

adjClust Package Vignette

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1 Introduction

This package implements the Adjacency Clustering algorithm designed by Chris Bolen.

2 Installation

Currently, installing AdjClust takes a bit of work. This will eventually be streamlined, but until the code is done, you'll need to download the source code from Bitbucket and compile it according to the instructions below.

2.1 Install required packages

```
> install.packages(c("gplots", "hexbin", "parallel", "png"))
> source("https://bioconductor.org/biocLite.R")
> biocLite(c("cytofkit", "flowCore"))
```

2.2 Building adjClust

The build instructions assume that you have downloaded the most recent version of AdjClust, and have started an RStudio project in the download directory.

1. Install build dependencies:

```
> install.packages(c("devtools", "roxygen2", "testthat", "knitr", "rmarkdown"))
```

2. Build with RStudio

- Build -> Configure Build Tools
- Check use devtools option
- Check use roxygen option
- Select configure roxygen options and check everything.
- Build -> Build and Reload

2.3 Compiling tSNE

Before tSNE can be used, you must compile the source code. AdjClust contains a function, `compileTSNE`, which will automatically compile the code and make it usable within R. To use your system's default c++ compiler, simply call `compileTSNE` using the default parameters:

```
> compileTSNE()
```

2.3.1 Compiling tSNE with OpenMP support

If you are planning to run the adjClust algorithm with large datasets, it is almost a necessity that the tSNE algorithm be run in parallel. To do this, we will need to compile the code with OpenMP support. On Linux and Windows systems, this should be as simple as adding a single parameter to the `compileTSNE` function call:

```
> compileTSNE(use_omp=TRUE)
```

OpenMP support on OSX

As of October 2015, the default compiler for Mac operating systems ("clang++") does not come with a working copy of OpenMP. That means that in order to enable OpenMP on a mac, you must install a different c++ compiler. There are many options available, but the Gnu compilers are one of the most popular. To install a recent version, follow the instructions below:

1. Install Homebrew (<http://brew.sh/>):

```
ruby -e "$(curl -fsSL https://raw.githubusercontent.com/Homebrew/install/master/install)"
```

2. Install gcc version 4.9:

Open up the terminal and run:

```
brew install gcc49
```

3. Confirm GCC 4.9 is installed:

```
g++-4.9 --version
```

Once gcc has been successfully installed, tSNE can be compiled with the newly installed compiler as follows:

```
> compileTSNE(compiler = "g++-4.9", use_omp = T)
```

```
Finding blas installation...Done!
```

3 The AdjClust workflow

3.1 Step 1: Reading in the FCS files

The first step in Adjacency Clustering is to perform dimensionality reduction with the tSNE algorithm. In the future we'll make this nice and streamlined, but for now it's gonna be a bit of a hassle. To do this, we need to read the data into R so that we can process it and get it into the format we want.

Let's say we have a folder filled with FCS files:

```
> fcsFolder = "~/fcsFiles"
> ##read all the filenames into a vector
> fcsFiles = dir(fcsFolder, full.names = T)
> fcsFiles

[1] "/Users/chris/fcsFiles/set1.poolA.1172.wk104.ns.CD8T.fcs"
[2] "/Users/chris/fcsFiles/set1.poolA.1172.wk104.pi.CD8T.fcs"
[3] "/Users/chris/fcsFiles/set1.poolA.1172.wk18.ns.CD8T.fcs"
[4] "/Users/chris/fcsFiles/set1.poolA.1172.wk18.pi.CD8T.fcs"
[5] "/Users/chris/fcsFiles/set1.poolA.1172.wk40.ns.CD8T.fcs"
[6] "/Users/chris/fcsFiles/set1.poolA.1172.wk40.pi.CD8T.fcs"
[7] "/Users/chris/fcsFiles/set1.poolA.1172.wk53.ns.CD8T.fcs"
[8] "/Users/chris/fcsFiles/set1.poolA.1172.wk53.pi.CD8T.fcs"
[9] "/Users/chris/fcsFiles/set1.poolA.ctrl1.NA.ns.CD8T.fcs"
[10] "/Users/chris/fcsFiles/set1.poolA.ctrl1.NA.pi.CD8T.fcs"
```

The first thing we need to do is read in all of these fcs files. To do this, we need to use the `flowCore` package from Bioconductor.

```
> library(flowCore)
> fcsDat = lapply(fcsFiles,function(fn){read.FCS(fn)})
```

We're then going to take each one of these FCS objects and extract the marker expression data from them:

```
> fcsMats = lapply(fcsDat, function(fcs){
+   #get marker vals
+   eset = exprs(fcs)
+
+   ##replace channel names with marker names
+   channels = fcs@parameters@data$desc
+   names(channels) = colnames(eset)
+   channels[is.na(channels)] = colnames(eset)[is.na(channels)]
+   colnames(eset) = channels
+ })
```

```

+
+   return(eset)
+ })
> fcsMats[[1]][1:10,1:6]

```

	Time	Event_length	BC1	BC2	BC3	BC4
[1,]	2442.604	35	4.435	18.247	7.484	95.764
[2,]	5441.979	43	4.255	26.072	7.239	126.849
[3,]	5826.432	40	0.000	36.553	7.302	90.253
[4,]	8387.188	38	0.000	17.719	7.852	95.326
[5,]	8915.117	37	0.000	20.914	7.599	60.876
[6,]	9489.388	30	0.000	29.219	9.105	26.002
[7,]	10560.703	27	1.199	26.466	8.894	171.672
[8,]	11107.930	61	0.000	29.771	9.983	112.359
[9,]	11484.792	30	0.050	23.987	8.285	89.709
[10,]	12380.052	44	1.430	41.458	10.950	83.965

