Spectral Clustering Notes

Much of this uses information from A Tutorial on Spectral Clustering by Ulrike von Luxburg and Proximity in Statistical Machine Learning by Michael Trosset.

The Ratio Cut Problem

Ratio Cut for k=2

It can be shown that in the relaxed case for k=2, minimizing:

$$W(k) = \sum_{i=1}^{k} (x_i - m_1)^2 + \sum_{i=k+1}^{n} (x_i - m_2)^2$$

where m_1 and m_2 are k-means centers, as perscribed by Luxburg results in the same clustering as by assigning clusters by minimizing

$$R(k) = \sum_{i=1}^{k} \left(x_i + \sqrt{\frac{n-k}{k}} \right)^2 + \sum_{i=k+1}^{n} \left(x_i - \sqrt{\frac{k}{n-k}} \right)^2$$

in \mathbb{R}^1 and where x_i s are ordered, i.e., $x_i \leq \cdots \leq x_n$. We also constrain the problem to $\sum_{i=1}^{n} x_i = 0$ and $\sum_{i=1}^{n} x_i^2 = n$.

The k-means centers in \mathbb{R} are

$$m_1 = \frac{1}{k} \sum_{i=1}^k x_i$$

$$m_2 = \frac{1}{n-k} \sum_{i=k+1}^n x_i$$

Note that m_1 and m_2 are functions of k.

Numerical results

Using an arbitrary vector $\overrightarrow{x} \in \mathbb{R}^n$ such that $\sum_i x_i = 0$ and $|\overrightarrow{x}|_2^2 = n$:

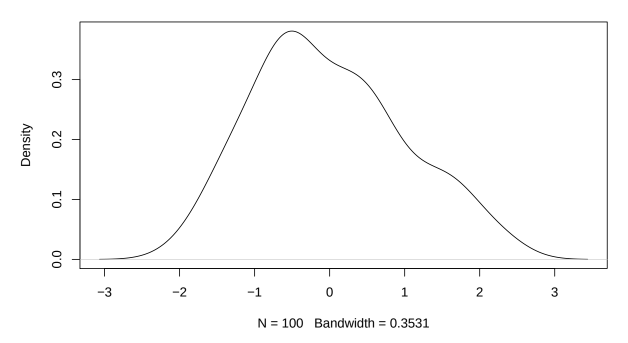
```
# packages
library(ggplot2)
import::from(magrittr, `%>%`, `%<>%`)
theme_set(theme_bw())
import::from(psych, tr)

# --- functions --- #

normalize <- function(x) {
   y <- x - mean(x)</pre>
```

```
z <- y / sqrt(mean(y ** 2))
 return(z)
k.means <- function(x) {
  x <- sort(x)
 n <- length(x)
  sapply(seq(n - 1), function(k) {
    m1 \leftarrow 1 / k * sum(x[seq(k)])
    m2 \leftarrow 1 / (n - k) * sum(x[seq(k + 1, n)])
    W \leftarrow sum((x[seq(k)] - m1)^2) + sum((x[seq(k + 1, n)] - m2)^2)
    return(W)
 })
ratio.cut <- function(x) {</pre>
  x <- sort(x)
  n <- length(x)</pre>
  sapply(seq(n - 1), function(k) {
    Ac \leftarrow sqrt((n - k) / k)
    A <- sqrt(k / (n - k))
    R \leftarrow sum((x[seq(k)] + Ac)^2) + sum((x[seq(k + 1, n)] - A)^2)
    return(R)
  })
# generate random data
n <- 100
k < -4
x \leftarrow c(rnorm(n / k, -1),
       rnorm(n / k, 0),
       rnorm(n / k, 1),
       rnorm(n / k, 3))
z <- normalize(x)</pre>
plot(density(z))
```

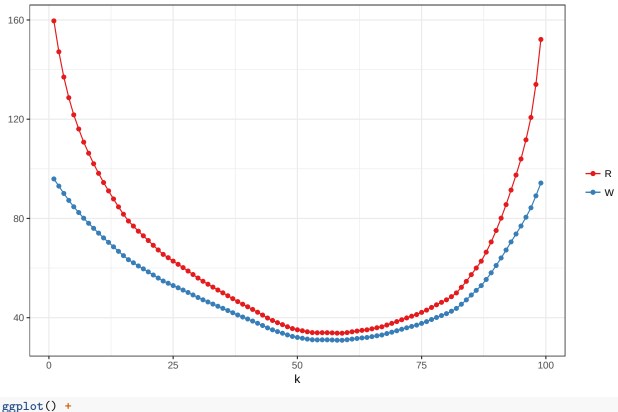
density.default(x = z)

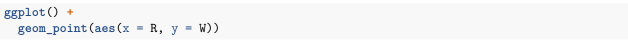


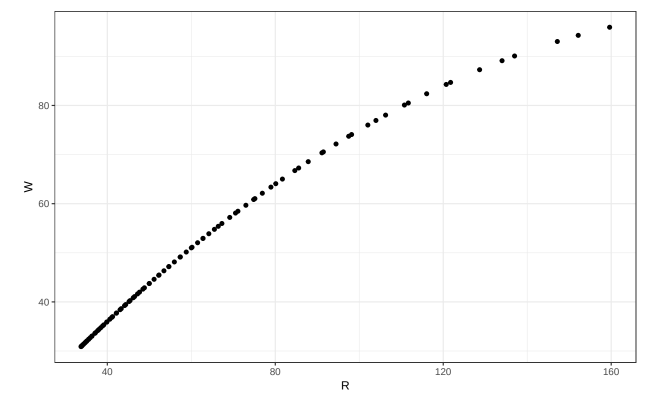
```
# compute W and R
W <- k.means(z)
R <- ratio.cut(z)

# visualizations
k <- seq(n - 1)

ggplot() +
    geom_point(aes(x = k, y = W, colour = 'W')) +
    geom_line(aes(x = k, y = W, colour = 'W')) +
    geom_point(aes(x = k, y = R, colour = 'R')) +
    geom_line(aes(x = k, y = R, colour = 'R')) +
    scale_colour_brewer(palette = 'Set1') +
    labs(x = 'k', y = NULL, colour = NULL)</pre>
```

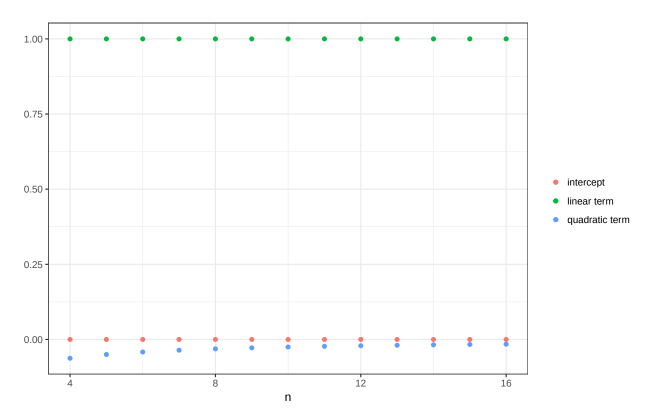






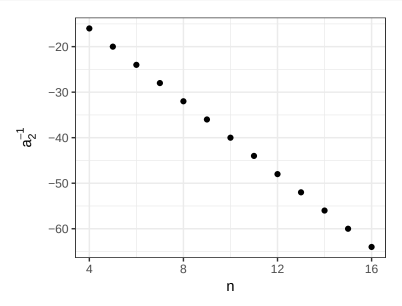
It appears that there is a definite relationship between W and R. Using quadratic regression:

```
summary(lm(W ~ R + I(R ** 2)))
Call:
lm(formula = W \sim R + I(R^2))
Residuals:
       Min
                   1Q
                          Median
                                          3Q
                                                     Max
-7.590e-14 -3.835e-15 -1.480e-16 3.780e-15 1.057e-13
Coefficients:
              Estimate Std. Error
                                      t value Pr(>|t|)
(Intercept) 4.570e-14 8.461e-15 5.401e+00 4.79e-07 ***
             1.000e+00 2.384e-16 4.195e+15 < 2e-16 ***
            -2.500e-03 1.413e-18 -1.769e+15 < 2e-16 ***
I(R^2)
___
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.498e-14 on 96 degrees of freedom
                         1, Adjusted R-squared:
Multiple R-squared:
F-statistic: 7.474e+31 on 2 and 96 DF, p-value: < 2.2e-16
\dots we get a perfect fit. We can try several values of the data size n:
N \leftarrow 2 ** 4 # number of obs to try
coefs.df <- lapply(seq(4, N), function(n) {</pre>
 x <- normalize(rnorm(n)) # generate data
  # compute results
  W <- k.means(x)
  R <- ratio.cut(x)</pre>
  # compute coefs for quadratic equation
  quad.coefs <- lm(W \sim R + I(R ** 2))$coefficients
  a0 <- quad.coefs['(Intercept)']</pre>
  a1 <- quad.coefs['R']
  a2 <- quad.coefs['I(R^2)']
  # compile into data frame
  dplyr::data_frame(n, a0, a1, a2)
}) %>%
  dplyr::bind_rows()
ggplot(coefs.df) +
  geom_point(aes(x = n, y = a0, colour = 'intercept')) +
  geom_point(aes(x = n, y = a1, colour = 'linear term')) +
  geom_point(aes(x = n, y = a2, colour = 'quadratic term')) +
  labs(colour = NULL, y = NULL)
```



