# Lab 08

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## Import the Data

```
url <- "https://tinyurl.com/UK-foods"
uk_foods <- read.csv(url, row.names = 1)</pre>
```

## **Explore Imported Data**

Question 1: How many rows and columns are in your new data frame named x? What R functions could you use to answer this questions?

```
dim(uk_foods)
```

```
## [1] 17 4
```

17 rows and 4 columns (17 food categories and 4 countries)

# Checking the Data

#### head(uk foods)

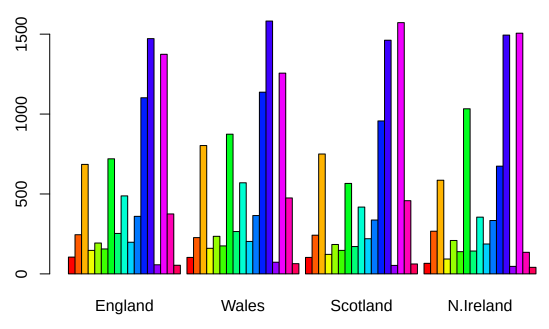
##	England	Wales	${\tt Scotland}$	${\tt N.Ireland}$
## Cheese	105	103	103	66
## Carcass_meat	245	227	242	267
## Other_meat	685	803	750	586
## Fish	147	160	122	93
## Fats_and_oils	193	235	184	209
## Sugars	156	175	147	139

Question 2: Which approach to solving the 'row-names problem' mentioned above do you prefer and why? Is one approach more robust than another under certain circumstances?

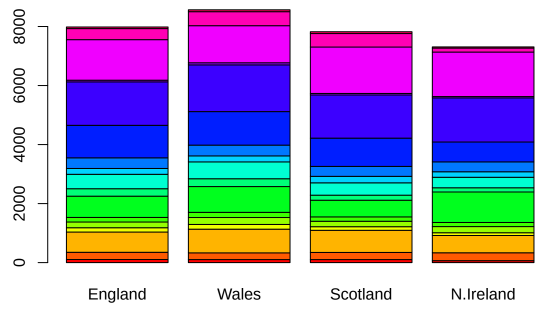
I prefer to read in the row-names when I read in the csv data (in *read.csv*). This is less dangerous since it does not mutate our data in the body of the code, and instead restricts it to when the data is read in.

# Looking for differences and trends

```
barplot(as.matrix(uk_foods), beside = T, col = rainbow(nrow(uk_foods)))
```



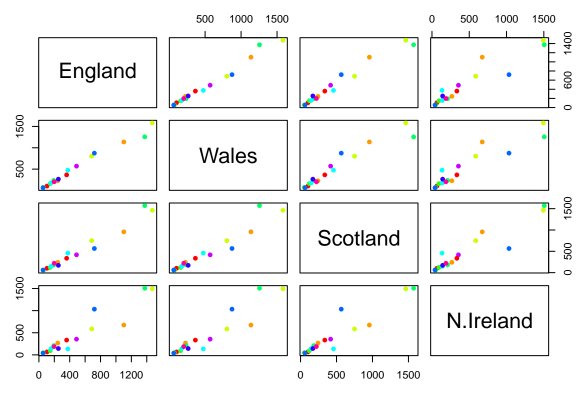
Question 3: Changing what optional argument in the above barplot() function results in the following plot?



Changing the  ${\tt beside}$  argument from TRUE to FALSE

Question 5: Generating all pairwise plots may help somewhat. Can you make sense of the following code and resulting figure? What does it mean if a given point lies on the diagonal for a given plot?

pairs(uk\_foods, col=rainbow(10), pch=16)



This code plots all pairwise comparisons of row values across the column groups. If a given point lies on the diagonal, this means that the value of the corresponding row is the same in both columns which are plotted. A point above the diagonal would be higher in the column plotted on the y-axis than in the column plotted on the x-axis. A point below the diagonal would be higher in the column plotted on the x-axis than in the column plotted on the y-axis.

Question 6: What is the main differences between N. Ireland and the other countries of the UK in terms of this data-set?

#### Without PCA

This is difficult to do without using a PCA, but a way could be to look at the maximum differences, pairwise, in log2 fold-changes between Northern Ireland and the other UK countries.