(side note: if you're working with lots of fcs files, you're gonna be using a lot of memory here. To mitigate this problem, remove objects as you go):

```
> rm(fcsDat)
```

Note that each one of these matrices are going to have different numbers of cells, but the column names of each of them should match:

```
> ##the number of cells
> sapply(fcsMats, nrow)
```

```
[1] 1093 1612 11135 8688 2063 1018 19310 13363 861 464
```

```
> ##the columns in each matrix
> sapply(fcsMats, colnames)[1:10,1:4]
```

	[,1]	[,2]	[,3]	[,4]
Time	"Time"	"Time"	"Time"	"Time"
Event_length	"Event_length"	"Event_length"	"Event_length"	"Event_length"
Pd104Di	"BC1"	"BC1"	"BC1"	"BC1"
Pd106Di	"BC2"	"BC2"	"BC2"	"BC2"
Pd108Di	"BC3"	"BC3"	"BC3"	"BC3"
In113Di	"BC4"	"BC4"	"BC4"	"BC4"
In115Di	"CD57"	"CD57"	"CD57"	"CD57"
La139Di	"CD16"	"CD16"	"CD16"	"CD16"
Ce140Di	"beads"	"beads"	"beads"	"beads"
Pr141Di	"IL8"	"IL8"	"IL8"	"IL8"

Now, optionally: if we have lots of fcs files, each with lots of cells, we may want to subsample the files to make sure the tSNE algorithm doesn't take forever. To do this, we're going to set the maximum number of cells for each matrix to 10,000 and randomly subsample any fcs files that are larger than that number:

```
> ##maximum number of cells in each matrix
> maxCell = 10000
> ##subsample each matrix
> fcsSubsample = lapply(fcsMats, function(mat){
+   #pick which rows to keep
+   i = sample(1:nrow(mat), min(nrow(mat), maxCell))
+   #return those rows
+   mat[i,]
+ })
>
```

And now we can combine them together into a single matrix:

```
> combinedEset = do.call(rbind, fcsSubsample)
> dim(combinedEset)
```

```
[1] 45799    49
```

Ideally, your final dataset should have less than 1 million cells. More than that is okay, but the processing time increases exponentially with the number of cells, and even on a fast computer 2 million cells would take more than a day to run. Additionally, the quality of the map will decrease when the number of cells is above 2.5 million, so it is recommended that you never go above that.

In this case, we have a bit under 46,000 cells, which is a very reasonable number. However, before we can run tSNE, there's three more things we need to do. First, select the markers we want to include in the map:

```
> markers = c("CD57", "IL8", "CD45RO", "perforin", "TNF", "CD25", "MIP1b", "CD103",
+             "CD107ab", "CD27", "CD45RA", "CXCR5", "CD28", "CD38", "CD69", "TGFB",
+             "IL2", "IFNG", "IntB7", "CCR7", "HLA-DR", "CD40L", "PD-1", "CD127")
> combinedEset = combinedEset[,markers]
```

This dataset only contains CD8+ T cells, so we are only going to include parameters relevant to that cell type.

Second, transform the data using the arcsinh transformation:

```
> combinedEset = apply(combinedEset, 2, arcsinhTransform())
```

And third, label which file each cell came from:

```

> ##calculate number of cells from each file
> cellCounts = sapply(fcsSubsample, nrow)
> ##make vector indicating source file for each cell
> sourceFile = rep(basename(fcsFiles), times=cellCounts)
> #store this in the rownames of combinedEset
> rownames(combinedEset) = sourceFile
> combinedEset[1:10, 1:4]

```

	CD57	IL8	CD45RO	perforin
set1.poolA.1172.wk104.ns.CD8T.fcs	5.3906711	1.1414645	2.6743522	5.9531878
set1.poolA.1172.wk104.ns.CD8T.fcs	1.2210972	0.8813736	0.8813736	2.6511672
set1.poolA.1172.wk104.ns.CD8T.fcs	5.8019100	2.3871005	0.8813736	4.4805739
set1.poolA.1172.wk104.ns.CD8T.fcs	4.7169704	0.8813736	0.8813736	4.7262266
set1.poolA.1172.wk104.ns.CD8T.fcs	2.9157235	4.5790343	0.8813736	1.3073930
set1.poolA.1172.wk104.ns.CD8T.fcs	0.8813736	2.5533300	0.8813736	0.8813736
set1.poolA.1172.wk104.ns.CD8T.fcs	2.7719971	3.0659396	0.8813736	0.8813736
set1.poolA.1172.wk104.ns.CD8T.fcs	2.1243564	2.4547809	0.8813736	0.8813736
set1.poolA.1172.wk104.ns.CD8T.fcs	0.8813736	0.8813736	0.8813736	2.4432922
set1.poolA.1172.wk104.ns.CD8T.fcs	3.4657125	5.8633495	0.8813736	0.8813736

Now we've finally finished processing the fcs files, and we can move on to running tSNE.

3.2 Step 2: Running tSNE

Again, I hope that in the future I can streamline this a lot more, but for now running tSNE takes a bit of work. Before running this section, make sure you have tSNE correctly installed and compiled according to the instructions in section 2.

