Asc-Seurat: Analytical single-cell Seurat-based web application

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Asc-Seurat (Analytical single-cell Seurat-based web application) is a web application based on Shiny¹. Pronounced as "ask Seurat", it provides a click-based, easy-to-install, and easy-to-use interface that allows the execution of all steps necessary for scRNA-seq analysis (See *Asc-Seurat workflow*). It integrates many of the capabilities of the Seurat² and Dynverse³ and also allows an instantaneous functional annotation of genes of interest using BioMart⁴.

Asc_seurat relies on multiple R packages. Please, visit the *references* and check the complete list of packages and their references.

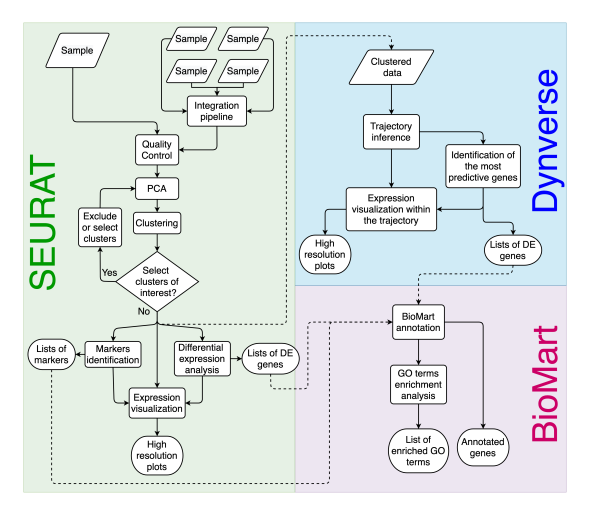


Fig. 1: **Asc-Seurat workflow overview.** Asc-Seurat is built on three analytical cores. Using Seurat, it is possible to explore scRNA-seq data of a population of cells to identify patterns that reflect the cell types of a sample(s) and identify markers and DEGs for each cell type/cluster. By incorporating Dynverse, Asc-Seurat allows the utilization of dozens of models to infer and visualize developmental trajectories (V and VI) and to identify genes differentially expressed on those trajectories (VII). Finally, using BioMart, Asc-Seurat allows immediate functional annotation and GO terms enrichment analysis for many species.

¹ shiny.rstudio.com/

² satijalab.org/seurat/

³ dynverse.org

⁴ www.biomart.org

INSTALLATION

1.1 Dependencies

Asc-Seurat relies on multiple R packages and their dependencies (See *References*). However, we provide a Docker image that contains all necessary software and packages.

To install Asc-Seurat, it is necessary to have Docker installed on the machine. Docker needs to be correctly installed and configured in the user's machine. Check the installation instructions provided by Docker at https://docs.docker. com/engine/install.

Warning: Single-cell RNA-seq data analysis can be resource-consuming. By default, Docker will use (allocate) only a fraction of your RAM memory. A minimum requirement of 8 Gb of RAM memory was necessary to analyze a dataset containing around eight thousand cells during our tests. Therefore, users need to adjust the amount of allocated memory according to their dataset. Please visit: https://docs.docker.com/docker-for-mac/space/ (MAC) or https://docs.docker.com/docker-for-windows/ (Windows) to learn how to make this adjustment.

1.1.1 Image download

After installing Docker, users can download the Docker image containing Asc-Seurat by executing the command below. The installation is quick and straightforward. After that, everything is set.

```
# Download the docker image:
docker pull kirstlab/asc_seurat
```

1.2 Starting Asc-Seurat

After downloading the image, users can start the app on their working directory. See below for the instructions on how to start the app in the different operational systems.

Note: During the first execution, some folders will be created in the working directory. They include the folders data/ and RDS_files/ that users will use to store their datasets, allowing Asc-Seurat to read them.

Always start the run inside the working directory to be able to use the data inside these folders.

1.2.1 For macOS and Linux

Tip: The code below will automatically update Asc-Seurat to the latest version. You can download and execute a specific version of Asc-Seurat by adding the version's tag to the image's name, i.e., replace kirstlab/asc_seurat by kirstlab/asc_seurat:v2.1 to use v2.1.

Note: After executing the "docker run" command, open your preferred web browser and paste the address http: //localhost:3838/. Asc-Seurat should be ready.

If users want to kill the Docker container, run the command below.

docker kill Asc_seurat

1.2.2 For Windows

To run Asc-Seurat on Windows via Docker, it is necessary to use Windows 10. Moreover, Windows Subsystem for Linux (WSL) needs to be installed. Before running Asc-Seurat, users must guarantee that Docker and its WSL 2 components are correctly installed and running. For that, check the two (sequential) tutorials below:

- 1. Docker installation info
- 2. Define windows WSL 2 as default (If you followed the link above correctly, you only need to execute step 5 of this tutorial).

The tutorials above contain all the necessary information to install Docker on Windows. However, it is also possible to find video tutorials on YouTube. Check the following link for an example: https://youtu.be/5nX8U8Fz5S0.

After certifying that everything is working, Asc-Seurat can be started using the commands below:

Tip: The code below will automatically update Asc-Seurat to the latest version. You can download and execute a specific version of Asc-Seurat by adding the version tag to the image's name, i.e., replace kirstlab/asc_seurat by kirstlab/asc_seurat:v.1.0 to use v1.0.

```
# Create the working directory
mkdir my_project
cd my_project
# If using Windows CMD
docker pull kirstlab/asc_seurat && docker run -v "%cd%:/app/user_work" -v /var/run/
idocker.sock:/var/run/docker.sock -v /tmp:/tmp -d --rm -p 3838:3838 kirstlab/asc_
idocker at the seurat
```

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Note: After executing the "docker run" command, open your preferred web browser and paste the address http: //localhost:3838/. Asc-Seurat should be ready.

If users want to kill the Docker container, run the command below.

docker kill Asc_seurat

REFERENCES

Asc-Seurat is built on the work of many other people and relies on a diversity of R packages. These packages, in turn, have many dependencies. Here, we list all packages that Asc-Seurat directly calls.

2.1 Analytical core

Three packages are the analytical core of Asc-Seurat. Below is listed their information.

- Seurat
 - web page: https://satijalab.org/seurat/
 - Publications:
 - * Satija, R. et al. (2015) Spatial reconstruction of single-cell gene expression data. Nat. Biotechnol., 33, 495–502.
 - * Stuart, T. et al. (2019) Comprehensive Integration of Single-Cell Data. Cell, 177, 1888–1902.e21.
- Dynverse (dynplot, dynwrap, and dynfeature)
 - web page: https://dynverse.org/
 - Publications:
 - * Saelens, W. et al. (2019) A comparison of single-cell trajectory inference methods. Nat. Biotechnol., 37, 547–554.
- biomaRt
 - web page: https://bioconductor.org/packages/release/bioc/html/biomaRt.html
 - Publications:
 - * Durinck, S. et al. (2009) Mapping identifiers for the integration of genomic datasets with the R/Bioconductor package biomaRt. Nat. Protoc., 4, 1184–1191.

2.2 Additional packages

2.2.1 CRAN

- circlize: https://cran.r-project.org/web/packages/circlize/
- DT: https://cran.r-project.org/web/packages/DT/
- dplyr: https://mran.microsoft.com/web/packages/dplyr/index.html

- future: https://cran.r-project.org/web/packages/future/index.html
- ggplot2: https://cran.r-project.org/web/packages/ggplot2/index.html
- ggthemes: https://cran.r-project.org/web/packages/ggthemes/index.html
- hdf5r: https://cran.r-project.org/web/packages/hdf5r/
- metap: https://cran.r-project.org/web/packages/metap/index.html
- patchwork: https://cran.r-project.org/web/packages/patchwork/index.html
- rclipboard: https://cran.r-project.org/web/packages/rclipboard/index.html
- reactable: https://cran.r-project.org/web/packages/reactable/index.html
- · scales: https://cran.r-project.org/web/packages/scales/index.html
- sctransform: https://cran.r-project.org/web/packages/sctransform/index.html
- SeuratObject: https://cran.r-project.org/web/packages/SeuratObject/index.html
- shiny: https://cran.r-project.org/web/packages/shiny/index.html
- · shinycssloaders: https://cran.r-project.org/web/packages/shinycssloaders/index.html
- shinyFeedback: https://cran.r-project.org/web/packages/shinyFeedback/index.html
- · shinyWidgets: https://cran.r-project.org/web/packages/shinyWidgets/index.html
- tidyverse: https://cran.r-project.org/web/packages/tidyverse/index.html
- utils: https://cran.r-project.org/web/packages/R.utils/index.html

2.2.2 Bioconductor

- ComplexHeatmap: https://bioconductor.org/packages/release/bioc/html/ComplexHeatmap.html
- glmGamPoi: https://bioconductor.org/packages/release/bioc/html/glmGamPoi.html
- multtest: https://bioconductor.org/packages/release/bioc/html/multtest.html
- SingleCellExperiment: https://bioconductor.org/packages/release/bioc/html/SingleCellExperiment.html
- slingshot: https://bioconductor.org/packages/release/bioc/html/slingshot.html
- topGO: https://bioconductor.org/packages/release/bioc/html/topGO.html
- tradeSeq: https://bioconductor.org/packages/release/bioc/html/tradeSeq.html

THREE

SESSION INFORMATION

3.1 Session information of the Asc-Seurat's environment (Docker image v2.2)

sessionInfo()

R version 4.0.1 (2020-06-06) Platform: x86_64-pc-linux-gnu (64-bit) Running under: Debian GNU/Linux bookworm/Matrix products: default

BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.10.0 LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0

locale:

LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C LC_TIME=en_US.UTF-8 LC_COLLATE=en_US.UTF-8 LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8 LC_PAPER=en_US.UTF-8 LC_NAME=C LC_ADDRESS=C LC_TELEPHONE=C LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

parallel stats4 grid stats graphics grDevices utils datasets methods base

other attached packages:

glmGamPoi_1.2.0 dynfeature_1.0.0 dynwrap_1.2.2 dynplot_1.1.2 topGO_2.42.0 SparseM_1.81 GO.db_3.12.1 AnnotationDbi_1.52.0 graph_1.68.0 biomaRt_2.46.3 multtest_2.46.0 slingshot_1.8.0 princurve_2.1.6 SingleCellExperiment_1.12.0 SummarizedExperiment_1.20.0 Biobase_2.50.0 GenomicRanges_1.42.0 GenomeInfoDb_1.26.7 IRanges_2.24.1 S4Vectors_0.28.1 BiocGenerics_0.36.1 MatrixGenerics_1.2.1 matrixStats_0.61.0 ComplexHeatmap_2.6.2 vroom_1.5.7 scales_1.1.1 hdf5r_1.3.5 DT_0.20 shinycssloaders_1.0.0 ggthemes_4.2.4 future_1.23.0 rclipboard_0.1.5 shinyFeedback_0.4.0 shinyWidgets_0.6.4 shiny_1.7.1 sctransform_0.3.3 reactable_0.2.3 circlize_0.4.13 patchwork_1.1.1 SeuratObject_4.0.4 Seurat_4.1.0 forcats_0.5.1 stringr_1.4.0 dplyr_1.0.7 purr_0.3.4 readr_2.1.2 tidyr_1.2.0 tibble_3.1.6 ggplot2_3.3.5 tidyverse_1.3.1

loaded via a namespace (and not attached):

