeneity among other biological factors. As a result, it is necessary to implement scRNA-seq-specific normalization and batch correction protocols (see (Luecken and Theis, 2019) for a review on this specifically). Clusters of cells (i.e., cells that share similar expression profiles) are constructed using community detection algorithms, which control the degree to which similar cells should be grouped together or stay separate based on preset parameters. To visualize the data, ‘dimensional reduction’ algorithms are applied to the data, typically via PCA, t-SNE or UMAP. However, a word of caution is necessary here: whereas PCA involves linear projections, t-SNE and UMAP are non-linear transformations introducing significant distortions (Chari and Pachter, 2023). Indeed, visualizing a synthetic dataset with tSNE or UMAP revealed that cluster distances and locations in discrete and trajectory simulations are inaccurately represented compared to the defined distances in the original data (Wu et al., 2018). These results indicate that cluster distances or locations cannot be used alone to draw biological conclusions. If different clustering outcomes (produced with customized parameters) seem equally plausible, it is appropriate to apply the next step - mapping clusters to cell types - to each possibility and then use that extra information to decide which clustering makes most sense from a biological point of view.

Cell cluster annotation

The annotation of clusters (meaning mapping each cluster to a cell type or state) is facilitated using manual or automated cell annotation methods (Figure 5). Manual annotation requires previous transcriptomic knowledge gained from as many different cell types as possible. Cell type marker genes for plant tissues in well-studied species can come from manually curated lists that are based on bulk RNA-seq data from purified cell populations (Jin et al., 2022) or from already annotated single-cell datasets. However, to date, databases containing this information are rare and restricted to few plant species and tissues (e.g. (Jin et al., 2022)). One can appeal to interspecies correspondences to support the functional annotations of the clusters, assuming that a substantial fraction of orthologous genes shares similar cell type transcriptional specificities. For instance, if an atlas for another species is available, one can attempt to extend the orthology between genes to orthology between cell types. Such an approach has been very successful, leading to atlases for instance in rice (Wang et al., 2021b; Li et al., 2023), *Medicago truncatula* (Cervantes-Pérez et al., 2022), Maize, Sorghum and Setaria (Guillotin et al., 2023). However,
this approach also has its limitations: genes often exist in multi-gene families rendering the orthology mapping ambiguous, and conservation of marker gene/cell type pairs is far from perfect (Movahedi et al., 2012). An additional limitation of these approaches is that they require multiple marker genes per cluster to ensure proper annotation.

Another option is so-called label-transfer: transferring cluster labels between existing single-cell expression atlases (e.g. for roots: https://rootcellatlas.org/, or other organs: http://neomorph.salk.edu:9000/) to unlabeled datasets. Common tools used are scmap (Kiselev et al., 2018), SingleCellNet (Tan and Cahan, 2019), SingleR (Aran et al., 2019) and Seurat (Stuart et al., 2019), which differ in their accuracy and their ability to handle sample- and protocol-related nested batch effect removal and the presence or absence of cell types in the reference atlas or target dataset (Luecken et al., 2022). The advantage of this approach is that already published data can be directly re-used. With both automated annotation approaches it is important to consult several tools and select one final annotation, for instance using a majority rule.

As an alternative or complementary approach, clusters can be annotated manually. To achieve this, marker gene expression of cells can be visualized in a UMAP plot or via a dot plot showing cluster specific expression of marker genes. Cell clusters with conflicting annotations or no annotation, due to e.g. low quality of the transcriptomic information or capture of uncharacterized cell (sub)type or cell transition state, should be marked as “unknown”.

Refining cell cluster annotation

Ideally each cluster will be clearly associated with one cell type, using any of the approaches described above. However, the current literature often mixes different levels of anatomical annotations. The ambiguity between classical anatomical descriptions and the new molecular characterizations of these cell types, tissues and structures make it even more challenging to navigate among these definitions. Hence, to resolve these ambiguities, multiple hierarchically structured annotations can be used in which cells/clusters can be annotated according to e.g. broad expression domains, the tissue or the cell type level (Michielsen et al., 2021). Until there is a consensus in the community, we recommend to separate the different level of annotations (cell type, tissue, cell cycle) to distinct plots. This will avoid ambiguity and misinterpretation of the data (Figure 5).