Saving data for tSNE

Once we have a matrix of cells, such as the `combinedEset` matrix that we generated in section 3.1, we can finally start using the functions in `adjClust`. First, we need to save the data in a format that tSNE understands. To do this, we will use the `saveTSNEdat` function. This takes as input the `combinedEset` matrix and the folder where the data should be saved.

```
> outFolder = "~/tsneRun"
> saveTSNEdat(combinedEset, outFolder)
```

Optionally, the `saveTSNEdat` function also takes a number of other parameters, including:

- `num_proc` - The number of processors to use. Default is the maximum number for your machine.
- `theta` and `perplexity` - Run parameters for tSNE. It is generally fine to leave `theta` at the default, but it can be very useful to increase the perplexity when running tSNE with large datasets. *When running tSNE with large datasets ($\geq 500k$ cells), the perplexity should be increased to 100.* This will improve the clustering of cells into similar groups.

Running tSNE

Once the data has been saved, the tSNE algorithm can be run using the `runTSNE` function. Be warned: for large datasets, this can take more than 24 hours!

```
> runTSNE(outFolder)

Step 1: Computing input similarities...
Step 2: Building tree...
|=====| 100%
Step 3: Learning embedding...
|=====| 100%
Finished!
```

Alternatively, tSNE can be run directly on the command line. The `runTSNE` function provides an option for returning just the commands for running tSNE, which can be pasted directly into a terminal to run tSNE.

```
> runTSNE(outFolder, commandOnly=T)

cd ~/tsneRun
/private/var/folders/w9/00pdmcy55hz3hj8wm03vf0b40000gn/T/RtmpY8qrk1/Rinst3ddf652ac3d9/adjClu
```

(NOTE: This command will be different for each system. Don't just copy the command above.)

Reading in the tSNE results

Finally, once the tSNE map has finished running, the final map can be read into R using the `loadTSNEdat` function. This returns an object of class `tmap`, which contains both the newly generated tSNE map and the original marker values, as well as most of the clustering information that we are about to generate.

```
> tmap = loadTSNEdat(outFolder)
```

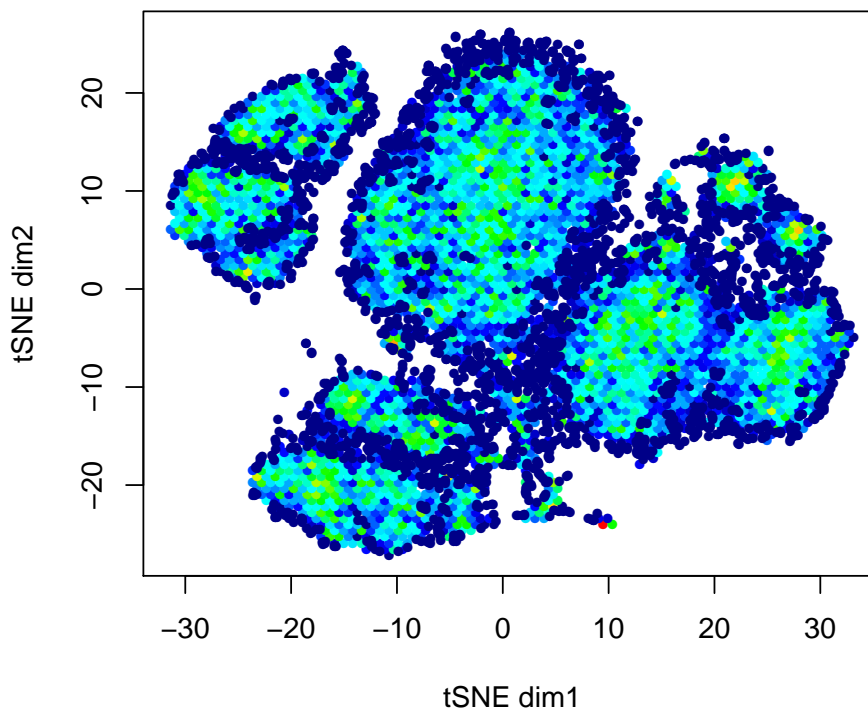
3.3 Analyzing the tSNE map

The `adjClust` package contains a number of functions for visualizing and analyzing a tSNE map. This section covers the basic plotting methods.

`hexplot`

Once the tSNE results has been loaded (via the `loadTSNEdat` function), the first step is to plot out the map.

```
> plot(tmap)
```



For `tmap` objects, the default plotting function calls `hexplot`, a highly customizable set of plotting functions specifically designed for visualizing high-density data. The default `hexplot` parameters split the plotting region up into a set of hexagonal bins, and everything in that bin is combined together into a single point, with the total number of points in each bin visualized using a color gradient. Low density bins are shown in dark blue, and higher density areas are represented using a color scale, going from cyan -> green -> yellow -> orange -> red.