rappdirs_0.3.3 scattermore_0.7 GA_3.2.2 bit64_4.0.5 irlba_2.3.5 DelayedArray_0.16.3 data.table_1.14.2 rpart_4.1-15 RCurl_1.98-1.5 generics_0.1.2 cowplot_1.1.1 RSQLite_2.2.9 RANN_2.6.1 carrier_0.1.0 bit_4.0.4 tzdb_0.2.0 spatstat.data_2.1-2 xml2_1.3.3 lubridate_1.8.0 httpuv_1.6.5 assertthat_0.2.1 viridis_0.6.2 fontawesome_0.2.2 hms_1.1.1 jquerylib_0.1.4 promises_1.2.0.1 fansi_1.0.2 progress_1.2.2 dbplyr_2.1.1 readxl_1.3.1 igraph_1.2.11 DBI_1.1.2 htmlwidgets_1.5.4 spatstat.geom_2.3-1 dyndimred_1.0.4 ellipsis_0.3.2 backports_1.4.1 RcppParallel_5.1.5 deldir_1.0-6 vctrs_0.3.8 remotes_2.4.2 Cairo_1.5-14 ROCR_1.0-11 abind_1.4-5 cachem_1.0.6 withr_2.4.3 ggforce_0.3.3 prettyunits_1.1.1 goftest_1.2-3 cluster_2.1.0 ape_5.6-1 lazyeval_0.2.2 crayon_1.4.2 babelwhale_1.0.3

pkgconfig_2.0.3 tweenr_1.0.2 nlme_3.1-148 rlang_1.0.0 globals_0.14.0 lifecycle_1.0.1 miniUI_0.1.1.1 BiocFile-Cache_1.14.0 modelr_0.1.8 cellranger_1.1.0 rprojroot_2.0.2 polyclip_1.10-0 lmtest_0.9-39 Matrix_1.4-0 zoo_1.8-9 reprex_2.0.1 ggridges_0.5.3 GlobalOptions_0.1.2 processx_3.5.2 png_0.1-7 viridisLite_0.4.0 rjson_0.2.21 bitops_1.0-7 KernSmooth_2.23-17 blob_1.2.2 shape_1.4.6 parallelly_1.30.0 memoise_2.0.1 magrittr_2.0.2 plyr_1.8.6 ica_1.0-2 zlibbioc_1.36.0 compiler_4.0.1 RColorBrewer_1.1-2 clue_0.3-60 fitdistrplus_1.1-6 cli_3.1.1 XVector_0.30.0 listenv_0.8.0 pbapply_1.5-0 ps_1.6.0 MASS_7.3-51.6 mgcv_1.8-31 tidyselect_1.1.1 stringi_1.7.6 yaml_2.2.2 askpass_1.1 ggrepel_0.9.1 sass_0.4.0 tools_4.0.1 future.apply_1.8.1 rstudioapi_0.13 foreach_1.5.1 gridExtra_2.3 farver_2.1.0 Rtsne_0.15 ggraph_2.0.5 proxyC_0.2.4 digest_0.6.29 dynparam_1.0.2 Rcpp_1.0.8 broom_0.7.12 later_1.3.0 RcppAnnoy_0.0.19 httr_1.4.2 colorspace_2.0-2 ranger_0.13.1 rvest_1.0.2 XML_3.99-0.8 fs_1.5.2 tensor_1.5 reticulate_1.24 splines_4.0.1 uwot_0.1.11 lmds_0.1.0 spatstat.utils_2.3-0 graphlayouts_0.8.0 plotly_4.10.0 xtable_1.8-4 jsonlite_1.7.3 tidygraph_1.2.0 R6_2.5.1 pillar_1.7.0 htmltools_0.5.2 mime_0.12 glue_1.6.1 fastmap_1.1.0 codetools_0.2-16 utf8_1.2.2 lattice_0.20-41 bslib_0.3.1 spatstat.sparse_2.1-0 curl_4.3.2 leiden_0.3.9 openssl_1.4.6 survival_3.1-12 desc_1.4.0 dynutils_1.0.6 munsell_0.5.0 GetoptLong_1.0.5 GenomeInfoDbData_1.2.4 iterators_1.0.13 haven_2.4.3 reshape2_1.4.4 gtable_0.3.0 spatstat.core_2.3-2

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FIVE

LOADING THE DATA OF AN INDIVIDUAL SAMPLE

5.1 Location of the dataset

For Asc-Seurat to read the datasets, they need to be located in a subdirectory inside the data/ directory. The data/ directory will be created during the installation and contains a subdirectory with an example dataset called example_PBMC/. This dataset is from the publicly available $10\times$'s Peripheral Blood Mononuclear Cells (PBMC) and contains 2700 cells.

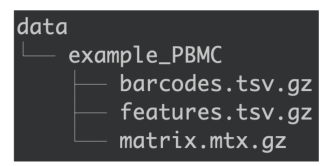


Fig. 1: Organization of the data/ directory.

Therefore, to add the dataset, create a subdirectory inside data/ containing the counts' matrix (*matrix.mtx.gz*), cell barcodes (*barcodes.tsv.gz*), and gene names (*features.tsv.gz*).

Asc-Seurat provides separated environments (tabs) to analyze a single sample and the integrated analysis of multiple samples.

5.2 Format of the dataset

Asc-Seurat can only read the input files in the format generated by Cell Ranger (10x genomics). However, it is possible to convert your counts' matrix to the acceptable format. For example, the function write10xCounts(), from the DropletUtils package, is an easy option to make this conversion.

Tip: Using write10xCounts(), users can provide as output the path to the data/ directory. In this way, Asc-Seurat can recognize the files automatically.

5.3 Loading the data

To analyze an individual sample, select the second tab in the web application, named One sample. Then, choose the sample to analyze and set the initial criteria to exclude cells that should not be load, as shown below.

After inserting the datasets in the data/ directory, the samples will be available to load in Asc-Seurat, as shown below.

Introduction	One sample	Integration of multiple samples	Trajectory inference	Advanced plots	BioMart ar	nnotation			
-		n save a bookmark (purple bo oad all selected parameters a	-			neter choices. Using the saved Ilts.			
Choose the samp	le to be analyz	ed and the initial requirements t	o load the data. Note t	hat cells that do no	t match the	e parameters will not be load.			
These parameters are used to exclude low-quality cells and allow the data to load quickly. Users can add more restrictive parameters after visualizing the distributions in the next section.									
After selecting t	he parameters	, click on the blue button to lo	oad the data.						
Select the sample	e to use	Project name	Min. number of cells	Min. number o	-	Common identifier of mitochondrial			
./data/example_	./data/example_PBMC								
			3	200	٢	^MT-			
						Load data of the selected sample			

Fig. 2: Example of how to load an individual sample for analysis and of the requested initial parameters.

In the first box to the left, it is possible to select the sample to use. However, there are a few parameters that need to provide before loading the data. This step is based on Seurat's functions CreateSeuratObject and PercentageFeatureSet. Between parenthesis, we list the name of the parameter in the CreateSeuratObject function.

Below is a description of these parameters:

- **Project name**: Sets the name for the project. The name will appear in some of the plots, but it is not required (project).
- Min. number of cells expressing a gene: Include genes only if they are detected in at least this many cells (min.cells).
- Min. number of genes a cell must express to be included: Include cells only if they expressed at least this number of genes (min.features).
- **Regex to identify mitochondrial genes**: Here, the regular expression (Regex) is a sequence of characters that is used to determine the genes belonging to the mitochondrial genome (pattern). For example, when using the human genome, this sequence should be "^MT-".

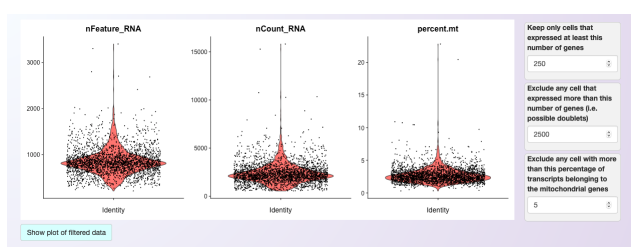
After setting the parameters described above, click on the button *Load data of the selected sample* to start the analysis. A violin plot showing the distribution of cells will appear. This plot can then be used to set more restrictive parameters for *quality control*.

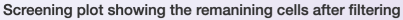
QUALITY CONTROL

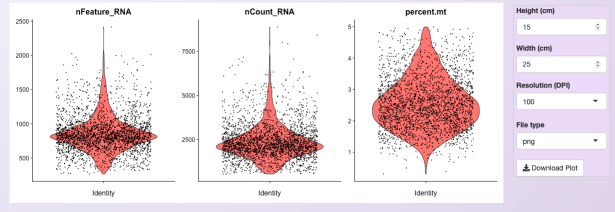
After loading the data, a violin plot will be generated showing the distribution of cells according to three parameters:

- nFeature_RNA: the number of genes detected in each cell
- nCount_RNA: the number of molecules detected per cell
- percent.mt: the percentage of transcripts that map to mitochondrial genes

After visualizing the distribution of cells, it is possible to set more restrictive parameters (on the right side of the plot) and filter cells based on the number of expressed genes per cell and the percentage of transcripts from mitochondrial genes. By clicking on *Show plot of filtered data*, users can see the distribution of cells after filtering and then readjust the parameters. The figure below shows the distribution of cells of the PBMC dataset before and after filtering.







Asc-Seurat allows users to download each of the plots with high-resolution by clicking on the Download plot button.

SEVEN

CLUSTERING

After filtering the data to remove low-quality cells, Asc-Seurat allows clustering the remaining cells according to their expression profiles. However, before clustering, a series of steps are executed, including normalization, scaling (if using LogNormalization), and dimensional reduction via PCA.

Moreover, users need to decide how many dimensions are to be used during the clustering after executing the PCA. Asc-Seurat provides an elbow plot to inform this decision. Below are instructions on how to perform the clustering depending on the normalization method of choice.

7.1 Normalization

7.1.1 LogNormalization

Asc-Seurat allows the normalization using Seurat's LogNormalize function. Users have the option to change the scaling factor if necessary, but it is typically not needed. In the same window (see the image below), users can select what method should be used to identify the most variable genes and how many of the most variable genes should be used during the dimension reduction (PCA).

The most variable genes exhibit high cell-to-cell variation in the dataset and therefore are more informative. We use Seurat's function FindVariableFeatures. The default setting should work well for the majority of cases.

Select the normalization method	Scale factor	Run the PCA analysis
 LogNormalize SCTransform 	10000	
	Select the method to detect the most variable genes	
	• vst	
	 mean.var.plot (mvp) 	
	o dispersion (disp)	
	Number of variable genes for PCA	
	3000	

7.1.2 SCTransform

The second option of normalization provided by Asc-Seurat is Seurat's Seurat's SCTransform. When using this normalization, it is unnecessary to set the scale factor or identify the most variable genes (See image below).



Note: Currently, the recommendation of Seurat's team is to use the standard "RNA" assay when performing differential expression (D.E) analysis and for data visualization, even when using SCTransform (See here). Therefore, Asc-Seurat will use the SCTransformed data ("SCT" assay) until the clustering step only.

To use the "RNA" assay after SCTransform, Asc-Seurat will automatically perform the LogNormalization and scaling of the data in the RNA assay by applying the default parameters.

7.2 Dimensional reduction (PCA)

The PCA will be executed using Seurat's function RunPCA and, after its conclusion, an elbow plot is generated automatically, to help users to decide how many PCs should be included to inform the clustering step.

Users can use this plot to select the PCs with the highest standard deviation (more informative PCs). Also, users should set the number of PCs to include during clustering in the windows at the plot's right side.

In the example below, only the first 10 PCs are selected. Not that the resulting plot will be slightly different depending on the normalization method. Below we show the plot obtained using the LogNormalization.



Fig. 1: Elbow plot provided to help to select the most informative PCs. For the PBMC dataset, and using the LogNormalization method, we chose the ten first PCs.

7.3 Clustering of cells

The next step is the clustering of the cells. For that, Asc-Seurat used both FindNeighbors and FindClusters functions of the Seurat package.