Validation of cell cluster annotation

Independent of the annotation approach used, we strongly recommend experimentally validating the main annotations by e.g. generating corresponding reporters, performing RNA in situ
hybridizations or complementing the data with spatial transcriptomics. This independent experimental approach is the only way to assess if differential gene expression observed in a high-throughput single-cell experiment is relevant in vivo, or an artefact introduced by one of the many steps, e.g. protoplasting.

Beyond these more classical approaches, spatial transcriptomics can be used to support the annotation of clusters identified in plant sc/snRNA-seq datasets (Guillotin et al., 2023; Lee et al., 2023; Nobori et al., 2023). The technologies that allow probing tissue gene expression and simultaneously retaining its spatial location can be divided into two main categories: targeted and untargeted. The division into those two categories is based on the type of approach applied to analyze the tissue gene expression information. Specifically, targeted methods (i.e., in situ sequencing (ISS) (Ke et al., 2013; Laureyns et al., 2021), MERFISH (Moffitt et al., 2016; Lee et al., 2023; Nobori et al., 2023), Xenium (Ke et al., 2013; Lee et al., 2015; Janesick et al., 2022; Liu et al., 2022a), NanoString CosMx (He et al., 2022), Molecular Cartography (Groiss et al., 2021; Guillotin et al., 2023; Yang et al., 2023) to list a few) require a priori knowledge on which genes to study, since these approaches use gene-specific probes to fluorescently visualize and count the gene transcripts of interest in the tissue. In contrast, untargeted methods (e.g. Visium (Giacomello et al., 2017; Liu et al., 2022a, 2022b; Peirats-Llobet et al., 2023), DBiT-Seq (Liu et al., 2020), Slide-seq v2/Curio (Stickels et al., 2021; Lee et al., 2023) and Stereo-seq (Xia et al., 2022)) leverage a localized capture of polyadenylated transcripts, thus allowing to obtain 2D whole transcriptome maps. Both targeted and untargeted approaches can aid in determining the spatial location of specific cell types or stages of interest (e.g. indeterminate and determinate SAM cells in Maize, defined by their expression levels of PLASTOCHRON1 (Laureyns et al., 2021)), annotate cluster identities obtained in sc/snRNA-seq experiments and/or validate marker genes (Guillotin et al., 2023; Lee et al., 2023; Nobori et al., 2023). As such, spatial transcriptomics and sc/snRNA-seq are complementary technologies for obtaining high resolution spatiotemporal expression data.

**Challenge 7: Application of trajectory inference in plant single-cell transcriptomics**

Trajectory inference can predict developmental or stress-response trajectories in sc/snRNA-seq datasets, allowing one to pinpoint e.g. cell cycle transitions or bifurcations when new specific cell identities branch off from another lineage. While trajectory interference is a very promising tool to identify novel biological phenomena and questions, it is important to understand that the pseudotemporal ordering of cells along a trajectory is purely based on transcriptome similarities, meaning that a sufficient, unbiased sampling of cells is required, as well as prior knowledge to
assign a developmental direction (see Tritschler et al., 2019) for general recommendations. Furthermore, no conclusions about the spatial organization can be drawn, and it is advised to confirm and complement the trajectory output with other methods. Multiple trajectory interference methodologies have been developed, benchmarked and used in the animal field (Saelens et al., 2019), but they show special promise in plants because plants have many continuous developmental programs and plant cells have remarkable capabilities for dedifferentiation and adaptation. Trajectory analysis gave significant insight into recreating cell (type)-specific transcriptional events during plant developmental processes, such as lateral root development (Serrano-Ron et al., 2021), stomata development (Kim et al., 2023), root hair (Denyer et al., 2019; Shulse et al., 2019), pistil (Li et al., 2023) or phloem development (Rozsak et al., 2021; Otero et al., 2022). For example, when combined with live-imaging, trajectory-predicted expression gradients and cell type specific transcriptional networks allowed a complete reconstruction of the developmental process and a precise, cell-by-cell lineage tracing during protophloem development (Rozsak et al., 2021). An interesting computational analysis that can be applied to verify predicted trajectories in plant tissues is making use of the ploidy increase in plant cells as they mature from meristematic cells with 2C/4C content into a more differentiated stage, marked by 8C/16C or higher content. In Arabidopsis root tissues, these ploidy states have been linked to specific markers, leading to predictions of the ploidy status of each individual cell. This increase in ploidy level allows to pinpoint or validate the more meristematic cells with lower ploidy levels and the more differentiated cells in a sc/snRNA-seq dataset with higher ploidy levels (Bhosale et al., 2018; Wendrich et al., 2020; Shahan et al., 2022).