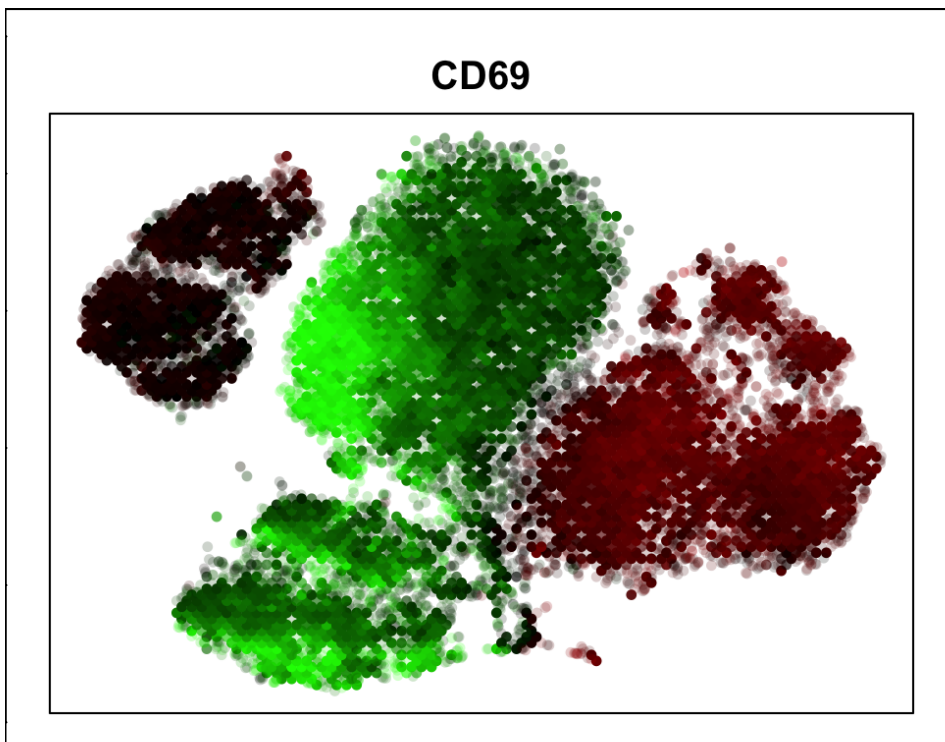
The `hexplot` function includes a number of options for making custom plots of the tSNE map, including parameters to control the colors, the number of bins, and how density information is displayed. Many of the most common customization options have been included in `adjClust` as separate functions, but it may eventually be useful to generate your own. Refer to the `hexplot` help file for more details.

plotMarkers

Now that we know what the map looks like, we can start to analyze where specific cell types have ended up on the map. We can do this using the `plotMarkers` function. This function uses parameter values from the original N-dimensional dataset to generate a custom-colored hexplot. Areas of the map with high expression of a specific parameter will be colored red, while areas with low expression will be colored green. Looking at these maps for each parameter individually makes it possible to find specific areas representing known cell types.

Below, we are plotting the expression of CD69, a marker of activation in T cells. We see that the map has three major groups: one large region in the center that is negative for CD69, and two smaller regions on the left and right sides of the map that are positive for CD69.

```
> plotMarkers(tmap, markerList=c("CD69"))
```



plotGroups

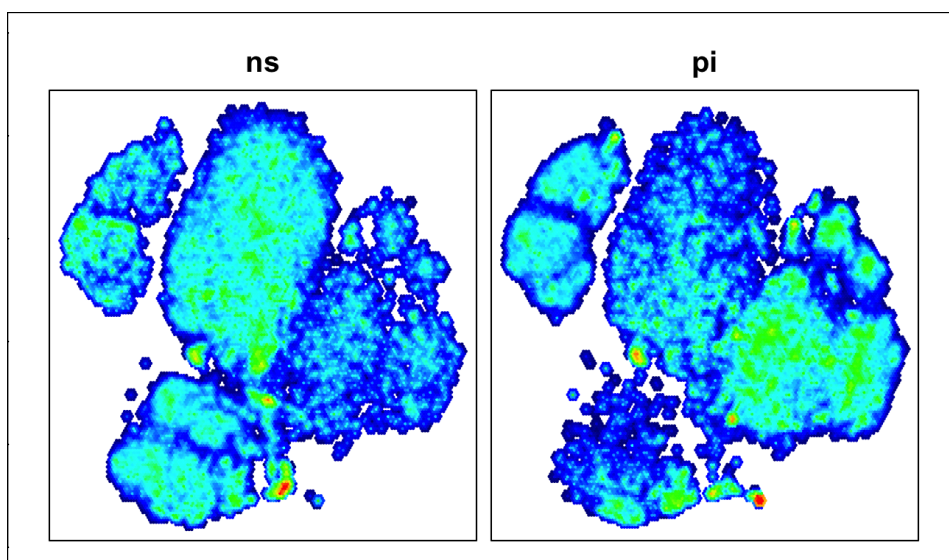
Another common task when examining tSNE maps is to compare sets of samples to each other. For example, our dataset contains ten total samples, half of which are unstimulated (ns), and the other half stimulated with PMA and Ionomycin (pi). We have information about each file stored in a data frame called `fileInfo`:

```
> fileInfo
```

	fn	stim	week	donor
1	set1.poolA.1172.wk104.ns.CD8T.fcs	ns	104	1172
2	set1.poolA.1172.wk104.pi.CD8T.fcs	pi	104	1172
3	set1.poolA.1172.wk18.ns.CD8T.fcs	ns	18	1172
4	set1.poolA.1172.wk18.pi.CD8T.fcs	pi	18	1172
5	set1.poolA.1172.wk40.ns.CD8T.fcs	ns	40	1172
6	set1.poolA.1172.wk40.pi.CD8T.fcs	pi	40	1172
7	set1.poolA.1172.wk53.ns.CD8T.fcs	ns	53	1172
8	set1.poolA.1172.wk53.pi.CD8T.fcs	pi	53	1172
9	set1.poolA.ctrl1.NA.ns.CD8T.fcs	ns	NA	ctrl
10	set1.poolA.ctrl1.NA.pi.CD8T.fcs	pi	NA	ctrl