Before the execution, however, users need to set a value for the resolution parameter. The resolution is an important parameter to evaluate because it determines the profile and number of clusters identified for a dataset. Selecting larger values will favor splitting cells into more clusters while choosing a smaller value has the opposite effect. Quoting from Seurat's tutorial: "We find that setting this parameter between 0.6-1.2 typically returns good results for single-cell datasets of around 3K cells. Optimal resolution often increases for larger datasets".

Tip: There is no easy way to define an optimal value for the resolution parameter. Users need to try different values and evaluate the resulting clusters according to the expectation for their cells population. Visualizing the expression profile of cell-type-specific markers can provide a hint if the chosen value is too small or too large.

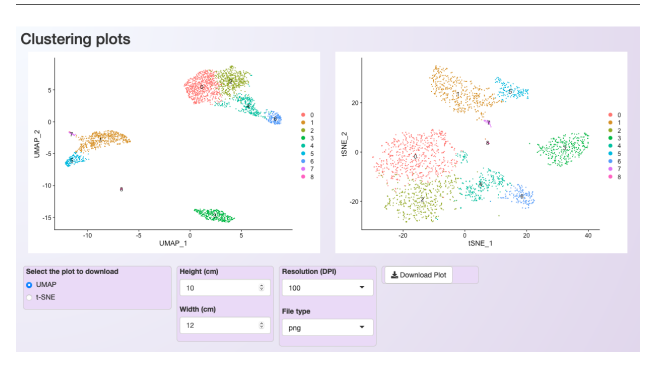


Fig. 2: Plot showing the clustering of the PBMC dataset after LogNormalization, using 10 PCs and a resolution value of 0.5.

After the execution of the clustering step, two plots are generated for cluster visualization. The first plot is generated using the Uniform Manifold Approximation and Projection (UMAP) technique (left). The second deploys the t-distributed Stochastic Neighbor Embedding (t-SNE) method (right).

7.3.1 Selecting clusters of interest

In some cases, it is interesting to select or exclude some clusters of cells from the dataset before executing the subsequent steps. This process is helpful, for example, when users desire to explore a developmental trajectory of a specific group of cell types.

Asc-Seurat makes this step simple. Users only need to select the cluster(s) to keep or exclude and start reanalysis of the remaining cells by clicking on *Reanalyze after selection/exclusion of clusters* (see below).

Do you want to select or exclude clusters of cells and reanalyze the data?	Do you want to select or exclude the clusters?	Choose clusters to select or exclude	Reanalyze after selection/exclusion of clusters	
• Yes	 Select 	0 -		
O No	• Exclude			

Fig. 3: Asc-Seurat makes it easy to select or exclude a cluster (or clusters) of cells. In this example, we exclude all cells belonging to cluster 0.

Asc-Seurat will then execute the steps with the new set of cells up to the PCA. Then, **users need to evaluate the elbow plot and decide the number of PCs to cluster the new set of cells**. Users can either keep the same value for the resolution parameter or modify it before clicking on *Run the clustering analysis* to start the clustering once more.

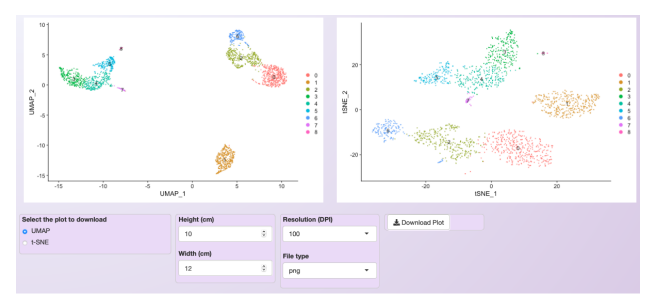


Fig. 4: Clustering of the PBMC dataset after excluding cells belonging to cluster 0 from the original dataset.

Warning: The cluster's numbering will change every time that cluster(s) are selected or excluded.

EIGHT

MARKERS IDENTIFICATION AND DIFFERENTIAL EXPRESSION ANALYSIS

After clustering the cells, users may be interested in identifying genes specifically expressed in one cluster (markers) or in genes that are differentially expressed among clusters of interest. Asc-Seurat can apply multiple algorithms to identify gene markers for individual clusters or to identify differentially expressed genes (DEGs) among clusters, using Seurat's functions FindMarkers and FindAllMarkers.

Asc-Seurat allows users to filter gene markers and DEGs by the fold change and minimal percentage of cells expressing a gene in the cluster(s). Moreover, users can define a significance level to exclude genes based on the adjusted p-value (see below).

Select the analysis to perform?	Select the (log) fold change threshold		
 Identify markers for all clusters 	0.25		
 Identify markers for one specific cluster Identify markers distinguishing a cluster from other(s) cluster(s) 	Select the minimal percentage of cells expressing a gene for this gene to be tested		
	0.1 🕀		
	Select the statistical test		
	wilcox -		
	Select the (adjusted) p-value threshold		
	0.05		
	Search for markers/D.E. genes		

Fig. 1: Example of Asc-Seurat's interface showing the settings to the search for gene markers for each of the clusters using the Wilcox test.

An iterative table will be available after executing the search for marker or DEGs, showing the significant genes. Moreover, users can download the list of significant markers or DEGs as a csv file.

The list of genes in the csv can then be used to visualize their gene expression in a series of plots, as shown in the section *Expression visualization*.

Select the analysis to perform?	Select the cluster of interest	Select the (log) fold change threshold
 Identify markers for all clusters 	0 -	0.25
 Identify markers for one specific cluster Identify markers distinguishing a cluster from other(s) cluster(s) 		Select the minimal percentage of cells expressing a gene for this gene to be tested
		0.1
		Select the statistical test
		wilcox
		Filter only positive markers?
		• yes
		○ no
		Select the (adjusted) p-value threshold
		0.05
		Search for markers/D.E. genes

Fig. 2: Example of Asc-Seurat's interface showing the settings to the search for markers for a specific cluster (cluster 0).

Select the analysis to perform?	Select the cluster of interest	Select the (log) fold cl	hange threshold
 Identify markers for all clusters 	0 -	0.25	٢
 Identify markers for one specific cluster Identify markers distinguishing a cluster from other(s) cluster(s) 	Select the cluster(s) to compare 2, 3	Select the minimal pe cells expressing a ger to be tested	-
		0.1	٢
		Select the statistical t	est
		wilcox	•
		Filter only positive ma	irkers?
		 yes 	
		O no	
		Select the (adjusted) threshold	p-value
		0.05	٢
		Search for marker	s/D.E. genes

Fig. 3: Example of Asc-Seurat's interface showing the settings to search for DEGs genes among clusters 0, 2, and 3.

geneID	cluster	p_val	avg_log2FC	pct.1	pct.2	p_val_adj
MS4A1	3	0	3.37780648904012	0.855	0.053	0
CD79A	3	0	4.30987416981298	0.936	0.041	0
CD79B	3	5.26049675440333e- 274	3.48110412099492	0.916	0.142	7.21424524898873 270
LINC00926	3	5.18018664334955e- 272	2.84260707176794	0.564	0.009	7.10410796268958 268
TCL1A	3	2.03777874670445e- 270	3.59137563045096	0.622	0.022	2.79460977323049 266
HLA-DQA1	3	6.07567789191485e- 266	3.05695403560438	0.89	0.118	8.33218466097202 262
VPREB3	3	5.40891671486496e- 237	2.42474808636795	0.488	0.007	7.4177883827658 233
HLA-DQB1	3	2.18079190672984e- 229	3.07591649859714	0.863	0.148	2.99073802088931 225
CD74	3	5.9732566244471e-1 85	2.91969016185635	1	0.821	8.19172413476675 181
HLA-DRA	3	2.72038439500985e- 183	2.7613963732939	1	0.495	3.73073515931651 179
10 of 397 rows S	how 10 •			Previ	ous 1 2 3 4	5 40 Ne

Fig. 4: List of the ten most significant markers identified for cluster 3 of the PBMC dataset (as defined in *Clustering*).

EXPRESSION VISUALIZATION

Asc-Seurat provides a variety of plots for gene expression visualization. From a list of selected genes, it is possible to visualize the average of each gene expression in each cluster in a heatmap. It also provides plots for the visualization of gene expression at the cell level. Moreover, violin plots and dot plots allow the visualization of each cluster's expression, emphasizing the inter-cluster comparison.

9.1 Format of the input file containing genes for expression visualization

Asc-Seurat expects as input a csv (comma-separated value) or a tsv (tab-separated value) file containing at least two columns. The first column must contain the gene ID as present in the dataset, and the second column is a grouping variable. An optional third column can contain the common names of each gene. Any additional column will be ignored. The output files generated by the differential expression analysis are already in the correct format to be used as input for the visualization.

Below is shown an example of an input file used for expression visualization. It contains ten markers identified for clusters 2 and 3. In this case, the dataset uses the gene name as an identifier, and this is the information on the first column. The second column is used to group de marker according to their clusters.

ing the required columns.				
IL32	Cluster_2			
LTB	Cluster_2			
LDHB	Cluster_2			
CD3D	Cluster_2			
IL7R	Cluster_2			
MS4A1	Cluster_3			
CD79A	Cluster_3			
CD79B	Cluster_3			
LINC00926	Cluster_3			
TCL1A	Cluster_3			

Table 1: Example of an input file for gene expression visualization showing the required columns.

After loading the input file, users can select what group(s) of genes to explore and choose specific genes from each group. Moreover, if a third column is provided in the input file, users can use the genes' common name instead of the gene IDs to select the genes to be shown.

9.2 Heatmap

Once users selected their genes of interest, they can generate a heatmap of the average of each gene expression in each cluster by clicking on the button *Show heatmap with the average of expression per cluster*. The heatmap will adjust its height according to the number of selected genes. Moreover, rows and columns will be organized by a hierarchical clustering algorithm. A high-resolution copy of the heatmap plot can be download in a diversity of formats.

Input the list of markers Browse PBMC_cluesters_mark Upload complete	Select the group of markers to test Cluster_3	Select the genes to show CD79A, CD79B, LINC00926, MS4/ •	Select the expression values to show counts normalized	Show heatmap
Does your file have a header?	Visualization options Show all genes Select genes to show		• normalized and scaled	
Load markers	Use IDs Use Name			

Heatmap

Note that the scale of colors of the heatmap is adjusted based on the expression of the selected genes

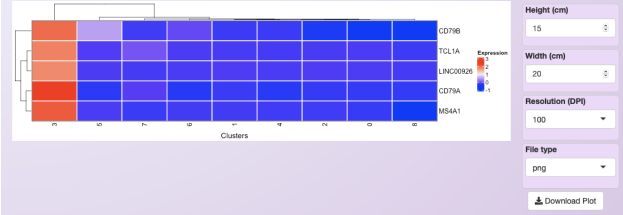


Fig. 1: Asc-Seurat's interface demonstrating the filtering options provided to select the genes for expression visualization. The heatmap shows the expression profile of the five most significant markers for cluster 3.

9.3 Gene expression at the cell level - Feature plots

From the list of genes on the heatmap, users can select genes to further explore by visualizing the expression at the cell level. For each selected gene, a couple of feature plots will be generated using Seurat's Feature plots function. The UMAP plot is shown side-by-side with the feature plots, so users can quickly compare the expression profile with the identified clusters.

