

Annotator

The widget provides an option to annotate cells with cell types based on marker genes.

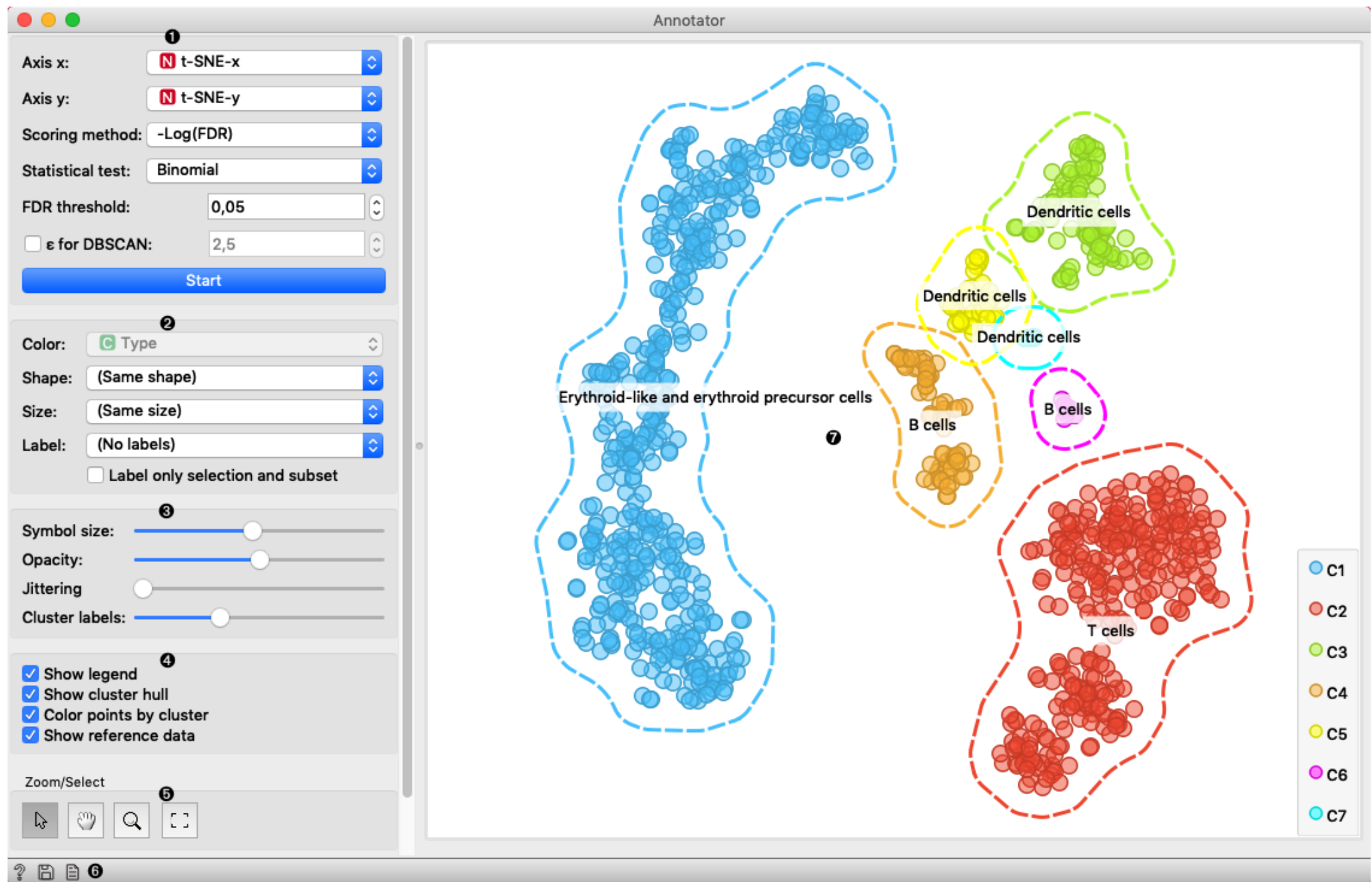
Inputs

- Reference Data: Data set with gene expression values.
- Secondary Data: Subset of instances (optional).
- Genes: Marker genes.

Outputs

- Selected Data: Instances selected from the plot.
- Data: Data with additional columns with annotations, clusters, and projection

This widget receives gene expression data together with mapping to a two-dimensional space and marker genes. It selects the most expressed genes for each cell with the Mann-Whitney U test and computes the p-value of each cell types for a cell based on the selected statistical test. It visualizes groups of cells and for each group, it shows few most present cell types.



1. This box contains settings for attribute selection and annotation:

- *Axis X* and *Axis Y* let you select attributes you will show in the widget. By default t-SNE-x and t-SNE-y are selected which are transformations to lower dimensional space with a method named t-SNE.
 - *Scoring method* allows you to select the method to score the cell type affiliation to the cell. It has the following options:
 - *-Log(FDR)* - negative of the logarithm of an FDR value
 - *Marker expression* - the sum of expressions of genes typical for cell type.
 - *Marker expression %* - the proportion of genes typical for a cell type expressed in the cell.
 - *Statistical test* allow the user to select between the binomial and hypergeometric statistical test for computing the p-value.
 - *FDR threshold* sets the threshold for FDR value. Cell types that have an FDR value below this threshold are selected.
 - *ϵ for DBSCAN* regulates the ϵ parameter of a DBSCAN algorithm which forms groups in the visualization. When unchecked algorithm estimates this parameter itself.
 - *Start* run the assigning process. Whenever you change any parameter in this box rerun the process with this button.
2. Set the color of the displayed points (you will get colors for discrete values and grey-scale points for continuous). Set label, shape, and size to differentiate between points.
 3. Set symbol size and opacity for all data points. Set **jittering** to prevent the dots overlapping. Set the number of labels shown for a cluster.
 4. Adjust *plot properties*:
 - *Show legend* displays a legend on the right. Click and drag the legend to move it.
 - *Show cluster hull* regulate whether the hull around cluster is present or not.
 - *Color points by cluster* colors the points with cluster specific color. When this option is checked setting color in box 2 is disabled.
 - *Show reference data* makes data from the *Reference data* input visible.
 5. *Select*, *zoom*, *pan*, and *zoom to fit* are the options for exploring the graph. The manual selection of data instances works as an angular/square selection tool. Double click to move the projection. Scroll in or out for zoom.
 6. This group of buttons allows you to get help about the widget, save the created image to your computer in a .svg or .png format, and create the report.
 7. The main view shows you data clustered data items. Each cluster is surrounded by a hull and has assigned *Cell type* labels shown.

Examples

In this example *Single Cell Datasets* widget provides *Bone marrow mononuclear cells with AML (sample)* gene expression dataset and *t-SNE* widget maps data into two-dimensional space. *Marker Genes* widget provides markers genes with cell types to the *Annotator* widget. The data table receives the output of the Annotator widget and shows items selected in the plot. The widget shows clustered cells projected to the two-dimensional plane. Clusters labels show annotation with the most common label in the cluster.

Single Cell Datasets

Info
30 datasets
1 dataset cached

Filter

Title	Size	Cells	Genes	Organism	Target
Bone marrow mononuclear cell...	345.0 KB	1000	1000	Human	C category
Dendritic cells and monocytes i...	19.4 MB	1140	26593	Human	
Dendritic cells and monocytes i...	18.1 MB	1244	26593	Human	
Bone marrow mononuclear cell...	2.7 MB	8390	1000	Human	C category
Pancreas cells in mouse	4.6 MB	1886	14881	Mouse	C category
Cell cycle in T-cells	4.4 MB	81	38293	Mouse	

Description

Bone marrow mononuclear cells with AML (sample) (2017), from 10x Genomics

Gene expressions in bone marrow mononuclear cells from a patient with acute myeloid leukemia (AML) and two healthy donors used as controls. The data includes a sample of 1000 cells and 1000 genes with the highest dispersion. This is a sample data that comes with [Loupe Cell Browser](#), and includes cells from three separate

☒ Send Data Automatically

Data Table

Entrez ID	Type	Acinar cells	Adipocytes	airway epithelial ce	airway goblet cells	alveolar macrophag	Astrocytes	B ce
1	healthy	4.21438	?	?	?	9.67762	6.8608	1
2	healthy	?	?	?	?	6.15625	4.50678	6
3	AML	3.84602	?	?	?	4.99512	3.51113	1
4	AML	?	?	?	?	?	?	1
5	AML	?	?	?	?	?	?	1
6	healthy	3.86441	?	?	?	9.07671	3.43955	2
7	healthy	?	?	?	?	?	4.67893	?
8	healthy	4.03232	?	?	?	13.7344	6.77734	5
9	healthy	?	?	?	?	5.4086	3.90313	9
10	healthy	?	?	?	?	15.3364	?	?
11	AML	?	5.42241	?	?	9.64045	4.02387	3
12	healthy	?	3.27153	?	?	5.80433	4.35405	6
13	AML	?	3.06088	?	?	?	?	6
14	healthy	4.3202	?	?	?	14.6279	7.49452	9
15	healthy	3.96572	?	?	?	18.7815	3.50999	1
16	healthy	?	?	?	?	14.6457	3.8883	1
17	healthy	4.18055	?	?	?	14.2353	?	3
18	healthy	4.35917	?	?	?	10.0768	?	6

Annotator

Axis x:

Axis y:

Scoring method:

Statistical test:

FDR threshold:

☐ ε for DBSCAN:

Start

Color:

Shape:

Size:

Label:

☐ Label only selection and subset

Symbol size:

Opacity:

Jittering: ☐

Cluster labels:

☒ Show legend
☒ Show cluster hull
☒ Color points by cluster
☒ Show reference data

Zoom/Select

Legend:

- C1
- C2
- C3
- C4
- C5
- C6
- C7

Annotations in plot:

- Erythroid-like and erythroid precursor cells
- Dendritic cells
- B cells
- T cells

```

graph LR
    A[Single Cell Datasets] --> B[t-SNE]
    B --> C[Annotator]
    C --> D[Data Table]
    E[Marker Genes] --> C
    F[Data] --> B
    G[Selected Data] --> D
  
```

The diagram illustrates the data workflow. It starts with 'Single Cell Datasets' and 'Data' inputs feeding into a 't-SNE' widget. The output of 't-SNE' goes to the 'Annotator' widget, which also receives 'Marker Genes' as input. The 'Annotator' then outputs 'Selected Data' to a 'Data Table' widget.

The second example shows how to use *Secondary Data* input of the widget. We load data with two *Single Cell Datasets* widgets. First widget loads *Cell cycle in mESC (Fluidigm)* data which are used as primary data. The second widget loads *Cell cycle in mESC (QuartzSeq)* dataset for secondary data input. After data are loaded, we normalize them with the *Single Cell Preprocess* widget and map the reference data to two-dimensional space with the t-SNE projection. Projecting the secondary data is not required since the Annotator widget projects them in the reference space. The Annotator window shows the mapping of secondary data (colored points) to clusters generated on the reference data.

