

Methods for Unsupervised Clustering

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2025-07-28





Supervised vs. Unsupervised Machine Learning

Machine learning algorithms are generally classified into two categories. In **Supervised** machine learning we use the outcomes in a training set to **supervise** the creation of our prediction algorithm.

In **unsupervised** machine we do not necessarily know the outcomes and instead are interested in discovering groups. These algorithms are also referred to as **clustering** algorithms since predictors are used to define **clusters**.





Supervised vs. Unsupervised Machine Learning

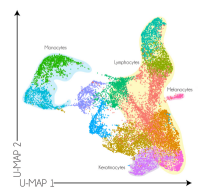
Sometimes clustering is not be very useful. For example, if we are simply given the heights we may not be able to discover two groups, males and females.





Unsupervised Machine Learning

However, there are applications in which unsupervised learning can be a powerful technique, such as an exploratory tool:







Unsupervised Machine Learning

There are many algorithms for unsupervised learning. We have already learned about **PCA** and **UMAP** for dimension reduction. Here we introduce two methods for clustering: **hierarchical clustering** and **k-means**.





Unsupervised Machine Learning and clustering

A first step in any clustering algorithm is defining a distance between observations or groups of observations.

Hierarchical clustering starts by defining each observation as a separate group, and distances are calculated between every group (distance matrix). Then the two closest groups are merged into a single group, and this new group (two observations) is represented by its centroid. Distances between this new group and the rest are calculated, and then the next two closest groups are merged. This process is repeated until there is just one group.



Consider the ratings of 50 movies from 139 different critics:

```
library(dslabs); data("movielens")
top <- movielens %>% group_by(movieId) %>%
  summarize(n=n(), title = first(title)) %>%
  top_n(50, n) %>% pull(movieId)
x <- movielens %>%filter(movieId %in% top) %>%
  group_by(userId) %>% filter(n() >= 25) %>%
  ungroup() %>% select(title, userId, rating) %>%
  spread(userId, rating)
row_names <- str_remove(x$title, ": Episode") %>% str_trunc(20)
x <- x[,-1] \%  as.matrix()
x \leftarrow sweep(x, 2, colMeans(x, na.rm = TRUE))
x \leftarrow sweep(x, 1, rowMeans(x, na.rm = TRUE))
rownames(x) <- row_names</pre>
```





We want to use these data to find out if there are clusters of movies based on the ratings from 139 movie raters. A first step is to find the distance between each pair of movies using the dist function:

 $d \leftarrow dist(x)$





With the distance between each pair of movies computed, we need an algorithm to define groups from these. The hclust function implements this algorithm and it takes a distance as input.

```
h <- hclust(d)
h
##
## Call:
## hclust(d = d)
##
## Cluster method
                     : complete
  Distance
                     : euclidean
  Number of objects: 50
```

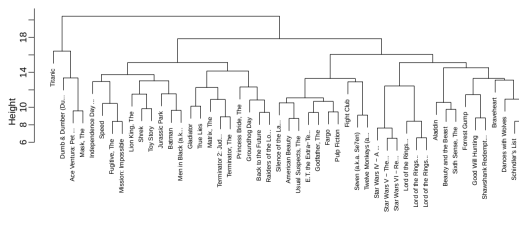




Apollo 13

Hierarchical clustering

We can see the resulting groups using a dendrogram.



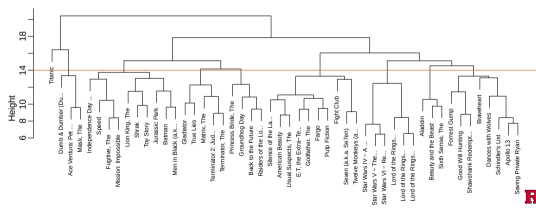


This graph gives us an approximation between the distance between any two movies. To find this distance we find the first location, from top to bottom, where these movies split into two different groups. The height of this location is the distance between these two groups. So, for example, the distance between the three *Star Wars* movies is 8 or less, while the distance between *Raiders of the Lost of Ark* and *Silence of the Lambs* is about 17.





To generate actual groups: 1) decide on a maximum distance to be in the same group or 2) decide on the number of groups. For example:





3 3 10 8 4 10 5 6 1

The function cutree can be applied to the output of hclust to perform either of these two operations and generate groups.

```
# Maximum Distance
groups <- cutree(h, h = 14)
table(groups)
## groups</pre>
```





Or we can do fewer groups:

```
#Number of groups
groups <- cutree(h, k = 10)
table(groups)

## groups
## 1 2 3 4 5 6 7 8 9 10
## 3 3 10 8 4 6 4 5 6 1</pre>
```





The clustering provides some insights, e.g., Group 4 appears to be blockbusters:

And group 9 appears to be fantasy/nerd movies:

```
mames(groups)[groups==9]
## [1] "Lord of the Rings..." "Lord of the Rings..."
## [4] "Star Wars IV - A ..." "Star Wars V - The..." "Star Wars VI - Re..."
```

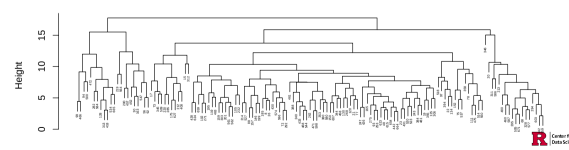




We can also explore the data to see if there are clusters of movie raters.

```
h_2 <- dist(t(x)) %>% hclust()
plot(h_2, cex = 0.35)
```

Cluster Dendrogram





We can also use the k-means algorithm. Here, we have to pre-define k, the number of clusters we want to define.