9.4 Visualization of the expression among clusters

For each selected gene, Asc-Seurat will also generate plots to visualize the distribution of cells within each cluster according to the expression of the gene (violin plot) and the percentage of cells in each cluster expressing the gene

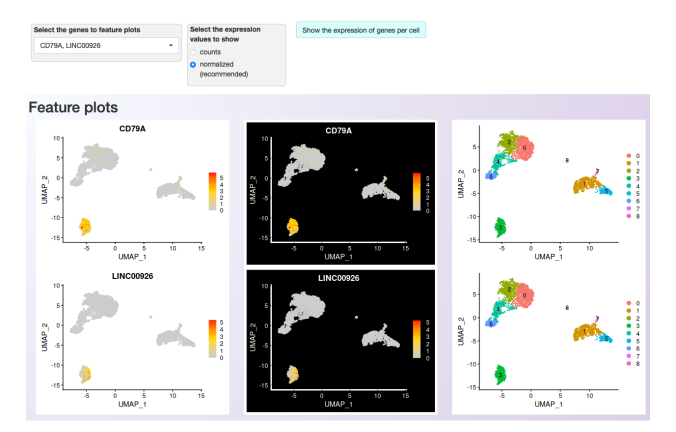
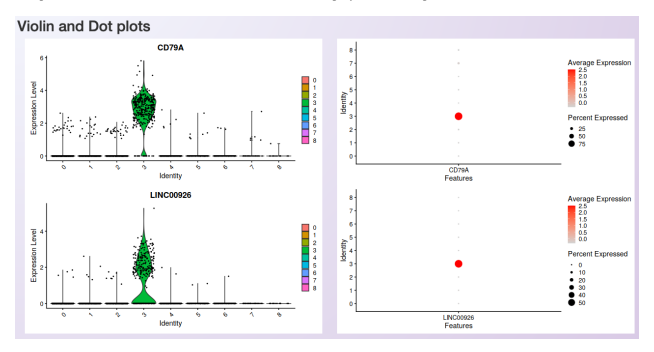


Fig. 2: Asc-Seurat's interface showing the filtering options provided to select the genes for expression visualization at the cell level. Two of the five genes shown on the heatmap were chosen for more detailed visualization.



(dot plot). Seurat's functions VlnPlot() and DotPlot() are deployed in this step.

Fig. 3: Visualization of the distribution of cells within each cluster according to the gene expression (violin plot; left) and the percentage of cells in each cluster expressing the gene (dot plot; right).

Tip: Sometimes, it is necessary to make fine adjustments to an image before publication. Saving the plots as a Scalable Vector Graphic (svg), allows the edition of all aspects of the plot by image edition software as Inkscape.

TEN

LOADING THE DATA AND INTEGRATION OF MULTIPLE SAMPLES

To analyze multiple samples, select the third tab in the web application, named Integration of multiple samples.

Note: The integration is based on Seurat's functions FindIntegrationAnchors and IntegrateData. For more information, see Seurat's integration tutorial and Stuart, T. et al. (2019).

10.1 Format of the dataset

Asc-Seurat can only read the input files in the format generated by Cell Ranger (10x genomics). However, it is possible to convert your counts' matrix to the acceptable format. For example, the function write10xCounts(), from the DropletUtils package, is an easy option to make this conversion.

Tip: Using write10xCounts(), users can provide as output the path to the data/ directory. In this way, Asc-Seurat can recognize the files automatically.

10.2 Location of the dataset

For the integration of multiple samples, the process is a little different. Users still need to add their datasets in the data/ directory, creating a subdirectory for each sample. However, users also need to provide a configuration file containing the parameter values for each sample. During the installation, an example file named *configuration_file_for_integration_analysis.csv* will be created in the directory and can be used as a model.

Note: The integration of samples can be biased if the parameters are not chosen appropriately. Therefore, it is recommended to explore each sample separately in the tab *One sample*, defining adequate parameters to remove deficient quality cells before the integration.

The user's configuration file must have six columns and a header (the column names are not restricted). They specify what cells should be kept for each sample while loading the data before the integration.

Also, the columns need to be in a specific order, as listed below.

1. **Subdirectory name**: The name of the subdirectories containing the datasets. Each sample must have a unique name for its subdirectory, even if they are replicates.

- 2. Sample name (any name you prefer): Your choice of name for each sample. If you have replicates and want them to be considered as one in the plots and analysis, use the same name for all replicates.
- 3. Min. number of cells expressing a gene: Include genes only if they are detected in at least this many cells.
- 4. Min. number of genes a cell must express to be included: Include cells only if they expressed at least this number of genes.
- 5. Maximum number of genes a cell can express and still be included: Remove cells that express more than this number of genes. Useful to remove cells that are suspected to be doublets.
- 6. Maximum percentage of genes belonging to the mitochondrial genome: Here, the regular expression (Regex) is a sequence of characters that is used to identify the genes belonging to the mitochondrial genome. For example, when using the human genome, this sequence should be "^MT-".

10.3 Loading the data and performing integration

To demonstrate the necessary steps to load and integrate multiple datasets using Asc-Seurat, we used two groups of cells from Kang et al., 2017, that are also used in Seurat's tutorial demonstrating the comparison of multiple samples. Two datasets are used, both containing peripheral blood mononuclear cells (PBMCs). However, the first dataset contains the cells of the control group (Control), while the second dataset contains cells treated with interferon-beta (Treatment).

The first step is two create two folders inside the data/ folder. The folders were named example_PBMC_control and example_PBMC_treatment, each containing the three necessary input files (shown in the image below).

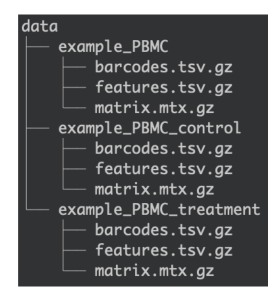


Fig. 1: Organization of the data/ folder the different datasets.

After that, it is necessary to create a configuration file in the csv format. During the installation, an example file named *configuration_file_for_integration_analysis.csv* is created in the directory. It can then be used as a model. For this example, the configuration file contains the information shown below.

	sumpres.				
Subdirectory	Sample name	Min. num-	Min. number	Max. num-	Max. per-
name (must be	(any name you	ber of cells	of genes a cell	ber of genes	centage of
inside data/)	prefer)	expressing a	must express	a cell can ex-	transcripts
		gene	to be included	press and still	belonging to
				be included	mitochondrial
					genome
example_PBMC_	contornation	3	250	2500	5
example_PBMC_	træatentent	3	250	2500	5

Table 1: Example of a configuration file for the integration of multiple samples.

Once the configuration file is ready, users only need to load it in the app and select the samples they want to integrate (see image below). Also, it is necessary to choose the normalization method, the regex string to detect mitochondrial transcripts, the number of Principal Components to be used during the integration (see below). Note that while default values are provided, users need to set these parameters based on their evaluation of the individual samples that are being integrated.

Run a new integration analysis or read a previously saved file?	Common identifier of mitochondrial genes	Select the normalization methodLogNormalize
 Run a new analysis Load file 	МТ	O SCTransform
Read the configuration file containing the samples' information	N of variable genes for integration	Scale factor
Browse confi_file_integration.et	2000 💿	Select the method to detect the most
Upload complete	N of components for the integration	variable genes vst
Select the samples to use	20 🔹	mean.var.plot (mvp)
example_PBMC_control, example		O dispersion (disp)
	Project name	
Load the integrated data or execute a new integration		

Fig. 2: Loading configuration file and defining parameters for the integration of multiple samples using LogNormalization.

10.4 Saving integrated data for reanalysis

The integration of multiple samples is a timing-consuming step of the analysis. The amount of time necessary to execute this step depends on the number of datasets and the number of cells in each dataset, and it can take several minutes to be concluded.

Therefore, Asc-Seurat allows users to save the integrated data and skip the integration step the next time users need to use the same dataset. To save the data, users can click on the button Download RDS object containing the integrated data. and save the rds file inside the RDS_files/folder.

Next time this data is necessary, users can select the option "Load file" and skip the integration step, as shown below.

Run a new integration analysis or read a previously saved file?Run a new analysisLoad file	Select the file containing the integrated data pbmc_integrated.rds	Load the integrated data or execute a new integration
	Inform the normalization method used to generate the integrated dataset	

ELEVEN

QUALITY CONTROL

After integrating the datasets, a violin plot will be generated showing the distribution of cells according to three parameters:

- nFeature_RNA: the number of genes detected in each cell
- nCount_RNA: the number of molecules detected per cell
- percent.mt: the percentage of transcripts that map to mitochondrial genes

After visualizing the distribution of cells, it is possible to set more restrictive parameters (on the right side of the plot) and filter cells based on the number of expressed genes per cell and the percentage of transcripts from mitochondrial genes. By clicking on *Show plot of filtered data*, users can see the distribution of cells after filtering and then readjust the parameters. The figure below shows the distribution of cells of the PBMC integrated (containing the Control and Treatment datasets, see *Loading the data and integration of multiple samples*) dataset before and after filtering.

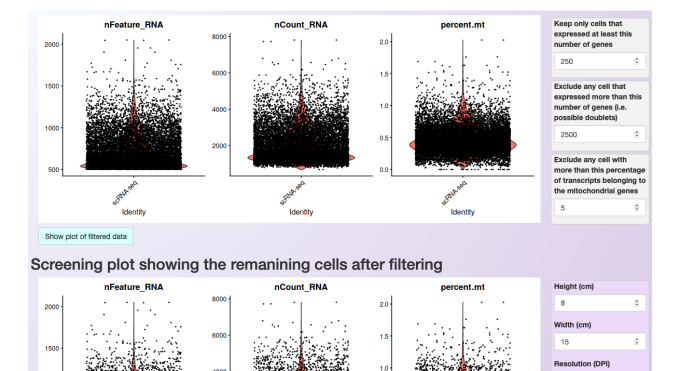
Asc-Seurat allows users to download each of the plots with high-resolution by clicking on the Download plot button.

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se RNA-sea

Identity



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Identity

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File type

png

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TWELVE

CLUSTERING

12.1 Normalization

When integrating multiple samples, the normalization is executing during the integration.

Note: Currently, the recommendation of Seurat's team is to use the standard "RNA" assay when performing differential expression (D.E) analysis and for data visualization, even when using SCTransform (See here). Therefore, Asc-Seurat will use the SCTransformed data ("SCT" assay) until the clustering step only.

To use the "RNA" assay after SCTransform, Asc-Seurat will automatically perform the LogNormalization and scaling of the data in the RNA assay by applying the default parameters.

12.2 Dimensional reduction (PCA)

The PCA will be executed using Seurat's function RunPCA and, after its conclusion, an elbow plot is generated automatically, to help users to decide how many PCs should be included to inform the clustering step.

Users can use this plot to select the PCs with the highest standard deviation (more informative PCs). Also, users should set the number of PCs to include during clustering in the windows at the plot's right side.

In the example below, the first 20 PCs are selected. Not that the resulting plot will be slightly different depending on the normalization method. Below we show the result obtained using LogNormalization.

12.3 Clustering of cells

The next step is the clustering of the cells. For that, Asc-Seurat used both FindNeighbors and FindClusters functions of the Seurat package.

Before the execution, however, users need to set a value for the resolution parameter. The resolution is an important parameter to evaluate because it determines the profile and number of clusters identified for a dataset. Selecting larger values will favor splitting cells into more clusters while choosing a smaller value has the opposite effect. Quoting from Seurat's tutorial: "We find that setting this parameter between 0.6-1.2 typically returns good results for single-cell datasets of around 3K cells. Optimal resolution often increases for larger datasets".

Tip: There is no easy way to define an optimal value for the resolution parameter. Users need to try different values and evaluate the resulting clusters according to the expectation for their cells population. Visualizing the expression profile of cell-type-specific markers can provide a hint if the chosen value is too small or too large.



Fig. 1: Elbow plot provided to help to select the most informative PCs. For the PBMC integrated dataset, and using the LogNormalization method, we chose the 20 first PCs.

After the clustering step's execution, three plots are generated for cluster visualization, all of them using the Uniform Manifold Approximation and Projection (UMAP) technique. The first plot shows the clustering of the whole dataset colored by cluster. The second plot shows the same plot, but cells are colored by sample. The third plot shows the clustering of the cells of each sample, with one subplot per sample.