The intercept term stays at 0 and the linear term stays at 1. Looking closer at the quadratic term:

```
ggplot(coefs.df) +
  geom_point(aes(x = n, y = a2 ** -1)) +
  labs(y = expression(a[2]^-1))
```



Then we arrive at the result $W = R - \frac{1}{4n}R^2$.

Analytic result

We can show:

$$W(k) = R(k) - \frac{(R(k))^2}{4n}$$

or $W(R) = R - \frac{1}{4n}R^2$. This function is strictly increasing for $R(k) \le 2n$. Expanding R(k), we get:

$$R(k) = 2n + 2\sqrt{\frac{n-k}{k}} \sum_{i=1}^{k} x_i - 2\sqrt{\frac{k}{n-k}} \sum_{i=k+1}^{n} x_i$$

It can be shown that R(k) is maximized at the endpoints.

Note that since $\sum_{i=1}^{n} x_i = 0$, $\sum_{i=1}^{k < n} x_i \le 0$, $x_n \ge 0$, and $x_i \le 0$.

k can range from 1 to n-1. If k=n-1, we get $R(n-1)=2n+2\sqrt{\frac{1}{k}}\sum_{i}^{n-1}x_{i}-2\sqrt{n-1}x_{n}$. The second term is ≤ 0 and the third term is ≥ 0 , so we get $R \leq 2n$. On the other hand, if k=1, $R(1)=2n+2\sqrt{n-1}x_{1}-\frac{2}{\sqrt{n-1}}\sum_{i=2}^{n}x_{i}=2n+2\sqrt{n-1}x_{1}-\frac{2}{\sqrt{n-1}}(-x_{1})=2n+x_{1}\left(2\sqrt{n-1}+\frac{2}{\sqrt{n-1}}\right)\leq 2n$.

Expanding W(k), we get:

$$W(k) = \sum_{i=1}^{k} x_i^2 - 2m_1 \sum_{i=1}^{k} x_i + km_1^2 + \sum_{i=k+1}^{n} x_i^2 - 2m_2 \sum_{i=k+1}^{n} x_i + m_2^2(n-k)$$

$$= n - \frac{2}{k} \left(\sum_{i=1}^{k} x_i\right)^2 + \frac{1}{k} \left(\sum_{i=1}^{k} x_i\right)^2 - \frac{2}{n-k} \left(\sum_{i=k+1}^{n} x_i\right)^2 + \frac{1}{n-k} \left(\sum_{i=k+1}^{n} x_i\right)^2$$

$$= n - \frac{1}{k} \left(\sum_{i=1}^{k} x_i\right)^2 - \frac{1}{n-k} \left(\sum_{i=k+1}^{n} x_i\right)^2$$

$$= n - km_1^2 - (n-k)m_2^2$$

Since $\sum_{i=1}^{n} x_i = 0$, $km_1 + (n-k)m_2 = 0$, or, $-nm_2 = k(m_1 - m_2)$. Then