If we want to see where the cells from "ns" and "pi" samples are located in the map, we can use the `plotGroups` function. This function takes a data frame `grp.info` which contains two columns. The first column is a list of filenames matching those in our dataset, and the second column is the group membership of that file. So if we want our two groups to be "ns" and "pi", we would run `plotGroups` as follows:

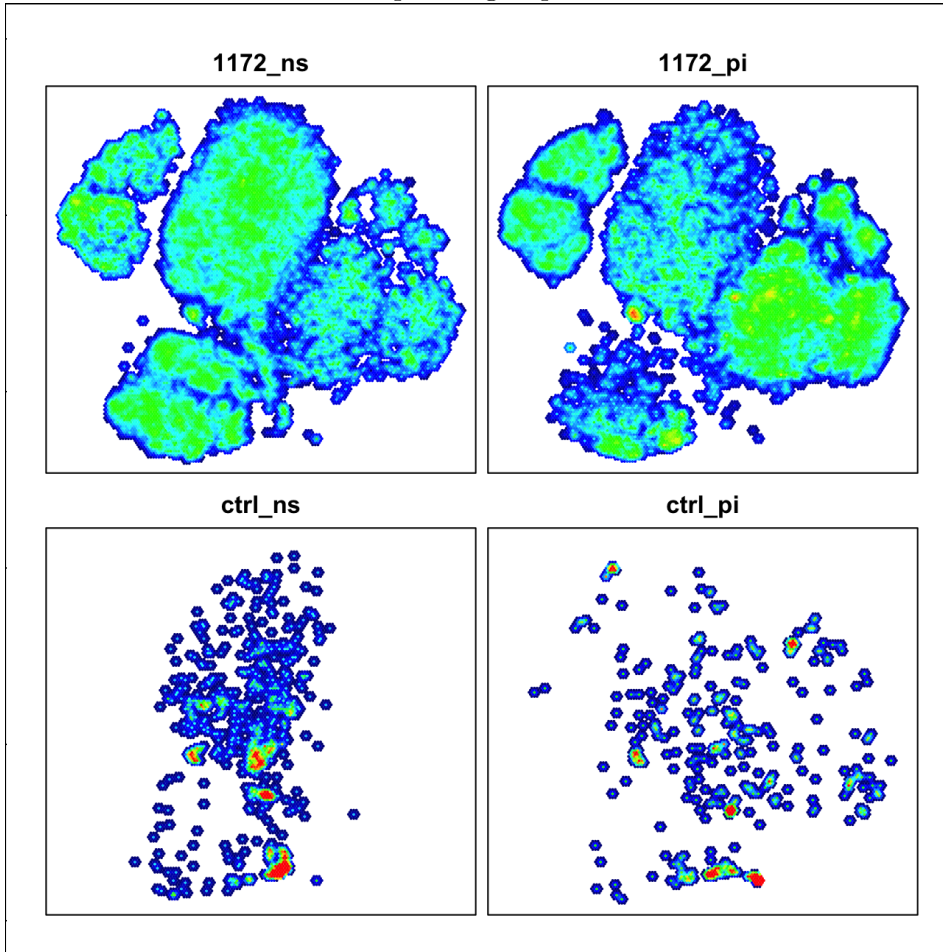
```
> grp.info = fileInfo[,c("fn", "stim")]
> plotGroups(tmap, grp.info = grp.info)
```



It is also possible to define the groups using an R formula. For example, if we want to separate out the controls from the actual samples, we can run `plotGroups` as follows:

```
> plotGroups(fn~donor+stim, data = fileInfo, tmap = tmap)
```

"fn" is name of the column containing the filenames in `fileInfo`, and "donor" & "stim" are the columns that we want to use to separate groups.



3.4 Adjacency Clustering

The stuff I describe in the previous chapters is nice, but the Adjacency Clustering algorithm is the reason this package exists. The Adjacency Clustering method is an adapted version of the same hierarchical clustering method used in *citrus*. This method clusters groups of cells based off of similar marker expression, but adds the limitation that those sets of cells must be adjacent to each other in the 2-dimensional tSNE map as well. The result is a set of smooth-bounded clusters that are both biologically relevant and easy to interpret.

This chapter describes how to use the clustering algorithm, how to tweak the parameters for optimal performance, and how to determine the markers that define those clusters.