12.3.1 Selecting clusters of interest

In some cases, it is interesting to select or exclude some clusters of cells from the dataset before executing the subsequent steps. This process is helpful, for example, when users desire to explore a developmental trajectory of a specific group of cell types.

Asc-Seurat makes this step simple. Users only need to select the cluster(s) to keep or exclude and start reanalysis of the remaining cells by clicking on *Reanalyze after selection/exclusion of clusters* (see below).

Asc-Seurat will then execute the steps with the new set of cells up to the PCA. Then, **users need to evaluate the elbow plot and decide the number of PCs to cluster the new set of cells**. Users can either keep the same value for the resolution parameter or modify it before clicking on *Run the clustering analysis* to start the clustering once more.

Warning: The cluster's numbering will change every time that clusters are selected or excluded.

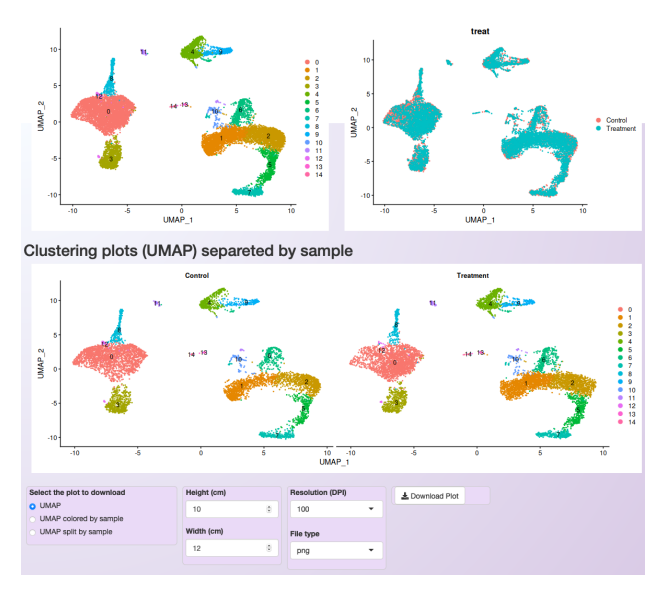
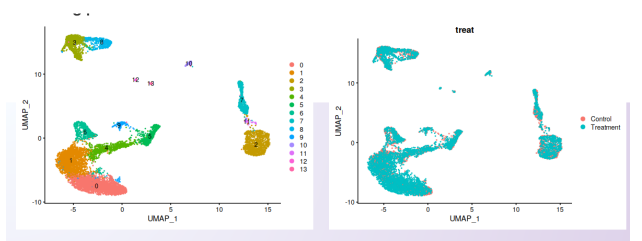


Fig. 2: Plot showing the PBMC integrated dataset clustering using 20 PCs, LogNormalization, and a resolution value of 0.5.

Do you want to select or exclude clusters of cells and reanalyze the data?	Do you want to select or exclude the clusters?	Choose clusters to select or exclude	Reanalyze after selection/exclusion of clusters	
• Yes	Select	•		
0 No	• Exclude			

Fig. 3: Asc-Seurat makes it easy to select or exclude a cluster (or clusters) of cells. In this example, we exclude all cells belonging to cluster 0.



Clustering plots (UMAP) separeted by sample

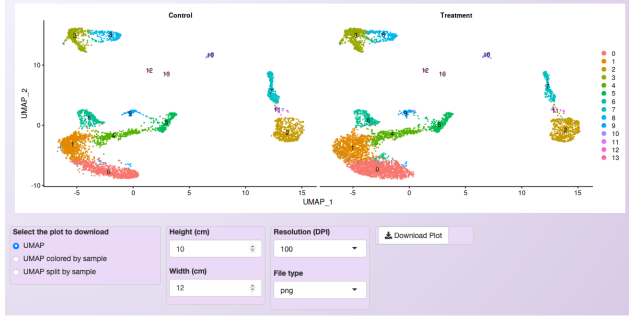


Fig. 4: Clustering of the PBMC integrated dataset after excluding cells belonging to cluster 0 from the original dataset.

THIRTEEN

MARKERS IDENTIFICATION AND DIFFERENTIAL EXPRESSION ANALYSIS

After clustering the cells, users may be interested in identifying genes specifically expressed in one cluster (markers) or in genes that are differentially expressed among clusters of interest. Asc-Seurat can apply multiple algorithms to identify gene markers for individual clusters or identify differentially expressed genes (DEGs) among clusters. Moreover, when using an integrated dataset containing multiple samples, it is possible to identify DEGs among samples for each cluster.

Note: When searching for markers of a cluster or DEGs among clusters using an integrated dataset, the search will attempt to find markers or DEGs conserved among samples.

Asc-Seurat allows users to filter gene markers and DEGs by the fold change and minimal percentage of cells expressing a gene in the cluster(s). Moreover, users can define a significance level to exclude genes based on the adjusted p-value (see below).

Detect conserved markers or D.E. genes	Select the analysis to perform?	Search for markers/D.E. genes
 Conserved markers 	 Identify conserved markers for all clusters 	
O D.E. genes between treat vs control	 Identify conserved markers for one specific cluster Identify conserved markers for one specific cluster in comparison with another cluster(s) 	

Fig. 1: Example of Asc-Seurat's interface showing the settings to the search for gene markers for each of the clusters and conserved among samples.

Detect conserved markers or D.E. genes	Select the analysis to perform?	Select the cluster of interest	Search for markers/D.E. genes
Conserved markers D.E. genes between treat vs control	 Identify conserved markers for all clusters Identify conserved markers for one specific cluster Identify conserved markers for one specific cluster in comparison with another cluster(s) 	0 •	

Fig. 2: Example of Asc-Seurat's interface showing the settings to the search for gene markers for a specific cluster and conserved among samples.

An iterative table will be available after executing the search for marker or DEGs, showing the significant genes. Moreover, users can download the list of significant markers or DEGs as a csv file.

The list of genes in the csv can then be used to visualize their gene expression in a series of plots, as shown in the section *Expression visualization*.

Detect conserved markers or D.E. genes	Select the analysis to perform?	Select the cluster of interest	Search for markers/D.E. genes
Conserved markers	O Identify conserved markers for all clusters	0 •	
D.E. genes between treat vs control	 Identify conserved markers for one specific cluster 	Select the cluster(s) to compare	
	 Identify conserved markers for one specific cluster in comparison with another cluster(s) 	1	

Fig. 3: Example of Asc-Seurat's interface showing the settings to search for DEGs genes among clusters 0 and 1.

Detect conserved markers or D.E.	Select the cluster of interest		Search for markers/D.E. genes
Conserved markers	0	•	
D.E. genes between samples	Select the sample/treatment of interest		
	Treatment	•	
	Select the sample/treatment of interest		
	Control	•	
	Select the statistical test		
	wilcox	•	
	Select the (adjusted) p-value threshold.		
	0.05	0	

Fig. 4: Example of Asc-Seurat's interface showing the settings to search for DEGs among samples for a specific cluster (cluster 0).

									Search	
geneID	cluster	Control_p_v al	Control_avg _log2FC	Control_pct. 1	Control_pct. 2	Control_p_v al_adj	Treatment_p _val	Treatment_a vg_log2FC	Treatment_p ct.1	Treatme ct.2
IGJ	4	0	1.801337491 35422	0.982	0.084	0	4.198947474 58671e-79	2.013165306 90708	0.128	0.01
MS4A1	4	7.144032132 85382e-263	3.021162764 87278	0.952	0.182	1.428806426 57076e-259	0	2.585376416 93091	0.474	0.01
CD79A	4	7.231286817 96333e-159	3.474233303 99099	0.891	0.35	1.446257363 59267e-155	0	3.321458944 53177	0.697	0.02
BANK1	4	3.057718755 06545e-267	1.303219199 51744	0.846	0.07	6.115437510 13089e-264	5.108717510 11766e-253	1.605406412 15536	0.25	0.00
TSPAN13	4	3.065264775 216e-250	0.280878039 603956	0.967	0.224	6.130529550 43201e-247	2.421113352 80203e-41	0.647817091 230995	0.121	0.02
CD74	4	1.170922211 71069e-172	1.962649891 25589	1	0.896	2.341844423 42139e-169	2.674951064 02733e-249	2.076550533 11759	0.995	0.66
NFRSF13B	4	7.826048436 39417e-246	1.443234912 69393	0.778	0.033	1.565209687 27883e-242	6.578718512 8867e-231	1.458355632 9649	0.194	0.00
FCRLA	4	7.021231392 50677e-241	0.973153972 557517	0.78	0.041	1.404246278 50135e-237	1.479959033 03173e-110	0.817668842 834523	0.107	0.00
ANXA1	4	1.370032254 23451e-191	-3.85455042 80957	0.722	0.987	2.740064508 46902e-188	7.831585073 39196e-201	-3.52280676 614149	0.096	0.81
APOBEC3B	4	1.714266721 63984e-197	-1.48695608 468927	0.197	0.951	3.428533443 27969e-194	8.244455115 88342e-15	-4.00998820 461381	0.027	0.1

Fig. 5: The ten most significant markers identified for cluster 4 of the PBMC integrated dataset (the clustering is shown in *Clustering*).

FOURTEEN

EXPRESSION VISUALIZATION

Asc-Seurat provides a variety of plots for gene expression visualization of the integrated data. From a list of selected genes, it is possible to visualize the average of each gene expression in each cluster in a heatmap. It also provides plots for the visualization of gene expression at the cell level. Moreover, violin plots and dot plots allow the visualization of each cluster's expression, emphasizing the inter-cluster comparison.

For the integrated dataset, besides identifying markers for each cluster and DEGs among clusters, it is also possible to identify DEGs among samples (See *Markers identification and differential expression analysis*). Below are shown examples of plots that Asc-Seurat generates to allow the expression visualization in all these cases.

14.1 Expression visualization of genes identified as markers

14.1.1 Format of the input file containing genes for expression visualization

Asc-Seurat expects as input a csv (comma-separated value) or a tsv (tab-separated value) file containing at least two columns. The first column must contain the gene ID as present in the dataset, and the second column is a grouping variable. An optional third column can contain the common names of each gene. Any additional column will be ignored. The output files generated by the differential expression analysis are already in the correct format to be used as input for the visualization.

Below is shown an example of an input file used for expression visualization. It contains ten markers identified for cluster 4 of the PBMC integrated dataset (Control and Treatment). In this case, the dataset uses the gene name as an identifier, and this is the information contained in the first column. The second column is used to group de marker according to their clusters.

ing the required	• or united	
MS4A1	Cluster_4	
CD79B	Cluster_4	
CD79A	Cluster_4	
BANK1	Cluster_4	
CD74	Cluster_4	
TNFRSF13B	Cluster_4	
ANXA1	Cluster_4	
KIAA0226L	Cluster_4	
BLNK	Cluster_4	
C7orf50	Cluster_4	

Table 1: Example of an input file for gene expression visualization showing the required columns.

After loading the input file, users can select what group(s) of genes to explore and choose specific genes from each group. Moreover, if a third column is provided in the input file, users can use the genes' common name instead of the

gene IDs to select the genes to be shown.

14.1.2 Heatmap

Once users selected their genes of interest, they can generate a heatmap of the average of each gene expression in each cluster by clicking on the button *Show heatmap with the average of expression per cluster*. The heatmap will adjust its height according to the number of selected genes. Moreover, rows and columns will be organized by a hierarchical clustering algorithm. A high-resolution copy of the heatmap plot can be download in a diversity of formats.

Warning: For the integrated dataset, the heatmap shows the average expression of all samples together. It is only helpful to identify if the cell types' markers make sense with the number of generated clusters.