Challenge 8: Documentation and publication of plant single-cell data sets

Recent advances in single-cell omics technologies in plants have enabled insights into diverse aspects of physiology and development and have been the centerpiece of a growing number of elegant studies. However, there is additional potential for single-cell resources through their re-use in integrative meta-analyses (showcased e.g. in (Leote et al., 2022)). The multiple applications would increase their power and depth through greater numbers of cells, a more comprehensive assessment of biological variation, and enhanced enrichment for different cell types or states that are targeted by individual studies. Popular methods enabled within software packages (e.g. Seurat, Monocle, Scanpy, Harmony, etc.) (Wolf et al., 2018; Korsunsky et al., 2019; Van den Berge et al., 2020; Hao et al., 2021) have streamlined the process of stitching datasets together across samples, studies, experimental platforms, even across tissues and species, allowing for a more expansive use of single-cell data for describing and understanding biological organization across scales.
To make optimal use of the generated data in the plant field, there is an urgent need for published single-cell data and their associated metadata to be more easily accessible and usable. First steps towards establishing a suitable infrastructure that allows data storage and comparison have been taken within the framework of the Plant Cell Atlas (Fahlgren et al., 2023). To a large extent, this FAIR (Findable, Accessible, Interoperable, Reproducible) principle has not been an issue for raw single-cell data. Unprocessed FASTQ files are deposited routinely through major data portals, such as the National Center for Biotechnology Information Short Read Archive (NCBI SRA; http://www.ncbi.nlm.nih.gov/sra), and accessibility to raw data is usually mandated by journals, funding agencies, and institutions. In most cases, well-indexed raw data is available and cited in journals. However, care must be taken for single-cell data generated through popular platforms in that only reads from the paired-end sequencing strategy, as well as indexed and/or UMI FASTQ must be deposited. These genomic reads alone are insufficient for reconstructing the single-cell counts matrix necessary for nearly all analysis steps. Processed data should also be stored in publicly accessible repositories such as the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo). Processed data includes the cell/gene counts matrix at minimum, but can also include more complex data objects such as those generated by the Seurat (R) (Hao et al., 2021) or AnnData (python) (Wolf et al., 2018) packages. The EBI Single-cell Expression Atlas is another repository that will accept processed count matrices and allows for deposition of cell metadata (see below).

Less prevalent is the public accessibility of metadata associated with single-cell studies. This metadata takes several forms: 1) experimental metadata describing how samples were treated, what tissues were harvested, how the samples were processed, and what platforms/versions single-cell sequencing was performed on them; and 2) imputed metadata describing attributes of cells defined by downstream analysis of the raw data, including the cell’s identifier (usually a sequence barcode), the number of distinct genes detected, total transcripts detected, assigned cell type or cluster number, or other features that are described in a study but not immediately available from the raw data. Experimental metadata, including protocols used to generate samples, should be well-documented in the manuscript that presents them, and we encourage depositing the protocol in a public repository. Protocols.io is emerging as a standard for this (see e.g. the repository for the Plant Cell Atlas; https://www.protocols.io/workspaces/plant-cell-atlas), and detailed sample information should go along with the raw data when submitting to a repository like GEO/SRA. Imputed metadata can be provided along with the raw data in the NCBI GEO as a simple machine-readable table (.csv or .tsv). All forms of metadata are vitally important for integrative analysis, as it is difficult, if not impossible, to recreate exactly the analysis steps performed from a published study to replicate results. In Table 1, we have listed important metadata parameters which should be accompanying all publications using sc/snRNA-
seq data to ensure transparency in data quality and will allow for a better interpretation of the results and their use in larger meta-analyses. Although it is challenging to give an exact number for these parameters due to the vast differences in experimental systems, species, tissues, and technologies, we do attempt to provide a range based on the collective author’s experience in processing and analyzing multiple species and organs to guide the less experienced user. An empty version of this table can be downloaded as Table S1 and freely used for publication. All co-authors collectively commit to start using this table in all publications.

Finally, analysis scripts, software environments, and (ideally) visualization portals should be made publicly available on established repositories/portals. Where possible, a well-documented code can be stored in dedicated repositories. Analysis environments should at minimum be well-documented (with versions specified for all software packages used for analysis).