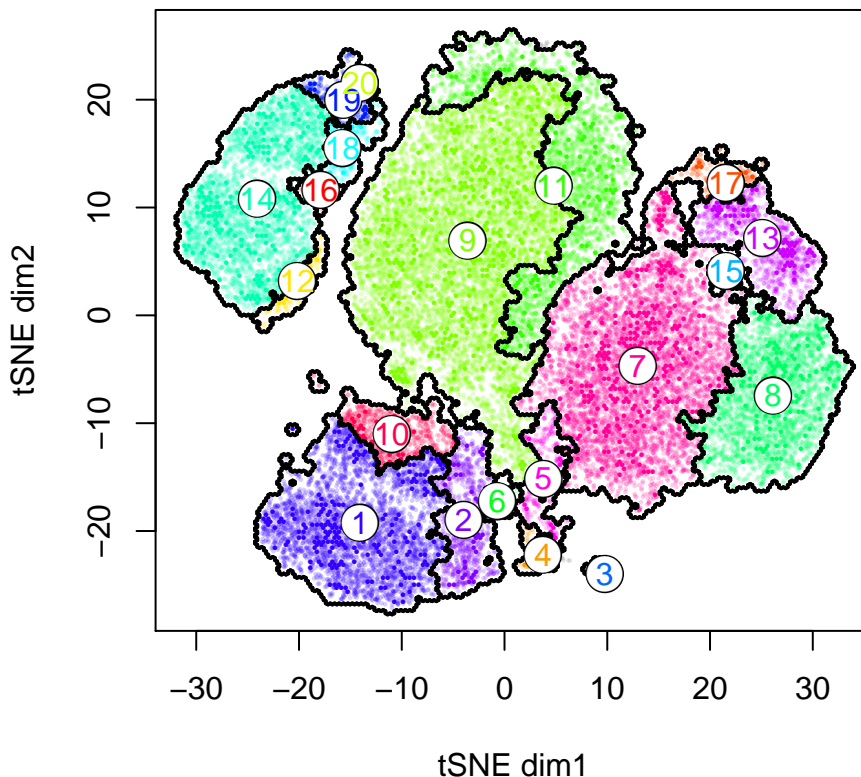
To start, we will attempt to cluster our tmap using the `cluster.tmap` function. In addition to taking the `tmap` object we loaded in using `loadTSNEdat`, this function also requires that you enter a number of clusters, `nclust`. There's no good rule for how many clusters to use, so we're going to start with a guess and change it as we see fit.

```
> tmap = cluster.tmap(tmap, nclust = 20)
```

(Note: The `cluster.tmap` function adds cluster information to our original `tmap` object, so we are overwriting the previous `tmap` object to save memory)

We can see how the clustering worked using the `plotClusters` function:

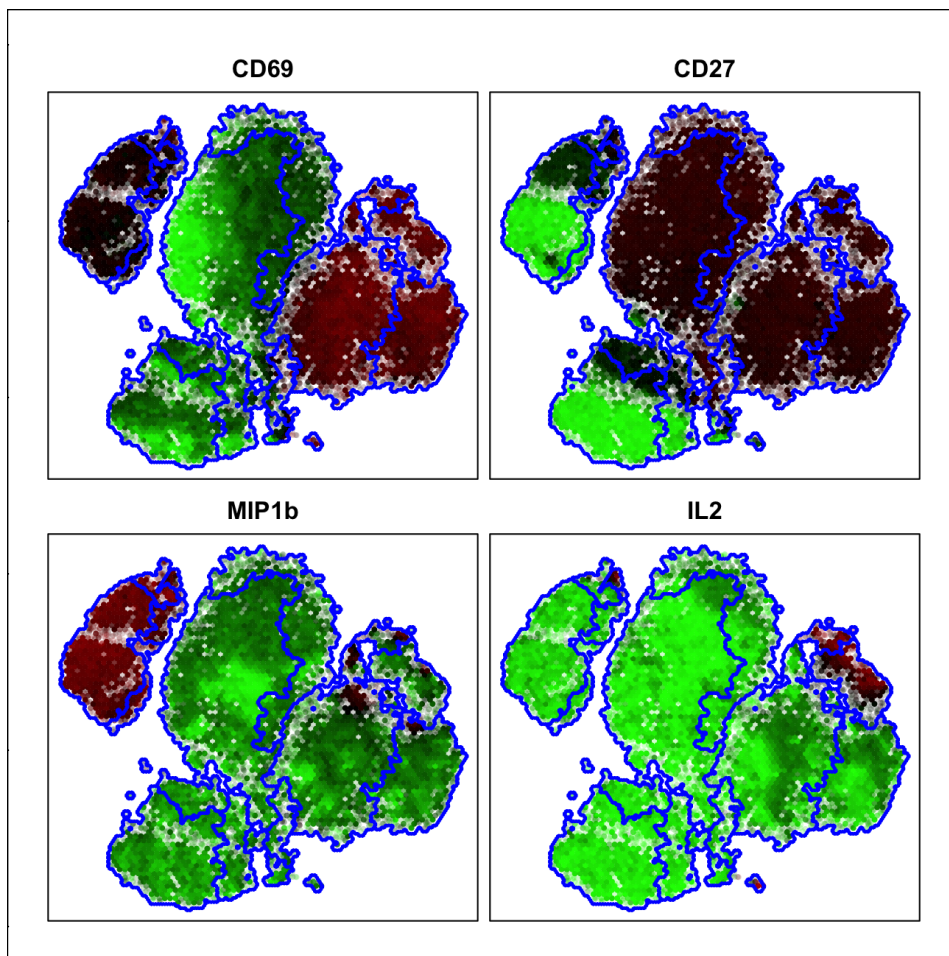
```
> plotClusters(tmap)
```



This function generates a `hexplot` with each cluster assigned a random color (if we run the function again, the colors of each cluster will change). Based on this map, we can see that the cluster edges generally do a good job at following the general topography of the tSNE map, but it's definitely not perfect.

To see how well these clusters match up against marker expression, we can again use the `plotMarkers` function.

```
> plotMarkers(tmap, markerList = c("CD69", "CD27", "MIP1b", "IL2"))
```



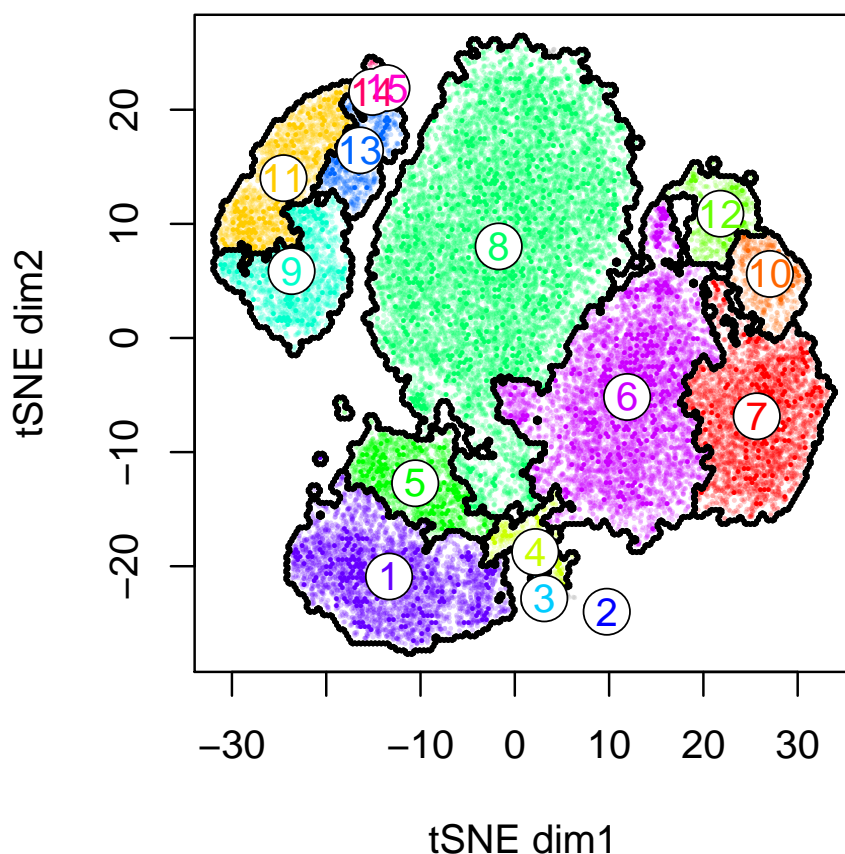
Above is just an example with four of the markers. Now that clustering has been performed, the cluster outlines have been automatically added to these plots (**Note: this will actually work for *all* the plotting methods we've used so far**). Adding the cluster outlines makes it much easier for us to see how the cells are clustering together, and whether or not the clusters accurately follow the marker expression.

Looking at these clusters, two things become apparent: First, that we chose to make too many clusters. There seems to be a large number of clusters that do not follow any marker, suggesting that boundaries have been defined based on very small changes in expression. Thus, lowering the number of clusters is probably appropriate. Second, some cluster edges seem to be following marker differences that are less important. For example, right in the middle of the

map there is a cluster border separating cells with low (green to black) expression of MIP1b from those with zero (bright green) expression. While this difference certainly exists, it is most likely uninteresting from a biological perspective, as both groups would most likely be considered negative for MIP1b expression. To rectify this, we can provide weights for specific parameters, which will allow the clustering algorithm to favor differences in some parameters over others. Specifically, we'll give less weight to the MIP1b parameter while also increasing the weight of those parameters which the clusters don't adequately follow.

After some iteration and studying the clusters, I've come up with weightings for many of the markers in our map. Here's our second attempt at clustering:

```
> markerWt = c(MIP1b=0.75, CD27=1.5, IL2=1.5, IntB7=0.75,
+              CXCR5=0.5, CD25=0.5, CD107ab=0.5, CD69=1.25,
+              IFNg=1.3, CD28=0.75, IL8=1.25, perforin=1.2)
> tmap = cluster.tmap(tmap, nclust = 15, markerWt = markerWt)
> plotClusters(tmap)
```



These clusters looks pretty good. We could play with it more, but they appear to follow the general trends in the data quite well, and hopefully they are much easier to interpret than the default clusters.

3.4.1 Defining cluster phenotypes

Most of the time, playing around with the marker weights is enough to learn what markers define each cluster. However, sometimes it is useful to dig a little deeper into individual clusters to see what makes them unique. The `adjClust` package contains a method, `clusterReport`, which gives a summary of the major differences in marker expression between neighboring clusters. To generate the cluster report, run the function as follows:

```
> report = clusterReport(tmap)
```

The report is a list containing reports for every individual cluster. To view the report for a specific cluster, we can look at that entry in the list:

```
> report[[8]]
```

```
[1] "Compared to 4: Low in CD45R0; High in CD45RA;"
[2] "Compared to 5: Low in CD57, perforin; High in CD27, CD28, CCR7, CD127;"
[3] "Compared to 6: Low in CD69;"
```

This report implies that cluster 8 is CD45RA+CCR7+ (i.e. naive), and CD69 low (resting). This lines up pretty well with what we saw from the marker maps, and makes it easy to characterize the phenotype of those cells. However, some of the clusters are harder to figure out. Cluster 14, for example, is small and seems to be difficult to characterize. Even worse, the cluster report doesn't appear to tell us anything about this cluster.

```
> report[[14]]
```

```
[1] "Compared to 13:" "Compared to 15:"
```