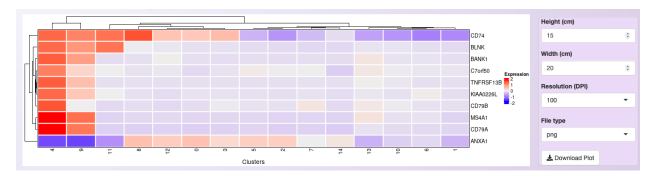


Fig. 1: Heatmap showing the expression profile of the then most significant markers for cluster 4 of the integrated datasets.

14.1.3 Gene expression at the cell level - Feature plots

From the list of genes on the heatmap, users can select genes to further explore by visualizing the expression at the cell level. For each selected gene, a feature plot showing each sample's profile will be generated using Seurat's Feature plots function. The UMAP plot is shown side-by-side with the feature plots, so users can quickly compare the expression profile with the identified clusters.

14.1.4 Visualization of the expression among clusters

For each selected gene, Asc-Seurat will also generate plots to visualize the distribution of cells within each cluster according to the expression of the gene (violin plot) and the percentage of cells in each cluster expressing the gene (dot plot) in each sample. Seurat's functions VlnPlot() and DotPlot() are deployed in this step.

14.2 Expression visualization of differentially expressed genes

14.2.1 Format of the input file containing genes for expression visualization

As before, a csv or tsv file is necessary as input for the expression visualization of DEGs. In this case, it contains ten genes identified as DEGs between the PBMC Treatment and PBMC Control datasets in cluster 4.

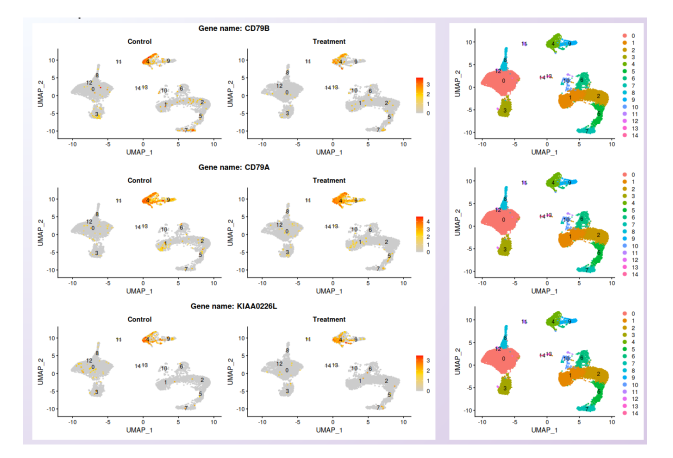


Fig. 2: Visualization of the expression profile of three of the genes shown on the heatmap in each sample.

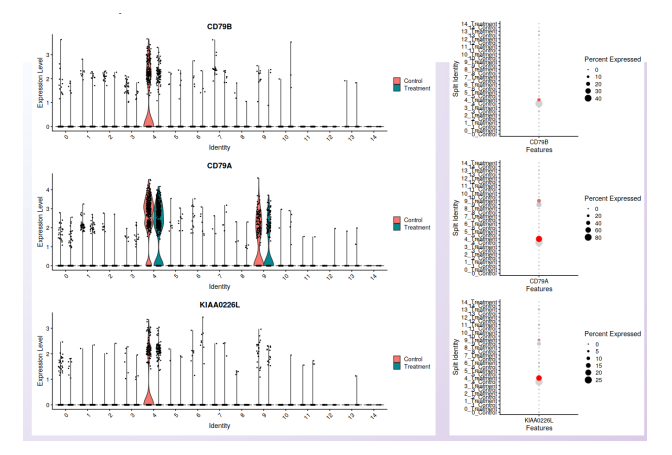


Fig. 3: Visualization of cells' distribution within each cluster according to the gene expression (violin plot; left) and the percentage of cells in each cluster expressing the gene (dot plot; right) in each sample. The three genes shown are the same used for the feature plots.

2200		
ISG15	DEGs Cluster_4	
IFIT3	DEGs Cluster_4	
IFI6	DEGs Cluster_4	
ISG20	DEGs Cluster_4	
IFIT1	DEGs Cluster_4	
MX1	DEGs Cluster_4	
LY6E	DEGs Cluster_4	
TNFSF10	DEGs Cluster_4	
IFIT2	DEGs Cluster_4	
B2M	DEGs Cluster_4	

Table 2: Example of an input file for gene expression visualization of DEGs.

14.2.2 Heatmap

All ten genes were selected for visualization in the heatmap. Once more, it is important to mention that the heatmap shows the average expression among all samples. However, by investigating the heatmap below, it is possible to notice that while these genes are the most significant DEGs between samples in cluster 4, they are widely expressed in other clusters too.

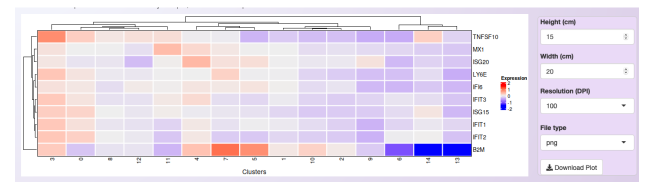


Fig. 4: Heatmap showing the expression profile of the ten most significant DEGs between Treatment and Control in cluster 4 of the integrated datasets.

14.2.3 Gene expression at the cell level - Feature plots

To compare the expression profile among samples, the visualization at the cell level is more relevant, as shown below.

From the list of genes contained on the heatmap, three genes were selected. While the expression is not localized in cluster 4, it is clear the increment of the expression in the Treatment dataset. The UMAP plot is shown side-by-side with the feature plots, allowing comparing the expression profile with the identified clusters.

14.2.4 Visualization of the expression among clusters

As in the feature plot, the violin and dot plots clearly show the increased level of expression in the cells of the PBMC Treatment sample compared to the PBMC control.

Tip: Sometimes, it is necessary to make fine adjustments to an image before publication. Saving the plots as a

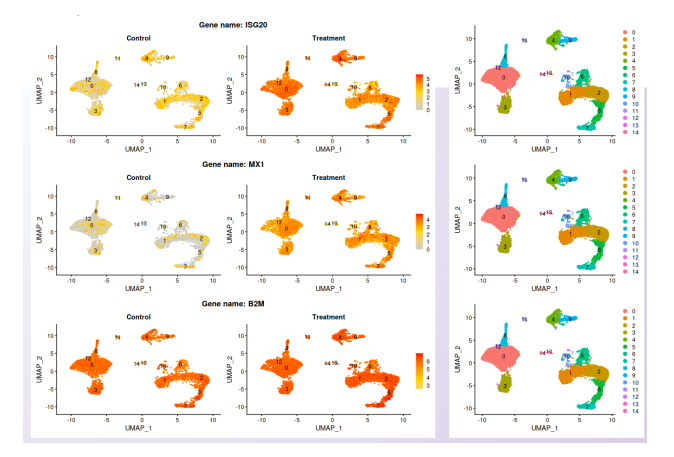


Fig. 5: Visualization of the expression profile of three of the genes shown on the heatmap in each sample.

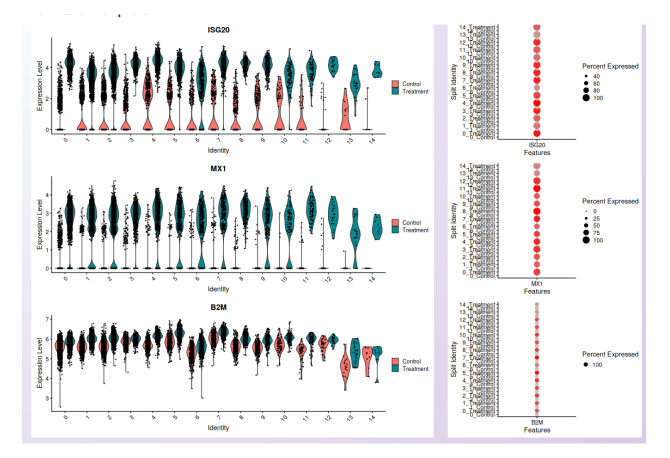


Fig. 6: Visualization of cells' distribution within each cluster according to the gene expression (violin plot; left) and the percentage of cells in each cluster expressing the gene (dot plot; right) in each sample. The three genes shown are the same used for the feature plots.

Scalable Vector Graphic (svg), allows the edition of all aspects of the plot by image edition software as Inkscape (free).

FIFTEEN

TRAJECTORY INFERENCE

For the trajectory inference analysis, users can either execute it through capabilities of the embedded slingshot (Bioconductor) package or select another model contained in dynverse, executed using a docker image provided by dynverse. In both options, users only need to choose the model and initial parameters (see below). However, the direct execution of slingshot is faster than executing models via dynverse's docker image.

To inform the model's choice, it is recommended the reading of Saelens et al., 2019 paper, that benchmarked a diversity of models. Different models will perform better or worst depending on the topology of the developmental trajectory of the dataset (that is unknown a priory). Therefore, users need to consider what topology they may expect for their dataset. For example, slingshot performs well for bifurcated, multifurcated, or "tree" like topologies, but not for cyclic or more complex disconnected trajectories.

Warning: Some of the models included in dynverse are computationally intensive. It is strongly recommended to check the requirements for a model before executing it on Asc-Seurat. You can use dynguidelines web application to investigate the necessary resources to analyze your dataset. The amount of resources also depends on the number of cells and the complexity of the dataset.

15.1 Executing the trajectory inference and trajectories visualization

To start the trajectory inference analysis, users need to save the clustered data in a specific folder automatically created during the installation (RDS_files/). Asc-Seurat recognizes the data automatically, and users can select the sample to be used. Next, users need to select the model to be used, inform if the data is composed of one or multiple integrated samples, and, optionally, inform the cluster(s) expected to be at the beginning and/or end of the inferred trajectory. After executing the analysis, three plots showing different inferred trajectory representations are generated. Moreover, when using an integrated dataset, users can also color the cells according to the sample of origin. To demonstrate these capabilities, we used the PBMC integrated dataset (containing two samples: Control and Treatment).

Note: The time to execute the trajectory varies from minutes to hours, depending on the complexity of the dataset and the chosen model. Visit dynguidelines web application for an estimative.

For the PBMC integrated dataset, slingshot was used to infer the developmental trajectory. Note that no cluster was select as the start or end of the trajectory, so slingshot makes this decision. If you know what cluster (or cell type) is expected at the beginning or end of the trajectory, providing this information will allow a better interpretation of the generated trajectory.

When users inform that multiple samples are used, Asc-Seurat offers coloring the cells by cluster identify or by samples. Both options are demonstrated below for the PBMC integrated dataset.

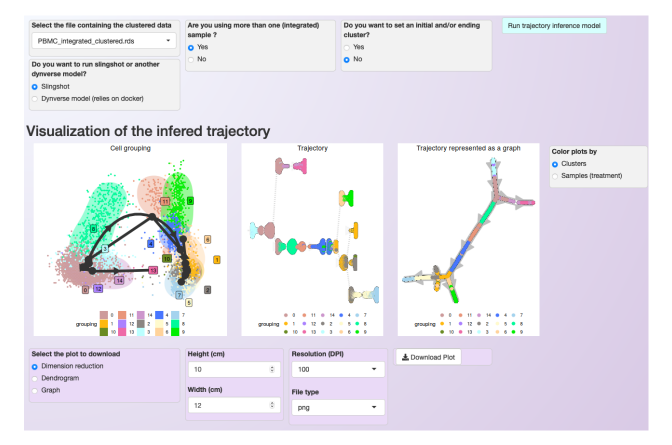


Fig. 1: Asc-Seurat provides multiple models for trajectory inference analysis and three options for trajectory visualization. In this case, cells are colored by clusters.