Many of these tools are only valid for single-cell transcriptomic or chromatin accessibility methods that use DNA sequence as an output and are thus incompatible with other data modalities such as proteomics or metabolomics. While these data types are outside the scope of this review, they too should adhere to FAIR principles (Wilkinson et al., 2016) to ensure that they can be integrated with other datasets when the computational infrastructure develops. These data management tools are also not guaranteed to be future-proof. New technologies may arise that will render existing datasets/tools obsolete. Thus, it is imperative that, as the field evolves, we do not forget about legacy data, and ensure that it is preserved in a way that will be useful before this becomes impossible.

**Conclusion and Outlook**

Even for more advanced users, it is challenging to keep up with the rapidly evolving field of sc/snRNA-seq. Continuous advancements are not limited to the actual technology, but also include the choice of which technology to use for a specific problem, which method to use to isolate high-quality cells or nuclei, and how to analyze, compare and store these vast amounts of data. In this commentary, we discuss recommendations regarding data generation, analysis, storage and documentation to ensure transparency in publication and optimal use of the generated data across experiments, tissues, species and laboratories. We collectively commit to following the guidelines and recommendations (**Table 1**). Future research and method maturation will allow us to fine-tune and expand these guidelines and recommendations.

**Supplemental Material**

Supplemental Table S1. Necessary reported information to allow evaluation and repetition of a plant single cell/nucleus experiment.
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Author contributions

CS, TE, JL, HN, KK, MTR, TG, ML and BDR conceived this initiative; CS, TE, DS, MN, LSB, YK, RS, HN, JL, SG, OCM, BC, KK, MTR, GP, TG, ML and BDR wrote a first draft version which was commented on and finished by all co-authors.

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Figure 1. Overview of plant single-cell and single-nuclei experiments.
A. Number of publications describing sc/snRNA-seq data in the plant field per year from 2019 until 2022.
B. Distribution of species used in these papers (n:46). C. Overview of the different single-cell/nucleus technologies and their usage in the plant field. Example references used: a. (Farmer et al., 2021; Cervantes-Pérez et al., 2022; Conde et al., 2022; Neumann et al., 2022; Sun et al., 2022; Liu et al., 2022b; Li et al., 2023; Guillotin et al., 2023); b. (Tian et al., 2020; Marand et al., 2021); c. (Wendrich et al., 2020; Graeff et al., 2021; Lopez-Anido et al., 2021; OrtizRamírez et al., 2021; Wang et al., 2021b; Apelt et al., 2022; Otero et al., 2022; Kim et al., 2023); d. (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulse et al., 2019; Turco et al., 2019; Zhang et al., 2019; Satterlee et al., 2020; Bezrutczyk et al., 2021; Chen et al., 2021; Gala et al., 2021; Kim et al., 2021; Liu et al., 2021; Ma et al., 2021; Yang et al., 2021; Zhang et al., 2021a, 2021b; Li et al., 2022; Shahan et al., 2022; Tao et al., 2022); e. (Kao et al., 2021; Picard et al., 2021; Sunaga-Franze et al., 2021; Abramson et al., 2022; Li et al., 2023); f. none; g. (Efroni et al., 2016; Lopez-Anido et al., 2021; Roszak et al., 2021; Serrano-Ron et al., 2021; Omary et al., 2022); h. (Zong et al., 2022); and i. (Nelms and Walbot, 2019; Song et al., 2020; Xie et al., 2022).
Figure 2. Challenges in plant single-cell and single-nuclei transcriptomics.
Overview of the different steps of performing sc/snRNA-seq in plant samples and summary of how the most important challenges discussed here influence this flow-chart.
Figure 3. Workflow for sc/snRNA-seq data analysis.

Figure 4. Overview of common problems in sc/snRNA-seq analysis and possible solutions.
Figure 5. Cluster annotation and appropriate labels. A common technique to represent single-cell RNA-seq data requires mapping of each cell’s transcriptome to a low (typically two-) dimensional domain (e.g. tSNE or UMAP), after which highly similar cells group together into initially unannotated clusters. Annotating cells to a cell type or developmental stage is important for further interpretation of transcriptome signatures. Traditional, manual annotation methods screen differentially expressed genes within each cluster for the presence of individual marker genes or by transferring knowledge on cluster annotation from a reference data set to an unannotated data set (reviewed in (Clarke et al., 2021)). Without prior knowledge of marker genes and reference data sets, automated annotation methods (e.g. eager, lazy and marker learning methods (Xie et al., 2021)) can aid in assigning labels based on comparing cell cluster-specific genes and their biological functions. Cells and clusters that cannot be annotated with high confidence to (only) one cell type must be analyzed carefully to determine if they represent a mix of sub-cell types and/or cell states. Although there is currently no standardized definition of a “cell type” or a “cell state” (discussed in more detail in (Amini et al., 2023)), it has been proposed that a cluster with homogenous marker gene expression among all cells likely represent a cell type, while expression gradients among the cells within a cluster represent cell states (Clarke et al. 2021). However, as these definitions are still evolving, caution should be taken to not mix different anatomical levels (cell types, tissues or organs) and cellular processes (e.g. cell division, cell cycle) within the same visual representation. To avoid confusion, we recommend using multiple figures with different levels of labels (e.g. separate cell types, cell division states, tissues and so on).
Table 1. Necessary reported information to allow evaluation and repetition of a plant single-cell/nucleus experiment. A downloadable empty version for use in publications can be found as Supplemental Table S1. If numerical values deviate from the recommended numbers provided below, an explanation should be provided. The numerical values in this table are derived from available studies in plants, most of which originated from Arabidopsis.

<table>
<thead>
<tr>
<th>Details</th>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biological material</strong></td>
<td></td>
</tr>
<tr>
<td>Species</td>
<td>e.g. <em>Arabidopsis thaliana</em>, Zea mays</td>
</tr>
<tr>
<td>Accession</td>
<td>e.g. Col-0</td>
</tr>
<tr>
<td>Genotype</td>
<td>e.g. WT or mutant background</td>
</tr>
<tr>
<td>Tissue type</td>
<td>e.g. root, leaf, stem, seed</td>
</tr>
<tr>
<td>Detailed growth conditions</td>
<td>e.g. temperature, light conditions, medium etc.</td>
</tr>
<tr>
<td>Harvest conditions</td>
<td>e.g. age of plants, time of day, amount harvested</td>
</tr>
<tr>
<td><strong>Sample preparation</strong></td>
<td></td>
</tr>
<tr>
<td>Isolation protocol</td>
<td>Short description of the way the sample was isolated</td>
</tr>
<tr>
<td>Tissue dissection</td>
<td>e.g. razor blades, needles, tissue homogenizer</td>
</tr>
<tr>
<td>Fixation</td>
<td>Short description of the way the sample was fixed if this was done</td>
</tr>
<tr>
<td>Cell/nuclei enrichment</td>
<td>e.g. sucrose gradient, FACS (incl. model, nozzle size &amp; temperature)</td>
</tr>
<tr>
<td>Total sample preparation time</td>
<td></td>
</tr>
<tr>
<td>For cells: &lt;90 min for Arabidopsis roots (from material harvest to cell loading)*</td>
<td></td>
</tr>
<tr>
<td>*Duration may increase depending on starting material and time needed for optimal tissue digestion</td>
<td></td>
</tr>
<tr>
<td>For nuclei: 30-60 min (depending whether a nuclei enrichment step is included)</td>
<td></td>
</tr>
<tr>
<td>Estimated cell/nuclei number loaded</td>
<td>An estimation of the amount of cells or nuclei loaded, based on the cell/nuclei concentration and volume that was loaded</td>
</tr>
<tr>
<td>Instrument/Method/Kit</td>
<td>e.g. 10X Genomics 3‘ v3.1, BD Rhapsody WTA</td>
</tr>
<tr>
<td>Cell viability test</td>
<td></td>
</tr>
<tr>
<td>For cells: Trypan blue, fluorescein diacetate, calcein, propidium iodide, 4´,6-Diamidino-2-phenylindol</td>
<td></td>
</tr>
<tr>
<td>For nuclei: Not applicable</td>
<td></td>
</tr>
<tr>
<td><strong>Libraries</strong></td>
<td></td>
</tr>
<tr>
<td>Library construction</td>
<td>Protocol and revision/version that was followed, e.g. CG000204 Rev D for 3‘ v3.1</td>
</tr>
<tr>
<td>Amplification method</td>
<td>e.g. number of PCR cycles used for cDNA amplification</td>
</tr>
<tr>
<td><strong>End bias</strong></td>
<td>E.g. 3' end mainly; excess of rRNA or TSO sequences</td>
</tr>
<tr>
<td><strong>Sequence results</strong></td>
<td><strong>Instrument/method</strong></td>
</tr>
<tr>
<td><strong>Library layout/paired-end</strong></td>
<td>Consider to use standardized library structures (Booeshaghi et al., 2023)</td>
</tr>
<tr>
<td><strong>N° sequenced reads</strong></td>
<td>20,000-50,000/cell for RNA or more 20,000-40,000/nucleus for RNA or more</td>
</tr>
<tr>
<td><strong>Raw data</strong></td>
<td><strong>Reference genome</strong></td>
</tr>
<tr>
<td><strong>Annotation version</strong></td>
<td>If custom annotation, also include .gtf/.gff/gff3 files</td>
</tr>
<tr>
<td><strong>Mapping method (incl. software, customized settings)</strong></td>
<td>E.g. STAR (cellranger)</td>
</tr>
<tr>
<td><strong>Mapping efficiency</strong></td>
<td>&gt;85% for Arabidopsis*  * value may be lower in other species</td>
</tr>
<tr>
<td><strong>Sequencing saturation</strong></td>
<td>&gt;50%</td>
</tr>
<tr>
<td><strong>Estimation of ambient RNA</strong></td>
<td>Fraction of reads in cells &gt; 60% for scRNA-seq  Fraction of reads in cells &gt; 50% for snRNA-seq</td>
</tr>
<tr>
<td><strong>Imputation method and settings</strong></td>
<td>If relevant</td>
</tr>
<tr>
<td><strong>Processed data</strong></td>
<td><strong>N° captured cells/nuclei</strong></td>
</tr>
<tr>
<td><strong>N° high quality cells/nuclei</strong></td>
<td>E.g. 20% of estimated number</td>
</tr>
<tr>
<td><strong>Filter criteria: % mitochondrial reads/cell or nucleus</strong></td>
<td>&lt;10% for scRNA-seq* close to 0% for snRNA-seq*  *value may deviate depending on biological context</td>
</tr>
<tr>
<td><strong>Filter criteria: % chloroplast reads/cell or nucleus</strong></td>
<td>&lt;5-10% for scRNA-seq* close to 0% for snRNA-seq*  *values may be higher depending on biological context</td>
</tr>
<tr>
<td><strong>Filter criteria: Minimum N° UMI/cell or nucleus</strong></td>
<td>&gt;1000 for scRNA-seq  &gt;400 for snRNA-seq</td>
</tr>
<tr>
<td><strong>N° total detected transcripts</strong></td>
<td>Dependent on the species but e.g. 60% of total number of transcripts in the annotated genome</td>
</tr>
<tr>
<td><strong>Doublet rate</strong></td>
<td>Estimates according to 10X Genomics user guide based on number of loaded cells/nuclei</td>
</tr>
<tr>
<td><strong>Replicate comparisons</strong></td>
<td>Provide coefficient correlation of the most variable genes or cluster-specific genes between independent replicates or compare pseudo-bulk from sc/snRNA-seq vs. bulk RNA-seq</td>
</tr>
<tr>
<td>--------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Batch correction method for merging (incl. reasoning for batch correction)</strong></td>
<td>e.g. Seurat, Harmony</td>
</tr>
<tr>
<td><strong>Additional processing</strong></td>
<td>e.g. removal of protoplast induced genes, cell cycle regression, noise (ambient RNA) removal, cluster membership bias between replicates, removal of low quality clusters</td>
</tr>
</tbody>
</table>
| **Validation** | **Method of automatic annotation of clusters**  
Method of manual annotation (markers, gene function info)  
Verification in planta (e.g. Number of markers used for validation)  
**Method of manual annotation (markers, gene function info)**  
**Verification in planta (e.g. Number of markers used for validation)** |
| **Data availability** | Analysis scripts & codes (GitHub)  
Excel Tables DEG for each cluster  
Objects/count matrix in repository (which one, where?)  
On-line tool/browser URL  
Cell-level metadata table  
**Additional** |
| **Data availability** | Analysis scripts & codes (GitHub)  
Excel Tables DEG for each cluster  
Objects/count matrix in repository (which one, where?)  
On-line tool/browser URL  
Cell-level metadata table  
**Additional** |
| **Data availability** | If relevant  
e.g. Lists for each cell type/developmental stage from FindMarkers (Seurat)  
e.g. use NCBI GEO to store count matrices and Seurat object  
List the URL if relevant  
Include cell type annotations for each cell barcode  
e.g. annotation/counting of rRNA, allow for rRNA estimation, consideration of intronic reads |
References