That's because the differences in this cluster are smaller. The `clusterReport` works by only examining those markers that are most different. By default, it will only report the 5% of markers that are most differentially expressed across all clusters, but we can change this parameter to make the cutoffs more relaxed.

```
> report = clusterReport(tmap, alpha = 0.1)
> report[[14]]
```

```
[1] "Compared to 13: High in CD69;"
[2] "Compared to 15: Low in TGFb, IL2;"
[3] "Compared to total map: High in TNF;"
```

This allows us to see the smaller differences between these clusters. However, beware: now that the cutoffs are lower, we may see differences that are much less important. For example, the expression of CD69 is higher in cluster 14 than cluster 13, but looking back at the marker map in section 3.3, we see that all of the clusters in this area are high for CD69, and cluster 14 is simply the highest.

3.5 Cluster Analysis

One of the primary goals for cluster analysis is to compare the relative frequencies of a cluster between two (or more) groups of samples. We've already shown in section 3.3 how to use the `plotGroups` function to visually compare sets of samples. In this section, we'll detail how to make statistical comparisons of cluster membership between sample sets and how to extract the cluster data so that you can perform your own analysis outside of the `adjClust` package.

3.5.1 Comparing groups

The `adjClust` package contains a function, `compareGroups`, which is designed to perform a quick automated comparison of two groups of samples. This function works more or less the same as the `plotGroups` function that we used in section 3.3. The function takes three parameters as input, our `tmap` object, the `grp.info` data frame, and a `contrast` string, which describes the comparison that we want to make. We're going to use the same `"grp.info"` object that we created in that section for this analysis:

```
> head(grp.info)

              fn stim
1 set1.poolA.1172.wk104.ns.CD8T.fcs ns
2 set1.poolA.1172.wk104.pi.CD8T.fcs pi
3 set1.poolA.1172.wk18.ns.CD8T.fcs ns
4 set1.poolA.1172.wk18.pi.CD8T.fcs pi
5 set1.poolA.1172.wk40.ns.CD8T.fcs ns
6 set1.poolA.1172.wk40.pi.CD8T.fcs pi
```

The comparison is then run as follows:

```
> cmpInfo = compareGroups(tmap, grp.info = grp.info, contrast = "pi-ns")
> head(cmpInfo)

  cluster statistic p.value  fdr
4         4         4 0.0947 0.444
6         6        21 0.0947 0.444
7         7        21 0.0947 0.444
5         5         6 0.2100 0.444
12        12        19 0.2100 0.444
10        10        18 0.2950 0.444
```

The table details the results of a two-sample test comparing the "ns" and "pi" groups, with info on p-values, test statistics, and fdr included for each cluster (In this case, the "statistic" column refers to the W statistic from the Wilcoxon rank-sum test. More extreme values—i.e. close to either 0 or $N1 \cdot N2$ —represent more significant tests, and thus will have more significant p-values. If we want to use a t-test instead, we can specify that `parametric=TRUE`, meaning that we should use a parametric test).

We see that although none of the p-values reach significance with an alpha of 0.05, the most significant clusters are numbers 4, 6 and 7; all of which are, based on the `clusterReport`, most likely activated cell types.

For more complicated group structures, it is possible to define groups using an R formula, just like we did for `plotGroups`. Below, we use this function to compare the ns and pi samples from a single donor, effectively dropping the controls from the analysis:

```
> cmpInfo = compareGroups(fn~donor+stim, data = fileInfo,
+                          contrast = "1172_pi-1172_ns", tmap = tmap)
> head(cmpInfo)
```

	cluster	statistic	p.value	fdr
4	4	2	0.112	0.421
5	5	2	0.112	0.421
6	6	14	0.112	0.421
7	7	14	0.112	0.421
3	3	4	0.186	0.558
1	1	4	0.312	0.642

Again, clusters 4, 6, and 7 come out as significant, although the p-value is slightly higher due to the reduced numbers of samples.

3.5.2 Cluster counts and percentages

There are far too many ways to analyze clustering data, and it would be silly to try and include them all in this package. However, the package provides two simple functions, `clusterCounts` and `clusterFract`, which return the cell counts and cell fractions, respectively, for each sample and cluster.

```
> cts = clusterCounts(tmap)
> cts[1:4,1:6]
```

	1	2	3	4	5	6
set1.poolA.1172.wk104.ns.CD8T.fcs	138	0	0	9	8	257
set1.poolA.1172.wk104.pi.CD8T.fcs	383	0	0	13	42	61
set1.poolA.1172.wk18.ns.CD8T.fcs	1253	15	6	251	1172	201
set1.poolA.1172.wk18.pi.CD8T.fcs	126	2	0	10	28	2523

```
> pct = clusterFract(tmap)
> signif(pct[1:4,1:4],3)
```

	1	2	3	4
set1.poolA.1172.wk104.ns.CD8T.fcs	0.1260	0.00000	0e+00	0.00823
set1.poolA.1172.wk104.pi.CD8T.fcs	0.2380	0.00000	0e+00	0.00806
set1.poolA.1172.wk18.ns.CD8T.fcs	0.1250	0.00150	6e-04	0.02510
set1.poolA.1172.wk18.pi.CD8T.fcs	0.0145	0.00023	0e+00	0.00115

These matrices can be used for whatever analyses you like, so do with them as you will.