$$W(k) = n - km_1^2 - (n - k)m_2^2$$

$$= n - km_1^2 - nm_2^2 + km_2^2$$

$$= n - km_1^2 + (nm_2)m_2 + km_2^2$$

$$= n - km_1 + k(m_1 - m_2)m_2 + km_2^2$$

$$= n + k(-m_1^2 + m_1m_2 - m_2^2 + m_2^2)$$

$$= n + km_1(m_2 - m_1)$$

Using the relationship $-nm_2 = k(m_1 - m_2) \implies m_2 - m_1 = \frac{nm_2}{k}$ from before, we can again rewrite W(k):

$$W(k) = n - (n - k)m_2 \frac{nm_2}{k}$$
$$= n \frac{(n - k)n}{k} m_2^2$$

And we use the same relationship again: $m_2 - m_1 = \frac{n}{k} m_2 \implies m_2 = (m_2 - m_1) \frac{k}{n}$.

Then we can finally write W(k) as:

$$W(k) = n - \frac{(n-k)n}{k} \frac{k^2}{n^2} (m_1 - m_2)^2$$
$$W(k) = n - \frac{(n-k)k}{n} (m_1 - m_2)^2$$

On the other hand, if we expand R(k):

$$R(k) = \sum_{i=1}^{k} x_i^2 + 2\sqrt{\frac{n-k}{k}} \sum_{i=1}^{k} x_i + n - k + \sum_{k=1}^{n} x_i^2 - 2\sqrt{\frac{k}{n-k}} \sum_{k=1}^{n} x_i + k$$
$$= 2n + 2\sqrt{k(n-k)}m_1 - 2\sqrt{k(n-k)}m_2$$

If we expand and simplify $-\frac{(R(k))^2}{4n}$, we get:

$$-\frac{(R(k))^2}{4n} = -n - \frac{k(n-k)}{n}m_1^2 - \frac{k(n-k)}{n}m_2^2 - 2m_1\sqrt{k(n-k)} + 2m_2\sqrt{k(n-k)} + \frac{2k(n-k)}{n}m_1m_2$$

Then noting that some terms cancel each other out, $R(k) - \frac{(R(k))^2}{4n}$:

$$n - \frac{k(n-k)}{n}m_1^2 - \frac{k(n-k)}{n}m_2^2 + 2\frac{k(n-k)}{n}m_1m_2$$
$$= n - \frac{k(n-k)}{n}(m_1^2 + m_2^2 - 2m_1m_2)$$
$$= n - \frac{k(n-k)}{n}(m_1 - m_2)^2$$

Which is exactly the same as our expression for W(k).

Arbitrary k

For arbitrary k, the methods as prescribed by Luxburg involve embedding the graph to \mathbb{R}^k and then performing k-means clustering. In this case, we cannot perform the same sort of analysis since there is no way to "order" the x_i 's.

Numerical experiments for k=2 and \mathbb{R}^2

Double spiral

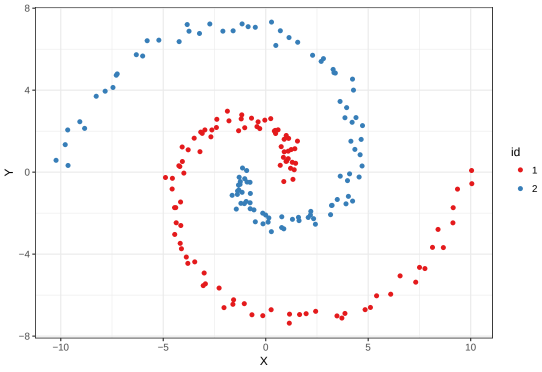
We will generate a "double spiral" to be partitioned into two clusters. This is an example that k-means would fail but is fairly distinguishable visually or intuitively. After generating the spiral, we will construct a k-nearest neighbors graph.

```
# borrow some functions from S675
source('http://pages.iu.edu/~mtrosset/Courses/675/manifold.r')
# parameters
set.seed(112358)
s <- 2 ** 5
eps <- 2 ** -2
k <- 10 # for constructing the knn graph
K <- 2 # number of clusters
cols2 <- colorRampPalette(c('blue', 'white', 'red'))(256)</pre>
rad.max <- 10
ang.max <- 2 * pi
angles <- seq(0, ang.max, length.out = 100)</pre>
radii <- seq(1, sqrt(rad.max), length.out = 100) ** 2</pre>
N \leftarrow 100 # number of times to try k-means clustering
# data
spiral.df <- dplyr::data_frame(X = radii * cos(angles),</pre>
                                Y = radii * sin(angles))
spiral.df <- dplyr::data_frame(X = radii * cos(angles),</pre>
                                 Y = radii * sin(angles))
neg.spiral.df <- dplyr::mutate(spiral.df,</pre>
                                 X = -X, Y = -Y,
                                 id = '2')
spiral.df %<>%
  dplyr::mutate(id = '1') %>%
  dplyr::bind_rows(neg.spiral.df) %>%
  dplyr::mutate(X = X + rnorm(n = n(), sd = eps),
                 Y = Y + rnorm(n = n(), sd = eps))
n <- nrow(spiral.df) # number of vertices</pre>
# viz
ggplot(spiral.df) +
  geom_point(aes(x = X, y = Y)) +
  coord_fixed() +
 labs(title = 'Double spiral')
```

Double spiral -4 -4 -10 -5 0 5 10

```
ggplot(spiral.df) +
geom_point(aes(x = X, y = Y, colour = id)) +
coord_fixed() +
scale_colour_brewer(palette = 'Set1') +
labs(title = '"Natural" clustering of the example data')
```

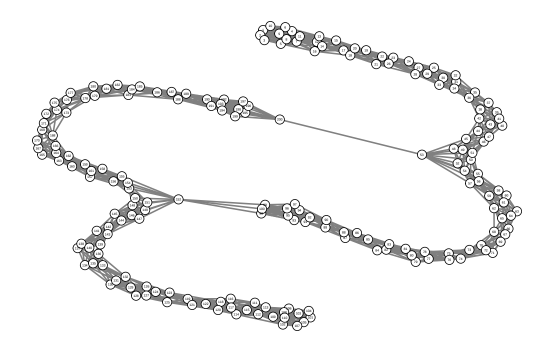
"Natural" clustering of the example data



Then we will construct a(n) 10-nearest-neighbors graph:

```
# construct W
W <- spiral.df %>%
    dplyr::select(X, Y) %>%
    as.matrix() %>%
    mds.edm1() %>%
    graph.knn(k) %>%
    graph.adj()

# viz
qgraph::qgraph(W)
```

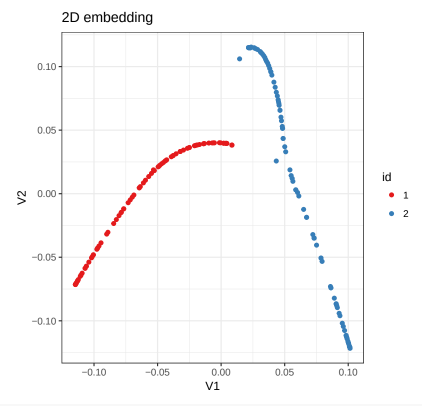


Here it is pretty obvious how we should cut the graph.

Then proceed with the clustering method as described by Luxburg:

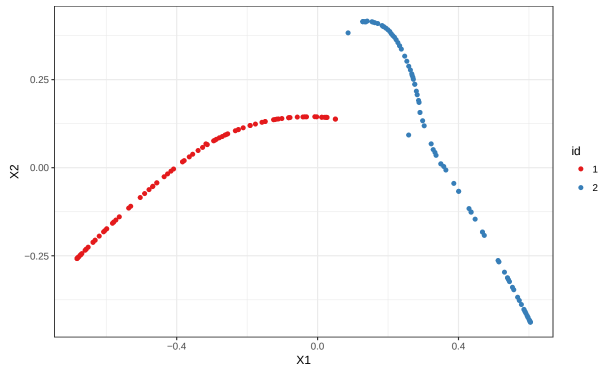
Then we can take a look at the projection to \mathbb{R}^2 :

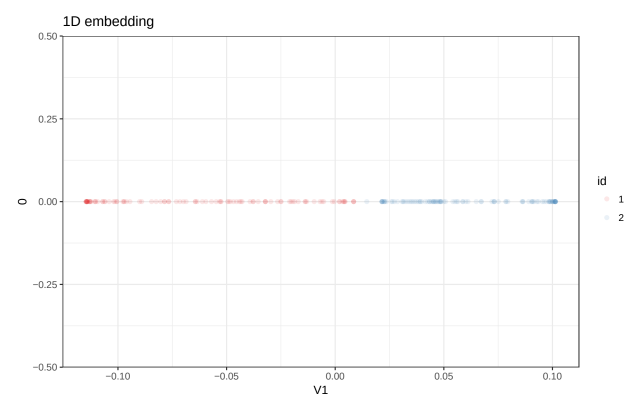
```
ggplot(two.eigen.df) +
geom_point(aes(x = V1, y = V2, colour = id)) +
coord_fixed() +
scale_colour_brewer(palette = 'Set1') +
labs(title = '2D embedding')
```



```
ggplot(two.eigen.df) +
  geom_point(aes(x = X1, y = X2, colour = id)) +
  coord_fixed() +
  scale_colour_brewer(palette = 'Set1') +
  labs(title = '2D embedding, scaled by eigenvalues')
```

2D embedding, scaled by eigenvalues





Next, we can take a look at how the three methods correspond to our original labels:

```
# run each clustering method N times and pick the best one
multiple.clusterings <- function(input.df, columns, clusters = 2, iter = 100) {</pre>
  # run the k-means algorithm many times
  clusterings <- lapply(seq(iter), function(i) {</pre>
    input.df %>%
      dplyr::select(columns) %>%
      dist() %>%
      kmeans(clusters)
  })
  # see which time was best
  tot.withinss <- sapply(clusterings, function(clustering) {</pre>
    clustering$tot.withinss
  })
  best.clust.ind <- which.min(tot.withinss)[1]</pre>
  return(clusterings[[best.clust.ind]])
}
unscaled.clustering <- multiple.clusterings(two.eigen.df, c('V1', 'V2'), K, N)
scaled.clustering <- multiple.clusterings(two.eigen.df, c('X1', 'X2'), K, N)</pre>
one.d.clustering <- multiple.clusterings(two.eigen.df, 'X1', K, N)
spiral.df %<>%
  dplyr::mutate(unscaled = as.character(unscaled.clustering$cluster),
                 scaled = as.character(scaled.clustering$cluster),
                 one.d = as.character(one.d.clustering$cluster))
```

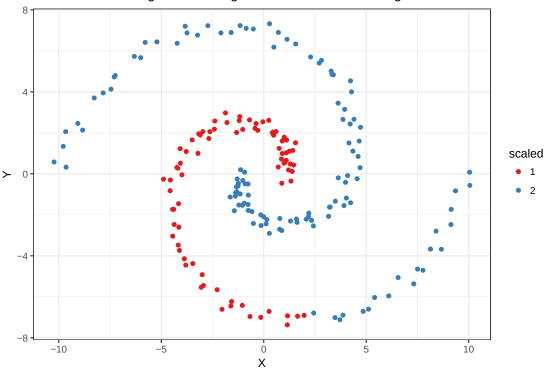
table(spiral.df\$id, spiral.df\$unscaled) 2 1 1 100 2 60 40 table(spiral.df\$id, spiral.df\$scaled) 2 1 80 20 0 100 table(spiral.df\$id, spiral.df\$one.d) 2 1 1 19 81 2 100 ggplot(spiral.df) + geom_point(aes(x = X, y = Y, colour = unscaled)) + coord_fixed() + scale_colour_brewer(palette = 'Set1') + labs(title = 'k-means clustering results using the unscaled 2D embedding') k-means clustering results using the unscaled 2D embedding unscaled

```
ggplot(spiral.df) +
geom_point(aes(x = X, y = Y, colour = scaled)) +
coord_fixed() +
```

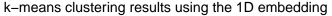
ó X

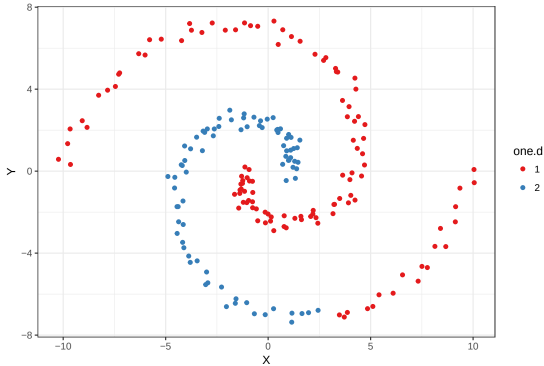
```
scale_colour_brewer(palette = 'Set1') +
labs(title = 'k-means clustering results using the scaled 2D embedding')
```

k-means clustering results using the scaled 2D embedding



```
ggplot(spiral.df) +
geom_point(aes(x = X, y = Y, colour = one.d)) +
coord_fixed() +
scale_colour_brewer(palette = 'Set1') +
labs(title = 'k-means clustering results using the 1D embedding')
```





In a way, we have "unraveled" the spirals, resulting in two curves. Eye-balling this representation gives us two obvious clusters, but k-means fails to separate them the way we would expect since they are not spherical clusters. Projection to the unscaled space seems to perform especially poorly, and the plots of the columns of H_{approx} reveal why.

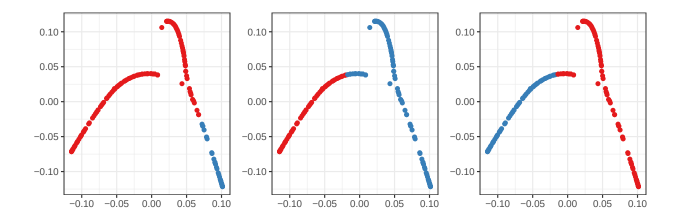


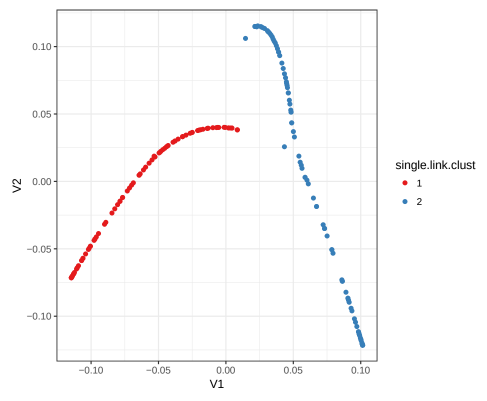
Figure 1: 2D embedding annotated by cluster assignments. From right to left: k-means on the unscaled embedding, k-means on the scaled embedding, and k-means on the 1D embedding.

In fact, it almost looks as if another clustering method is more appropriate here, in particular, single-linkage.

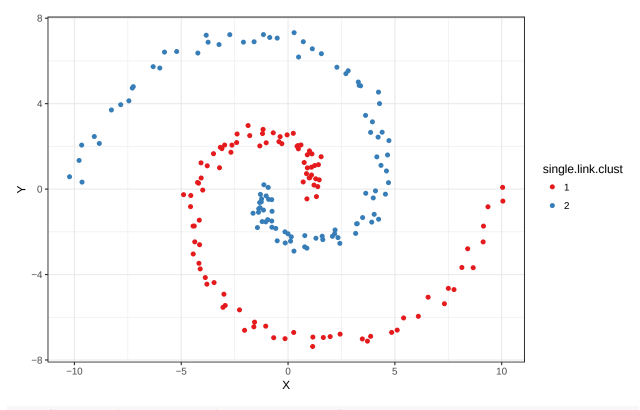
```
single.link.clust <- two.eigen.df %>%
    dplyr::select(V1, V2) %>%
    dist() %>%
    hclust(method = 'single') %>%
    cutree(2)

two.eigen.df %<>%
    dplyr::mutate(single.link.clust = as.character(single.link.clust))
spiral.df %<>%
    dplyr::mutate(single.link.clust = as.character(single.link.clust))

ggplot(two.eigen.df) +
    geom_point(aes(x = V1, y = V2, colour = single.link.clust)) +
    coord_fixed() +
    scale_colour_brewer(palette = 'Set1')
```



```
ggplot(spiral.df) +
geom_point(aes(x = X, y = Y, colour = single.link.clust)) +
coord_fixed() +
scale_colour_brewer(palette = 'Set1')
```



table(spiral.df\$id, spiral.df\$single.link.clust)

```
1 2
1 100 0
2 0 100
```

We should also check that this clustering is actually better according to the metrics from the Luxburg tutorial. We try this two ways:

- 1. H_{approx} should be as close of an approximation to H as possible. Here, H_{approx} is the k-dimensional embedding via spectral decomposition while H defines the actual best clustering. In this comparison, we fix H_{approx} and construct H according to the resulting clustering of H_{approx} (e.g., via k-means or single-linkage), then compare the constructed H to H_{approx} via the Frobenious norm.
- 2. The exact clustering should minimize $Tr(H^TLH)$ where H is defined by the particular clustering. If single-linkage is indeed better than k-means in this case, then the H constructed from single-linkage clustering should result in a lower metric than the one constructed from k-means clustering.

```
# construct Happrox
H.approx <- two.eigen.df %>%
    dplyr::select(V2, V1) %>% # i put this in the wrong order
    as.matrix()

#' @title construct H function
#' @description H is constructed based on a vector that designates the clusters
#' @param clustering (numeric) A vector of cluster assignments
#' @return (matrix) H based on the cluster assignments
construct.H <- function(clustering) {
    # this function is limited to nonempty clusters
    # e.g., if there are 3 clusters, they must be assigned as 1, 2, 3</pre>
```

```
clusters <- unique(clustering)</pre>
  if (length(clusters) != max(clustering)) {
    stop(simpleError('there are empty clusters'))
  }
  if (min(clustering) < 1) {</pre>
    stop(simpleError('cluster indexing starts at 1'))
  # find |A_k|
  cluster.sizes <- sapply(clusters, function(i) {</pre>
    length(clustering[clustering == i])
  })
  # construct H
  H <- sapply(clustering, function(i) {</pre>
   h <- rep(0, length(clusters))</pre>
    h[i] <- 1 / sqrt(cluster.sizes[i])</pre>
    return(h)
 }) %>%
    t()
 return(H)
}
# construct H for k-means (unscaled)
H.kmeans <- construct.H(unscaled.clustering$cluster)</pre>
# construct H for single linkage
H.singlelink <- construct.H(single.link.clust)</pre>
# frobenius norm comparison
norm(H.approx - H.kmeans, type = 'F')
[1] 1.497036
norm(H.approx - H.singlelink, type = 'F')
[1] 1.704416
# making sure ordering doesn't matter
\# this could get messy for large k
norm(H.approx - H.kmeans[, 2:1], type = 'F')
[1] 2.400969
norm(H.approx - H.singlelink[, 2:1], type = 'F')
[1] 2.257204
# minimizing trace comparison
# here order doesn't matter
tr(t(H.kmeans) %*% L %*% H.kmeans)
[1] 0.46875
tr(t(H.singlelink) %*% L %*% H.singlelink)
[1] 0.08
```

```
# trace for Happrox
tr(t(H.approx) %*% L %*% H.approx)
```

[1] 0.1049757

Here, we can see that single linkage produces a H matrix that is closer to H_{approx} and results in a better clustering metric.

For completeness, let's check whether single-linkage produced a better clustering according to the k-means metric:

```
# function for computing total within SS
within.ss <- function(X, clustering) {
   clusters <- unique(clustering)
   sapply(clusters, function(i) {
     X.clust <- X[clustering == i, ]
     sum(dist(X.clust) ** 2) / 2 / nrow(X.clust)
}) %>%
   sum()
}

# total within-ss from k-means
within.ss(dplyr::select(two.eigen.df, V1, V2), unscaled.clustering$cluster)
```

[1] 0.5689028

```
# total within-ss from single linkage
within.ss(dplyr::select(two.eigen.df, V1, V2), single.link.clust)
```

[1] 0.6209851

And indeed, the k-means clustering result resulted in a better k-means metric. This conclusively tells us that k-means on the embedding is not the best clustering method for this example.

However, this may be a very particular case. From the embedding, we can see that while there is a very obvious clustering, the clusters are not ellipsoids, which is what k-means is good at. Perhaps in most graph embeddings of this type, we do end up with ellpsoid clusters, and we just happened to get long strands in this example.