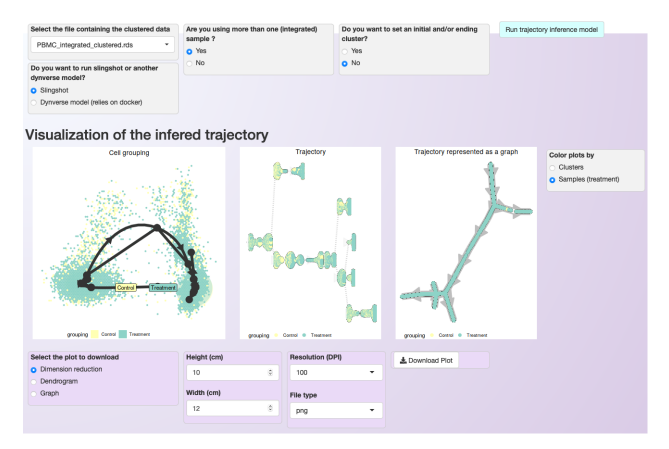


Fig. 2: Asc-Seurat provides multiple models for trajectory inference analysis and three options for trajectory visualization. In this case, cells are colored by sample.

Tip: Suppose you are interested in studying the developmental trajectory of a subgroup of clusters only. In that case, it is better to exclude the other clusters than to try to infer the trajectory using the whole dataset. The model will execute quicker and provide a better resolution of the trajectory since the complexity of the dataset is reduced. Asc-Seurat allows the exclusion of clusters from your dataset, see *Selecting clusters of interest* (one sample) or *Selecting clusters of interest* (integrated dataset).

15.2 Expression visualization within the trajectory and identification of DEGs in the trajectory

After inferring the developmental trajectory, it is possible to visualize the expression of genes of interest in the cells within the trajectory. Asc-Seurat provides two options for this visualization, 1) a heatmap displaying the expression of genes in each cell, ordered by the cell position within the trajectory, and 2) the visualization of the same three trajectory's representation shown above but colored by the gene expression.

Users can either load their list of genes of interest or identify DEGs within the trajectory for the visualization.

15.2.1 Visualizing the expression of a list of selected genes

To visualize the expression of specific genes, the process is similar to the described on *Expression visualization*. Asc-Seurat expects as input a csv (comma-separated value) file containing at least two columns. The first column must contain the gene ID as present in your dataset, and the second column is a grouping variable. An optional third column can contain the common names of each gene. Any additional column will be ignored. **No header is allowed for this file**.

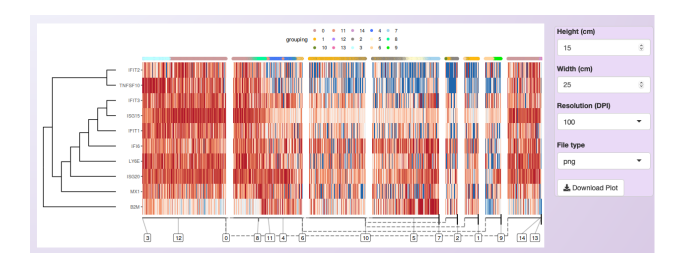
After loading the input file, users can then select what group(s) of genes to explore, as well as select specific genes from each group. Moreover, if a third column is provided in the input file, users can use the common name of the genes instead of the gene IDs to select the genes to be shown.

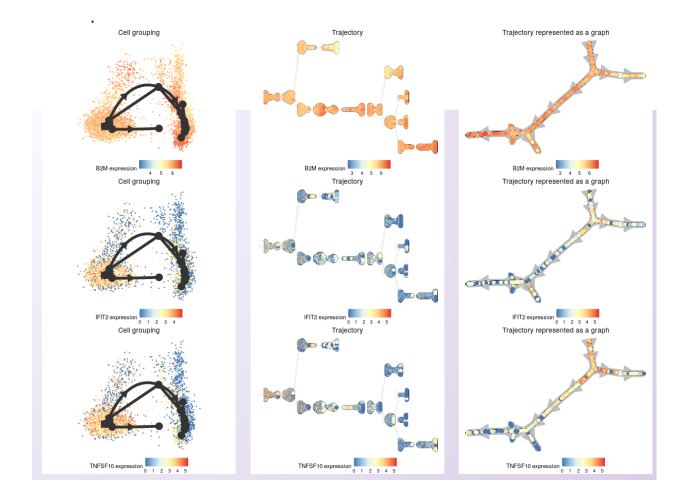
Input the list of markers	Select the group of markers to test	Select the genes to show	Select the expression values to show	Show heatmap
Browse PBMC_cluesters_mark	Cluster_2, Cluster_3 -	CD3D, CD79A, CD79B, IL32, IL7R, •	 counts normalized 	
Does your file have a header?	 Visualization options Show all genes Select genes to show 		 normalized and scaled 	
Load markers	ID options Use IDs Use Name			

After choosing the genes, a heatmap showing the expression in the cells sorted by their position in the inferred trajectory is shown. Then, users can select genes for individual visualization.

As an example, it is shown the expression of the same ten DEGs identified for cluster 4 in the comparison of Control and Treatment for the PBMC integrated dataset (see *Markers identification and differential expression analysis*).

Next, three of those genes were selected to show the expression on the cells in the inferred trajectory.





15.2.2 Identification of DEGs in the trajectory

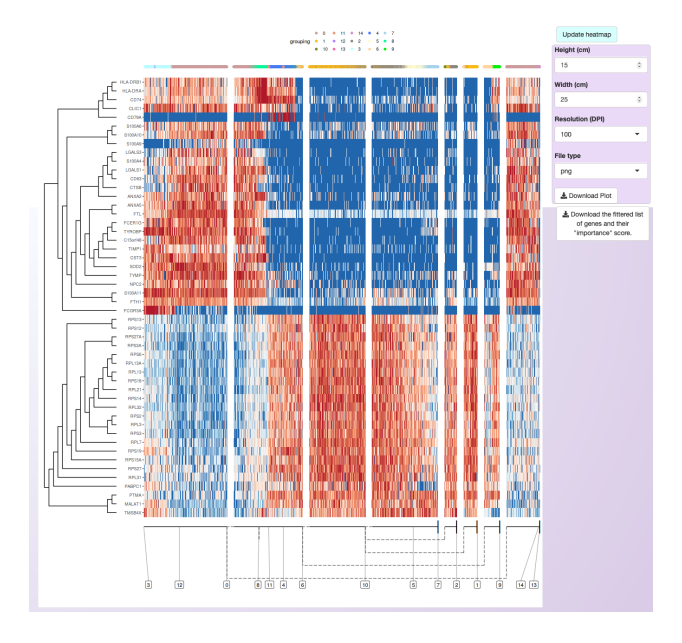
To identify differentially expressed genes, Asc-Seurat deploys the dynfeature, part of dynverse's collection of packages. Here we provide a short introduction to these methods. Please, visit dynverse's Trajectory differentially expression page for a demonstration of each method.

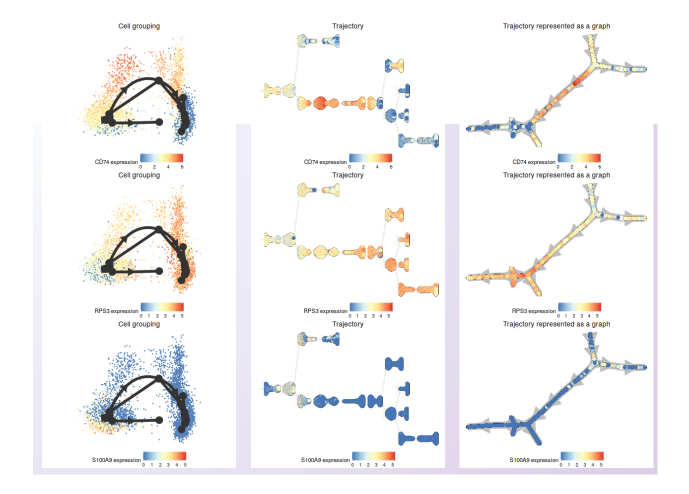
Asc-Seurat allows the search for DEGs within the whole trajectory, in a branch of the trajectory between two clusters or in a branching point. Each of these methods will rank all genes of the dataset. Therefore, users need to select the number of genes (ranked by the most important genes) to visualize in the heatmap. Also, users can download the list of all genes and their "importance values".

Select the type of analysis: Vizualization of a list of genes		
 Idenfitication of important genes using Dynfeature 		
Expression of markers		
For each of the below analyzes, it is possible to s	save the list of all expressed genes	with their "importance" values. It is also possible to save the list of the top N genes.
For the branch/linage analysis, you can provide t is blank	he beginning (from) and end (to) of	he branch/linage of interested. You can also select only one, either from or to, as long as the other box
For the bifurcation analysis, you can provide the the "importance" values for each branching point		ching happened to select the genes more relevant to this point. If left in blank, the analysis will identify
Select the analysis to perform	Defines the most important genes	
global overview of the most predictive genes Lineage/branch markers Genes important at bifurcation points	Download the list of all important genes in all branches and their "importance" score.	
Set the number of genes to be used in the heatmap (top relevant genes)		
50		

As an example, for the PBMC integrated dataset, we opted to show the 50 most significant DEGs within the trajectory, as ranked by their "importance" value on explaining the inferred trajectory.

From those, a few genes were selected for expression visualization on the trajectory.





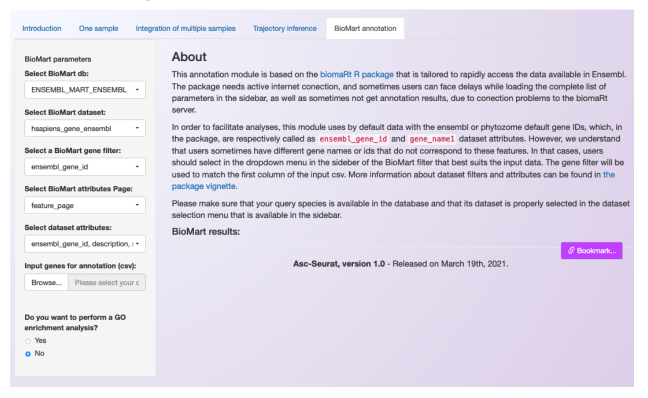
SIXTEEN

BIOMART ANNOTATION

The annotation module of Asc-Seurat is based on the **biomaRt** package (Bioconductor). BiomaRt is designed to facilitate the functional annotation of genes available for various species through the **BioMart** databases. To date, the primary databases in BioMart are the ones provided by Ensembl. Fortunately, biomaRt provides direct access to these datasets, and they can all be accessed via Asc-Seurat. Moreover, due to its importance for plant species, we also incorporated access to the Phytozome's BioMart database.

16.1 Functional annotation of genes

The annotation module of Asc-Seurat was designed to be simple to use (See image below). Nonetheless, a basic understanding of how BioMart queries are built is required so that users can select the filters and attributes needed. Please, visit biomaRt's vignettes for an overview.



As shown in the image above, Asc-Seurat contains a sidebar on which users can select the best parameters for annotating their genes. Initially, users should select the database to use (Phytozome or one of Ensembl's databases). Then, Asc-Seurat will load it and display the datasets (species) available for the selected database. After selecting the species' dataset to use, users can define the filter and attributes of the query. In summary, the filter corresponds to the dataset being used as input and, for most cases, will be the gene IDs or the gene names. The attributes are the information users want to extract from the database, e.g., description of the gene function, Gene Ontology (GO) terms, Pfam domains, etc. Please check this section of biomaRt's vignettes for an example.

After defining the filter and the attributes, users can provide a csv file containing a list of gene ids (or gene names) and start the query. Moreover, users can select only a subset of the genes listed in the csv file, reducing the time necessary for the annotation.