The k-means algorithm is iterative. The first step is to define k centers. Then each observation is assigned to the cluster with the closest center to that observation. In a second step the centers are redefined using the observation in each cluster: the column means are used to define a **centroid**. We repeat these two steps until the centers converge.





The kmeans function included in R-base does not handle NAs. For illustrative purposes we will fill out the NAs with 0s. In general, the choice of how to fill in missing data, or if one should do it at all, should be made with care.

```
x_0 <- x; x_0[is.na(x_0)] <- 0
set.seed(0)
k <- kmeans(x_0, centers = 10)</pre>
```





The cluster assignments are in the cluster component:

```
groups <- k$cluster
table(groups)</pre>
```

```
## groups
## 1 2 3 4 5 6 7 8 9 10
## 1 9 3 3 4 3 3 5 6 13
```





[1] "Aladdin"

[4] "E.T. the Extra-Te..." "Forrest Gump"

This yields some interesting groups:

"Apollo 13"

```
names(groups) [groups==4]
## [1] "Ace Ventura: Pet ..." "Dumb & Dumber (Du..." "Mask. The"
names(groups)[groups==6]
## [1] "Lord of the Rings..." "Lord of the Rings..." "Lord of the Rings..."
names(groups)[groups==7]
## [1] "Star Wars IV - A ..." "Star Wars V - The..." "Star Wars VI - Re..."
names(groups)[groups==9]
## [1] "Braveheart"
                              "Dances with Wolves"
                                                      "Godfather, The"
## [4] "Good Will Hunting"
                              "Schindler's List"
                                                      "Shawshank Redempt..."
names(groups) [groups==10]
```

"Beauty and the Beast"

"Gladiator"





Note that because the first center is chosen at random, the final clusters are random. We impose some stability by repeating the entire function several times and averaging the results. The number of random starting values to use can be assigned through the nstart argument.

 $k \leftarrow kmeans(x 0, centers = 10, nstart = 25)$





Heatmaps

A powerful visualization tool for discovering clusters or patterns in your data is the heatmap. The idea is simple: plot an image of your data matrix with colors used as the visual cue and both the columns and rows ordered according to the results of a clustering algorithm. We will demonstrate this with the tissue_gene_expression dataset. We will scale the rows of the gene expression matrix.

The first step is compute:

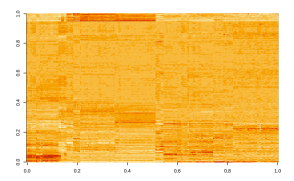




Heatmaps

Now we can use the results of this clustering to order the rows and columns.

```
image(x[h_1$order, h_2$order])
```



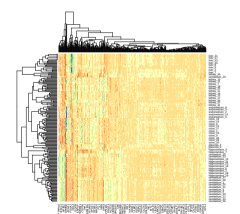




Heatmaps

But there is heatmap function that does it for us:

```
heatmap(x, col = RColorBrewer::brewer.pal(11, "Spectral"))
```







Filtering features

If the information about clusters in included in just a few features, including all the features can add enough noise that detecting clusters becomes challenging. One simple approach to try to remove features with no information is to only include those with high variance. In the movie example, a user with low variance in their ratings is not really informative: all the movies seem about the same to them.





Filtering features

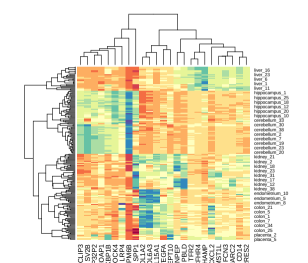
For example, if we include only features (genes) with highest variance.

```
library(matrixStats)
sds <- colSds(x, na.rm = TRUE)
o <- order(sds, decreasing = TRUE)[1:25]
heatmap(x[,o],
    col = RColorBrewer::brewer.pal(11, "Spectral"))</pre>
```





Filtering features







Session Info

sessionInfo()

```
## R version 4.5.1 (2025-06-13)
## Platform: aarch64-apple-darwin20
## Running under: macOS Sequoia 15.5
##
## Matrix products: default
          /Library/Frameworks/R.framework/Versions/4.5-arm64/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.5-arm64/Resources/lib/libRlapack.dylib: LAPACK version 3.12.1
##
## locale:
## [1] en US.UTF-8/en US.UTF-8/en US.UTF-8/C/en US.UTF-8/en US.UTF-8
##
## time zone: America/New York
## tzcode source: internal
##
## attached base packages:
## [1] stats
                graphics grDevices utils
                                               datasets methods
                                                                   base
## other attached packages:
    [1] matrixStats 1.5.0 dslabs 0.8.0
                                            lubridate 1.9.4 forcats 1.0.0
    [5] stringr_1.5.1
                         dplyr_1.1.4
                                            purrr_1.1.0
                                                              readr_2.1.5
    [9] tidyr_1.3.1
                         tibble 3.3.0
                                            tidvverse 2.0.0 caret 7.0-1
## [13] lattice 0.22-7
                          ggplot2 3.5.2
## loaded via a namespace (and not attached):
```