Identify the log2 fold change and identify the component with the maximum absolute change

```
n_ireland_v_england <- log(uk_foods$N.Ireland, base = 2)/log(uk_foods$England, base = 2)
n_ireland_v_wales <- log(uk_foods$N.Ireland, base = 2)/log(uk_foods$Wales, base = 2)
n_ireland_v_scotland <- log(uk_foods$N.Ireland, base = 2)/log(uk_foods$Scotland, base = 2)
names(n_ireland_v_england) <- row.names(uk_foods)
names(n_ireland_v_wales) <- row.names(uk_foods)
names(n_ireland_v_scotland) <- row.names(uk_foods)
which.max(abs(n_ireland_v_england))
## Fresh_potatoes
## 7
which.max(abs(n_ireland_v_wales))
## Carcass_meat
## 2</pre>
```

```
which.max(abs(n_ireland_v_scotland))
## Fresh_potatoes
## 7
```

## **PCA**

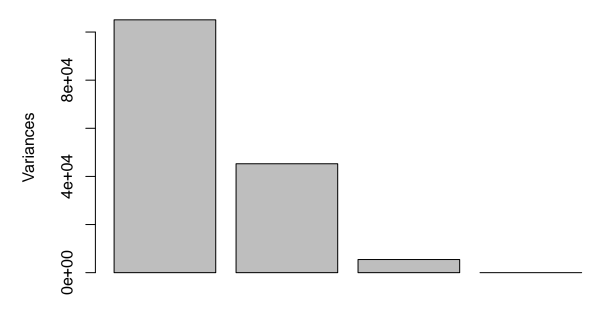
#### Conduct PCA with the UKFoods data set

```
uk_foods_pca <- prcomp(t(uk_foods))</pre>
summary(uk_foods_pca)
## Importance of components:
                                         PC2
                                                             PC4
##
                                PC1
                                                  PC3
## Standard deviation
                           324.1502 212.7478 73.87622 3.176e-14
                             0.6744
                                      0.2905 0.03503 0.000e+00
## Proportion of Variance
## Cumulative Proportion
                             0.6744
                                      0.9650 1.00000 1.000e+00
```

### Inspect PCAs

plot(uk\_foods\_pca)

# uk\_foods\_pca

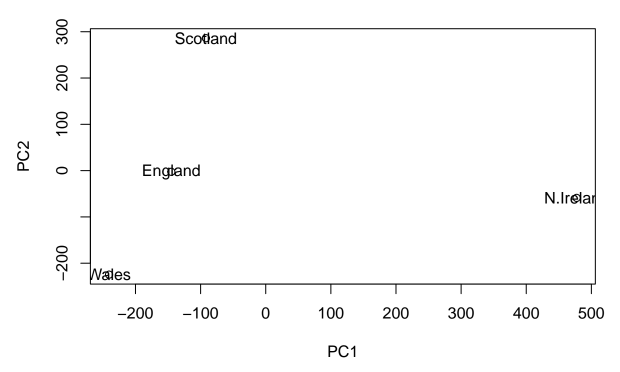


### Plot the data using PCs

Use the two PCs which explain the most variance to plot the UKFoods data set (PC1 and PC2)

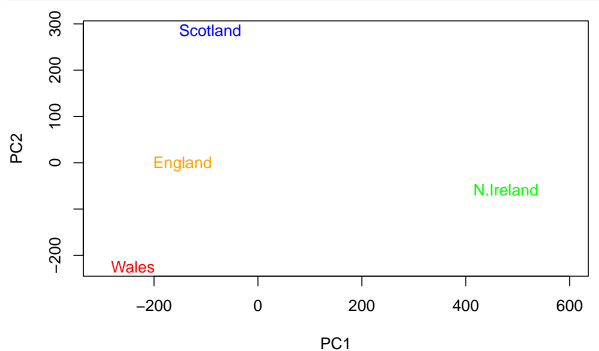
Question 7:

```
plot(uk_foods_pca$x[,1:2])
text(uk_foods_pca$x[,1:2], colnames(uk_foods))
```



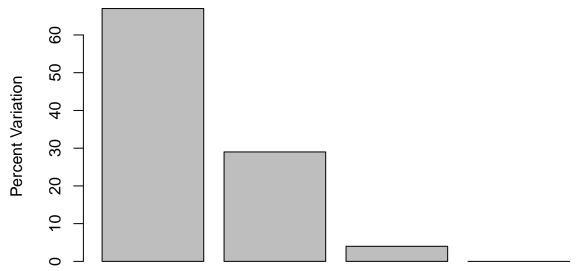
Question 8: Customize your plot so that the colors of the country names match the colors in our UK and Ireland map and table at start of this document.

```
color_vector <- c("orange", "red", "blue", "green")
plot(uk_foods_pca$x[,1:2], col = "#DDDDDD", xlim = c(-300, 600))
text(uk_foods_pca$x[,1:2], colnames(uk_foods), col = color_vector)</pre>
```



#### Plot the proportions of variences explained by each PC

```
variance_per <- round(((uk_foods_pca$sdev^2)/sum(uk_foods_pca$sdev^2)) * 100)
barplot(variance_per, xlab="Principal Component", ylab="Percent Variation")</pre>
```



# **Principal Component**

### Dig-

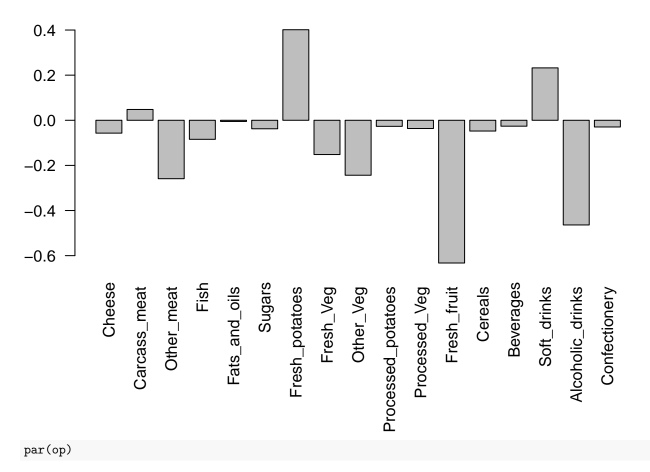
ging Deeper: Variable Loadings

uk\_foods\_pca\$rotation

```
##
                                PC1
                                             PC2
                                                         PC3
                                                                      PC4
                       -0.056955380
                                    0.016012850 0.02394295 -0.694538519
## Cheese
                                    0.013915823
## Carcass_meat
                        0.047927628
                                                 0.06367111
                                                              0.489884628
## Other_meat
                       -0.258916658 -0.015331138 -0.55384854
                                                              0.279023718
                       -0.084414983 -0.050754947 0.03906481 -0.008483145
## Fish
## Fats and oils
                       -0.005193623 -0.095388656 -0.12522257
                                                             0.076097502
## Sugars
                       -0.037620983 -0.043021699 -0.03605745
                                                             0.034101334
## Fresh_potatoes
                       0.401402060 -0.715017078 -0.20668248 -0.090972715
## Fresh_Veg
                       -0.151849942 -0.144900268 0.21382237 -0.039901917
## Other Veg
                       -0.243593729 -0.225450923 -0.05332841
                                                             0.016719075
## Processed_potatoes -0.026886233 0.042850761 -0.07364902 0.030125166
## Processed_Veg
                       -0.036488269 -0.045451802 0.05289191 -0.013969507
## Fresh_fruit
                       -0.632640898 -0.177740743 0.40012865 0.184072217
## Cereals
                       -0.047702858 -0.212599678 -0.35884921
                                                             0.191926714
## Beverages
                       -0.026187756 -0.030560542 -0.04135860
                                                             0.004831876
## Soft drinks
                        0.232244140
                                    0.555124311 -0.16942648
                                                              0.103508492
## Alcoholic_drinks
                                    0.113536523 -0.49858320 -0.316290619
                       -0.463968168
## Confectionery
                       -0.029650201
                                    0.005949921 -0.05232164 0.001847469
```

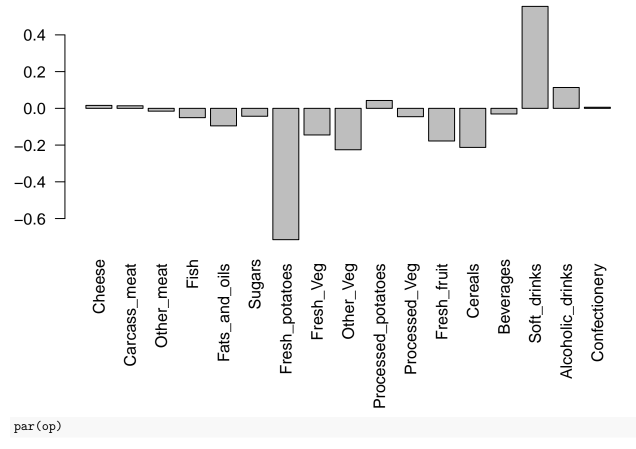
Since PC1 accounts for the most variance, the contribution of each component on the distribution of countries on this PC will be the most helpful.

```
op <- par(mar=c(10,3,0.35,0))
barplot(uk_foods_pca$rotation[,1], las = 2)</pre>
```



Question 9: Generate a similar 'loadings plot' for PC2. What two food groups feature prominantely and what does PC2 maninly tell us about?

```
op <- par(mar=c(10,3,0.35,0))
barplot(uk_foods_pca$rotation[,2], las = 2)</pre>
```



The two most prominent features are Fresh potatoes and soft drinks. This PC is mainly telling us which features most contribute to the differences between the other three UK countries (England to Scotland to Wales) since it is on this PC that these countries are differentiated upon. Therefore, these features (fresh potatoes and soft drinks) are the components which contribute to the differences among these three countries.

#PCA of RNA-Seq Data

## Import the Data

```
url2 <- "https://tinyurl.com/expression-CSV"</pre>
rna_data <- read.csv(url2, row.names=1)</pre>
head(rna_data)
##
           wt1 wt2
                    wt3
                          wt4 wt5 ko1 ko2 ko3 ko4 ko5
## gene1
          439
              458
                    408
                          429 420
                                    90
                                        88
                                            86
                                                 90
                                                     93
               200
   gene2
          219
                    204
                          210
                              187 427 423 434 433 426
         1006 989
                   1030
                         1017 973 252
                                       237 238 226 210
   gene3
                    829
                                       856 835
   gene4
           783 792
                          856
                              760 849
          181 249
                    204
                          244 225 277 305 272 270 279
  gene5
## gene6
          460 502
                    491
                          491 493 612 594 577 618 638
     Question 10: How many genes and samples are in this data set?
dim(rna_data)
```

## [1] 100 10

100 genes and 10 samples in the data set.

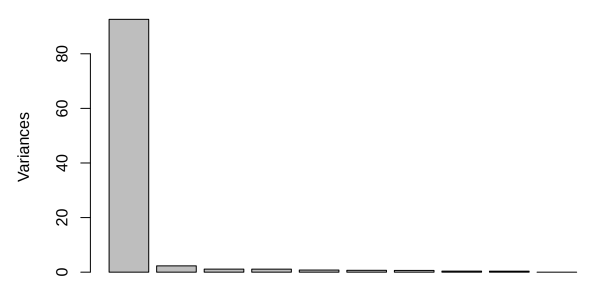
## PCA for RNA-Seq Data

```
rna_pca <- prcomp(t(rna_data), scale = TRUE)</pre>
summary(rna_pca)
## Importance of components:
                                             PC3
##
                             PC1
                                     PC2
                                                     PC4
                                                              PC5
                                                                      PC6
                                                                              PC7
                           9.6237 1.5198 1.05787 1.05203 0.88062 0.82545 0.80111
## Standard deviation
## Proportion of Variance 0.9262 0.0231 0.01119 0.01107 0.00775 0.00681 0.00642
## Cumulative Proportion 0.9262 0.9493 0.96045 0.97152 0.97928 0.98609 0.99251
                                       PC9
##
                               PC8
                                                PC10
## Standard deviation
                           0.62065 0.60342 3.457e-15
## Proportion of Variance 0.00385 0.00364 0.000e+00
## Cumulative Proportion 0.99636 1.00000 1.000e+00
```

Scree plot of the PCs helps to show how much variance is explained by each PC. Here it is clear that PC1 dominates compared to the others, explaining >92% of the total variance.

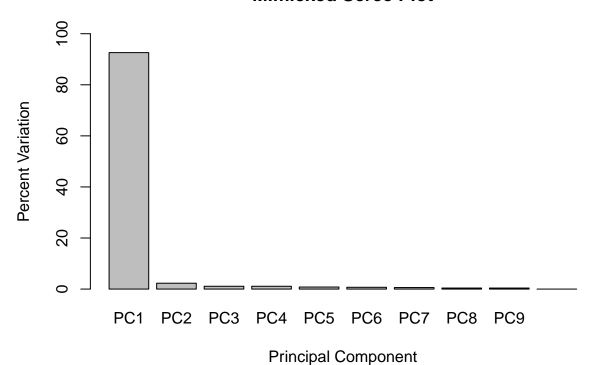
```
plot(rna_pca, main = "Scree plot")
```

# **Scree plot**



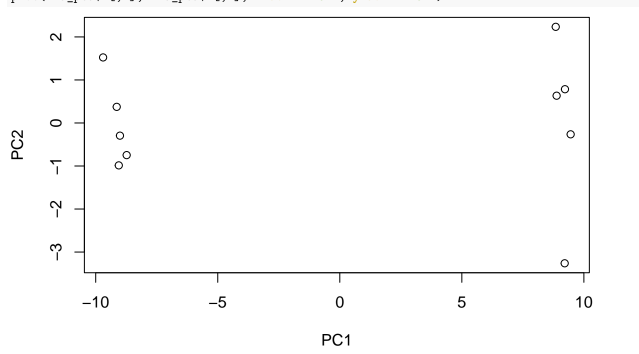
Replicate the Scree plot method for the pca object with base R plotting.

# **Mimicked Scree Plot**



Plot the data using PC1 and PC2  $\,$ 

```
plot(rna_pca$x[,1], rna_pca$x[,2], xlab = "PC1", ylab = "PC2")
```



Check if the groupings make sense

```
color_vector <- colnames(rna_data)
color_vector[grep("wt", color_vector)] <- "red"
color_vector[grep("ko", color_vector)] <- "blue"</pre>
```

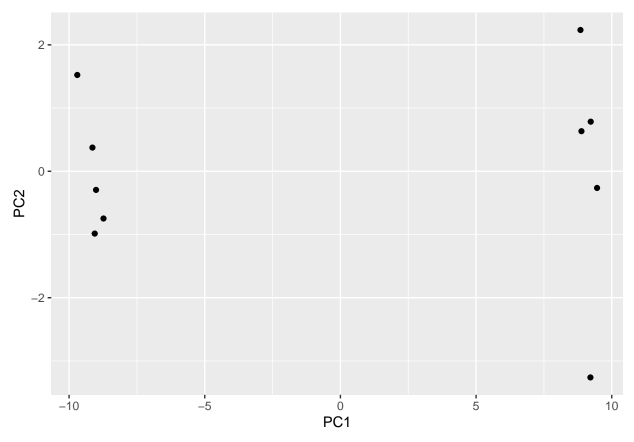
```
plot(rna_pca$x[,1], rna_pca$x[,2], col=color_vector, pch=16,
    xlab=paste0("PC1 (", rna_pca_varience_percent[1], "%)"),
    ylab=paste0("PC2 (", rna_pca_varience_percent[2], "%)"))
text(rna_pca$x[,1], rna_pca$x[,2], labels = colnames(rna_data), pos=c(rep(4,5), rep(2,5)))
                                                                        ko1 •
             wt1
                                                                        k895.
     0
                                                                          ko3 •
                                                                         ko2 •
          -10
                            -5
                                              0
                                                               5
                                                                                10
                                       PC1 (92.6%)
```

# Plotting PCA using ggplot

```
# Load the ggplot2 library
library(ggplot2)

# Convert PCA results to a data frame for ggplot
rna_pca_df <- as.data.frame(rna_pca$x)

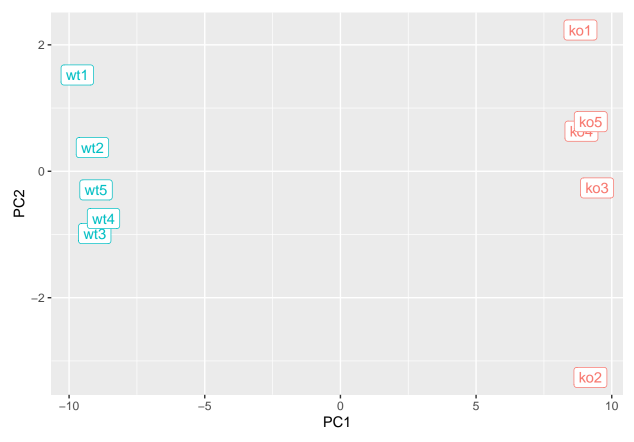
# Plot samples on PC1 and PC2
ggplot(rna_pca_df) +
   aes(x = PC1, y = PC2) +
   geom_point()</pre>
```



To make the figure clearer, plot the labels instead of just a point, and map color to sample condition.

```
# Create columns in the data frame for labeling samples and conditions
rna_pca_df$samples <- colnames(rna_data)
rna_pca_df$conditions <- substr(colnames(rna_data),1,2)

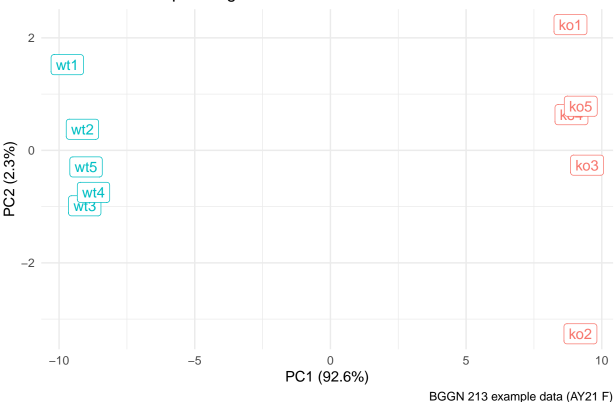
ggplot(data = rna_pca_df) +
   aes(x = PC1, y = PC2, label = samples, color = conditions) +
   geom_label(show.legend = FALSE)</pre>
```



Refine the ggplot figure with themes and labels

```
ggplot(data = rna_pca_df) +
  aes(x = PC1, y = PC2, label = samples, color = conditions) +
  geom_label(show.legend = FALSE) +
  labs(
    title = "PCA of RNA-sequencing data",
    x = paste0("PC1 (", rna_pca_varience_percent[1], "%)"),
    y = paste0("PC2 (", rna_pca_varience_percent[2], "%)"),
    caption = "BGGN 213 example data (AY21 F)") +
  theme_minimal()
```

# PCA of RNA-sequencing data



#### Gene loadings

Explore which genes contribute most to PC1 to determine what the key, differentiating genes are between the two condition groups.

```
# Save gene loading scores for PC1
gene_loading_scores <- rna_pca$rotation[,1]

# Rank by absolute value from high to low
ranked_absolute_loading_scores <- sort(abs(gene_loading_scores), decreasing=TRUE)

# Print the gene names with the first 10 (top 10) scores
names(ranked_absolute_loading_scores[1:10])

## [1] "gene100" "gene66" "gene45" "gene68" "gene98" "gene60" "gene21"

## [8] "gene56" "gene10" "gene90"</pre>
```

## Session Info

```
sessionInfo()

## R version 4.1.1 (2021-08-10)

## Platform: x86_64-pc-linux-gnu (64-bit)

## Running under: Arch Linux

##

## Matrix products: default

## BLAS: /usr/lib/libblas.so.3.10.0

## LAPACK: /usr/lib/liblapack.so.3.10.0
```

```
##
## locale:
  [1] LC CTYPE=en US.UTF-8
                                   LC NUMERIC=C
  [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=en_US.UTF-8
                                   LC_MESSAGES=en_US.UTF-8
## [7] LC PAPER=en US.UTF-8
                                   LC NAME=C
## [9] LC ADDRESS=C
                                   LC TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats
                graphics grDevices utils
                                               datasets methods
                                                                   base
## other attached packages:
## [1] ggplot2_3.3.5
##
## loaded via a namespace (and not attached):
## [1] knitr_1.33
                          magrittr_2.0.1
                                            tidyselect_1.1.1 munsell_0.5.0
## [5] colorspace 2.0-2 R6 2.5.0
                                                              fansi 0.5.0
                                            rlang 0.4.11
                                                              tools_4.1.1
## [9] dplyr_1.0.7
                          stringr_1.4.0
                                            highr_0.9
## [13] grid 4.1.1
                          gtable 0.3.0
                                                              utf8 1.2.1
                                            xfun 0.24
## [17] DBI_1.1.1
                          withr_2.4.2
                                            htmltools_0.5.1.1 ellipsis_0.3.2
## [21] assertthat_0.2.1 yaml_2.2.1
                                            digest_0.6.27
                                                              tibble_3.1.2
## [25] lifecycle_1.0.0
                                            farver_2.1.0
                                                              purrr_0.3.4
                          crayon_1.4.1
## [29] vctrs 0.3.8
                          glue 1.4.2
                                            evaluate 0.14
                                                              rmarkdown 2.11
                                                              pillar_1.6.1
## [33] labeling_0.4.2
                          stringi_1.7.2
                                            compiler_4.1.1
## [37] generics_0.1.0
                          scales_1.1.1
                                            pkgconfig_2.0.3
```