Note: The input csv file should contain one or more columns, separated by commas. A header is required, but users are free to use their choice of column(s) name(s). The only required information is the gene ids, or gene names, one entrance per line. Asc-Seurat will ignore other columns that might be present. The csv files generated within Asc-Seurat are adequate as input for the annotation.

To execute the annotation, users need to click on *Annotate selected genes!*. An iterative table containing the requested information will be generated. Also, users can download the list of annotated genes as a csv or an Excel file (see below).

Introduction One sample Integration of multiple samples	Trajectory inference BioMart annotation		
BioMart parameters Select BioMart db:	About This annotation module is based on the biomaRt R package that is tailored to rapidly access the data available in Ensembl. The package needs active internet	t conection, and som	netimes users can
ENSEMBL_MART_ENSEMBL -	face delays while loading the complete list of parameters in the sidebar, as well as sometimes not get annotation results, due to conection problems to the bio		
Select BioMart dataset:	In order to facilitate analyses, this module uses by default data with the ensembl or phytozome default gene IDs, which, in the package, are respectively callect gene_name1 dataset attributes. However, we understand that users sometimes have different gene names or ids that do not correspond to these features. In		
hsapiens_gene_ensembl •	dropdown menu in the sideber of the BioMart filter that best suits the input data. The gene filter will be used to match the first column of the input csv. More in attributes can be found in the package vignette.	nformation about dat	aset filters and
Select a BioMart gene filter:	Please make sure that your query species is available in the database and that its dataset is properly selected in the dataset selection menu that is available in	n the sidebar	
ensembl_gene_id -	BioMart results:	rino oldobali	
Select BioMart attributes Page:	Annotate selected genesl		
feature_page •	Copy CSV Excel Show 5 6 entries		
Select dataset attributes:	ensembl_gene_id description	start_position	end_position
ensembl_gene_id, description, start_position, end_position -	ENSG00000007312 CD79b molecule [Source:HGNC Symbol;Acc:HGNC:1699]	63928740	63932336
Input genes for annotation (csv):	ENSG0000008517 interleukin 32 [Source:HGNC Symbol;Acc:HGNC:16830]	3065297	3082192
Browse genes_IDS_top50.csv Upload complete	ENSG00000011600 transmembrane immune signaling adaptor TYROBP [Source:HGNC Symbol;Acc:HGNC:12449]	35904401	35908295
Select genes to annotate:	ENSG00000019582 CD74 molecule [Source:HGNC Symbol;Acc:HGNC:1697]	150400041	150412969
Select genes to annotate.		88643289	88651054
ENSG00000235915, ENSG00000230791, ENSG0000023551 +	ENSG00000051523 cytochrome b-245 alpha chain [Source:HGNC Symbol;Acc:HGNC:2577]		
ENSG00000235915, ENSG00000230791, ENSG0000023551 - Do you want to perform a GO enrichment analysis?		2 3 4 5	20 Next

16.2 GO terms enrichment analysis

Asc-Seurat also provides an option to execute the GO terms enrichment analysis using topGO, a Bioconductor package.

This analysis aims to identify genes over/under-represented in the set of genes being annotated (known as target) compared to a broader set of genes (known as the universe). The universe can be a set of all genes expressed in the dataset or any set of genes that users desire to compare with the set of genes being annotated.

If users choose to execute this analysis, they need to provide a second csv file containing the list of genes to be used as the "universe" of the analysis.

Warning: Both sets of genes should contain the same type of identifier (i.e., gene ID). Also, be aware of extra spaces or any discrepancy between the two sets of genes' IDs.

At the end of the GO enrichment analysis, an iterative table containing all enriched GO terms is generated, which can be downloaded in the csv format or as an Excel file. Moreover, a plot showing the most significant GO terms is

generated. Users can adjust the number of significant GO terms shown for each GO category in the plot (see below for an example using 5 GO terms per category).

Run GO enrichment analysis! Copy CSV Excel Show 5 enrichment	tries							
GO.ID ¢ Term ¢	Annotated 🔶	Significant 🕴	Expected 🕴	Rank in classicFisher	classicFisher	KS 🛉	weightFisher	Macro
GO:0006955 immune response	34	1	0.08	8	0.0778	0.0000038	0.00308	Biological Process
GO:0002376 immune system process	39	1	0.09	9	0.0888	0.000032	1	Biological Process
GO:0000151 ubiquitin ligase complex	10	0	0.24	9	1.000	0.0085	1	Cellular Componen
GO:0008270 zinc ion binding	246	2	2.2	33	0.64813	0.065	0.00194	Molecular Function
GO:0000145 exocyst	1	0	0.02	8	1.000	0.1652	1	Cellular Componen
Showing 1 to 5 of 30 entries						Previous 1	2 3 4	5 6 Nex
GO En	richment Analy	sis						
immune response immune system process lipid metabolic process biological_process proteolysis				 			•	Biological Process
ubiquitin ligase complex - exocyst - integral component of membrane - intrinsic component of membrane -	8							cellular Componer
zinc ion binding structural constituent of ribosome actin binding lysozyme activity calcium ion binding	•••			 				Note cular Function
0				2 Enrichment score -log lines drawn at equivalents of		4		
itatistical test to plot	Width (cm)			Resolution (DPI)		± D	ownload Plot	
Kolmogorov-Smirnov 🔻	10			300		-		
lot (N) top GOs per category	Height (cm)			File type				
5	10			svg		•		

CHAPTER SEVENTEEN

ADVANCED PLOTS

As shown in the sections describing the expression visualization tools (*here* and *here*), Asc-Seurat provides a diversity of plots to explore your dataset. However, it focuses on exploring each gene individually, not providing tools to visualize the expression of multiple genes at once.

Starting on v2.0, Asc-Seurat also provides the capacity of generating dot plots and "stacked violin plots" comparing multiple genes.

Using an rds file containing the clustered data as input, users must provide a csv or tsv file in the same format described in the *expression visualization* section. Next, using the grouping variable, column two of the csv (or tsv), select the sets of genes to be used in the plot. Both violing and dot plot will be generated.

17.1 Stacked Violin plot

Stacked violin plots are a popular way to represent the expression of gene markers but are not provided by Seurat. Asc-Seurat's version of the stacked violin plot is built by adapting the code initially posted on the blog "DNA CONFESSES DATA SPEAK", by Dr. Ming Tang.

Note that the genes (y-axis) **will be displayed following the order of the grouping variable (column two of your file) selected by the user**. Once the plot is generated, users can choose the order of the clusters to show on the x-axis. For example, we show the expression profile of the three most significant gene markers identified for each cluster of the PBMC dataset.

An arbitrary order of the clusters is used in the plot, demonstrating how users can customize the result.

17.2 Multiple-genes Dot plot

A multiple-genes dot plot will be generated following the same order selected for the stacked violin plot.

Warning: Please be aware that Asc-Seurat uses multiple R packages and that many of those are in continuous development. While the docker version of Asc-Seurat is stable, it may become outdated as the packages on wich it relies on are updated. *Here* you can find a list of the packages used by Asc-Seurat and their versions.

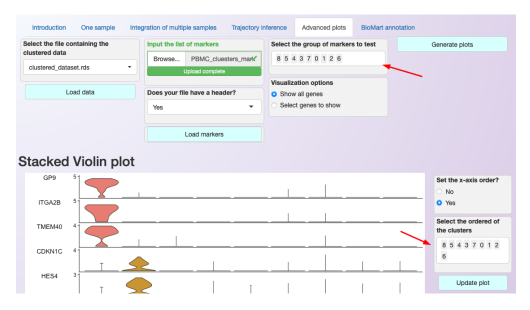


Fig. 1: Interface for generating multiple genes plot. Note that users can select the order that genes (y-axis) and clusters (x-axis) are shown; see the red arrows in the image.

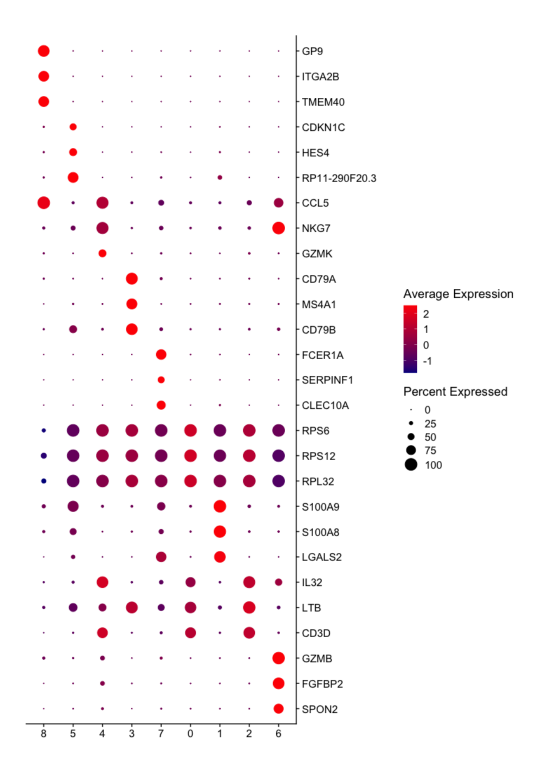


Fig. 3: Multiple-genes dot plot showing the three most significant markers of each cluster of the PBMC dataset. Observe that some of the significant markers are not specific for the cluster but present a higher level of expression than the other clusters.

EIGHTEEN

RELEASE NOTES

Tip: To use an old version of Asc-Seurat, check here.

NINETEEN

V2.2 - RELEASE ON FEBRUARY 8TH, 2022

- Add the capacity to load a clustered dataset in the tab for the individual sample analysis.
- Add the capacity to load a clustered dataset in the tab for the integrated sample analysis.
- Genes identified as mitochondrial genes via the regex expression are now shown to the users.
- Changes the color scheme of the dynverse's plots to match the color scheme used by Seurat's plots.
- Small changes in the interface to improve usability.
- Fix a bug in the download of markers identified for multiple clusters in the integrated dataset. If a gene was identified as a marker in multiple clusters, a number was appended in the gene's name.
- Fix a bug that caused the app to crash when searching for conserved markers in an integrated dataset, and the gene was not expressed in one or more of the samples.
- Fix a bug where plots were exported with a dark background.
- Fix a bug in the advanced plots that caused expressed genes not to be identified. When using integrated datasets, the function now looks for the RNA assay instead of the integrated assay.
- Fix a bug where the app would crash when downloading the plots generated in the trajectory inference tab.

TWENTY

V2.1 - RELEASED ON MAY 26TH, 2021.

- Changes the assay used for differential expression analysis and visualization to "RNA" when using SCTransform normalization. Therefore, "SCT" assay is used for the steps until clustering the data.
- Changes the output of the differential expression analysis to the format required for the visualization tools.

TWENTYONE

V2.0 - RELEASED ON MAY 19TH, 2021.

- Inclusion of SCTransform normalization
- Addition of stacked violin plots
- Addition of multiple-genes dot plot
- Improvements on the user interface
- Improvements in the app stability
- Fix of minor bugs.

TWENTYTWO

V1.0 - RELEASED ON MARCH 19TH, 2021.

• Release of Asc-Seurat.

CHAPTER TWENTYTHREE

REFERENCE

[1] Pereira WJ, Almeida FM, Balmant KM, Rodriguez DC, Triozzi PM, Schmidt HW, Dervinis C, Pappas Jr. GJ, Kirst M. Asc-Seurat: analytical single-cell Seurat-based web application. BMC Bioinformatics 22, 556 (2021).

Pereira WJ, Almeida FM, Balmant KM, Rodriguez DC, Triozzi PM, Schmidt HW, Dervinis C, Pappas Jr. GJ, Kirst M. Asc-Seurat – Analytical single-cell Seurat-based web application. BioRxiv, 2021.

TWENTYFOUR

SUPPORT CONTACT

Have any questions or suggestions? Please contact us at GitHub.

Footnotes: