Lecture 6 Split-Apply-Combine

BIOF 339 October 24, 2016

Goals for today

- Learn how to merge data sets
- Learn how to reshape data sets from wide to long
- Split-apply-combine
 - Split a dataset into a list of several datasets
 - Do something to each dataset
 - Put the results back together
- Use it for
 - Running tests for many variables
 - Visualizing data with p-value annotation

The data for today

This data set is taken from a breast cancer proteome database available here and modified for this exercise.

- Clinical data: CSV | XLSX
- Proteome data: CSV | XLSX

The data for today

```
# Excel
library(readxl)
clinical_data <- read_excel('lecture6_data/BreastCancer_Clinical.xlsx')</pre>
# CSV
clinical_data <- read.csv('lecture6_data/BreastCancer_Clinical.csv',</pre>
                            stringsAsFactors=F)
     Complete.TCGA.ID Gender Age.at.Initial.Pathologic.Diagnosis ER.Status
#
                                                                  Negative
#
        TCGA-A2-A0CM FEMALE
        TCGA-BH-A18Q FEMALE
                                                              56 Negative
     PR.Status HER2.Final.Status Tumor Tumor..T1.Coded Node Node.Coded
     Negative
                       Negative
                                                              Negative
                                       T_Other NO
                                   T2
     Negative
                       Negative
                                                             Positive
                                  T2
                                            T Other N1
    Metastasis Metastasis.Coded AJCC.Stage Converted.Stage
#
             M<sub>O</sub>
                       Negative Stage IIA
                                             Stage IIA
                       Negative Stage IIB No_Conversion
     Survival.Data.Form Vital.Status Days.to.Date.of.Last.Contact
               followup
                            DECEASED
                                                              754
```

The data for today

```
expression_data <- read.csv('lecture6_data/BreastCancer_Expression.csv', stringsAsFactors=F)
head(expression_data[,1:5],2)

# TCGA_ID NP_958782 NP_958785 NP_958786 NP_000436
# 1 TCGA-A0-A12D 1.096131 1.111370 1.111370 1.107561
# 2 TCGA-C8-A131 2.609943 2.650422 2.650422 2.646374
```

Merging data

Note that

- 1. The names of the ID columns are different
- 2. They are in different orders

The merge function takes care of both of these issues

Merging data

```
final_data = merge(clinical_data,
                    expression_data[,1:5],
                    by.x = 'Complete.TCGA.ID',
                    by.y = 'TCGA_ID')
head(final_data,2)
    Complete.TCGA.ID Gender Age.at.Initial.Pathologic.Diagnosis ER.Status
#
        TCGA-A2-A0CM FEMALE
                                                               Negative
#
                                                               Negative
        TCGA-A2-A0D2 FEMALE
    PR.Status HER2.Final.Status Tumor Tumor..T1.Coded Node Node.Coded
     Negative
                      Negative T2
                                     T Other NO
                                                          Negative
     Negative
                      Negative
                                     T_Other NO
                                                           Negative
                               T2
    Metastasis Metastasis.Coded AJCC.Stage Converted.Stage
                      Negative Stage IIA Stage IIA
#
            M<sub>O</sub>
                      Negative Stage IIB Stage IIA
#
            MΘ
    Survival.Data.Form Vital.Status Days.to.Date.of.Last.Contact
              followup
                          DECEASED
                                                            754
              followup
                             LIVING
                                                           1027
                                                                 7/31
    Days.to.date.of.Death OS.event OS.Time PAM50.mRNA
```

Merging data

```
dim(clinical_data)
   [1] 77 30
dim(expression_data)
  [1] 83 11
dim(final_data)
  [1] 80 34
```

Note that we have extra rows, which usually means duplication of rows/ids. Something to check on.

R usually can split data on rows.

If we want to split on variables (columns), we have to transform the data from *wide* to *long* so that each row is the data for one individual *for one variable*.

First, of course, we have to select the variables we want to include.

Selecting variables

head(final_data,2)

```
Complete.TCGA.ID Gender Age.at.Initial.Pathologic.Diagnosis ER.Status
#
        TCGA-A2-A0CM FEMALE
                                                                  Negative
                                                                  Negative
        TCGA-A2-A0D2 FEMALE
    PR.Status HER2.Final.Status Tumor Tumor..T1.Coded Node Node.Coded
                                              T_Other
                       Negative
                                                              Negative
     Negative
                                    T2
                                                         NO
     Negative
                       Negative
                                               T Other
                                                              Negative
                                    T2
                                                         NO
    Metastasis Metastasis.Coded AJCC.Stage Converted.Stage
                                             Stage IIA
                        Negative Stage IIA
#
            M<sub>O</sub>
                        Negative Stage IIB
                                             Stage IIA
#
            M0
    Survival.Data.Form Vital.Status Days.to.Date.of.Last.Contact
               followup
                           DECEASED
                                                              754
               followup
#
                              LIVING
                                                             1027
    Days.to.date.of.Death OS.event OS.Time PAM50.mRNA
#
                       754
                                        754 Basal-like
                                       1027 Basal-like
    SigClust.Unsupervised.mRNA SigClust.Intrinsic.mRNA miRNA.Cluster/31
                                                    -13
```

Selecting variables

We have two goals here:

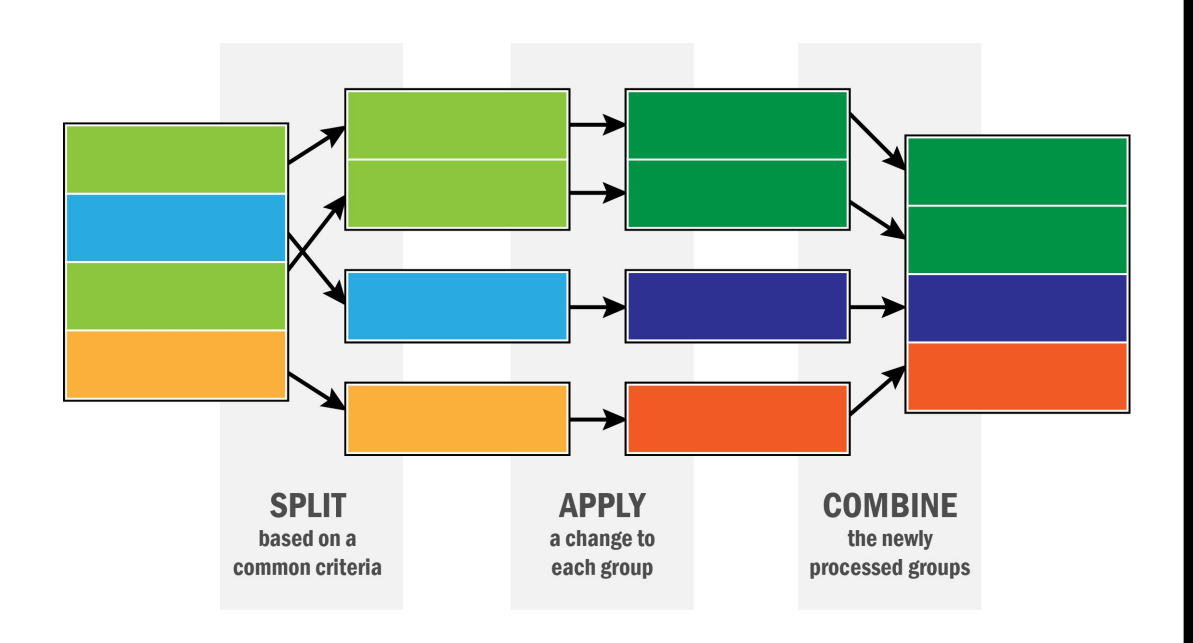
- 1. Keep the ER status aligned with each expression level, so we would need to repeat the column
- 2. Make each row have the ER status for an individual and the corresponding expression for one protein

```
library(reshape2)
reshaped_data <- melt(selected_data,</pre>
                       id.vars=c('Complete.TCGA.ID','ER.Status'))
head(reshaped_data, 5)
    Complete.TCGA.ID ER.Status variable
                                              value
#
        TCGA-A2-A0CM Negative NP_958782 0.6834035
#
        TCGA-A2-A0D2 Negative NP_958782
#
                                          0.1074909
        TCGA-A2-A0EQ Negative NP_958782 -0.9126703
#
#
  4
        TCGA-A2-A0EV Positive NP_958782 0.4529859
# 5
        TCGA-A2-A0EX Positive NP_958782 1.1851082
```

reshaped_data <- arrange(reshaped_data, Complete.TCGA.ID) # from dplyr head(reshaped_data,5)

```
# Complete.TCGA.ID ER.Status variable value
# 1 TCGA-A2-A0CM Negative NP_958782 0.6834035
# 2 TCGA-A2-A0CM Negative NP_958785 0.6944241
# 3 TCGA-A2-A0CM Negative NP_958786 0.6980976
# 4 TCGA-A2-A0CM Negative NP_000436 0.6870771
# 5 TCGA-A2-A0D2 Negative NP_958782 0.1074909
```

Split-apply-combine



Splitting data by protein

```
split_data <- split(reshaped_data, reshaped_data$variable)</pre>
class(split_data)
  [1] "list"
length(split_data)
# [1] 4
names(split_data)
   [1] "NP_958782" "NP_958785" "NP_958786" "NP_000436"
```

Splitting data

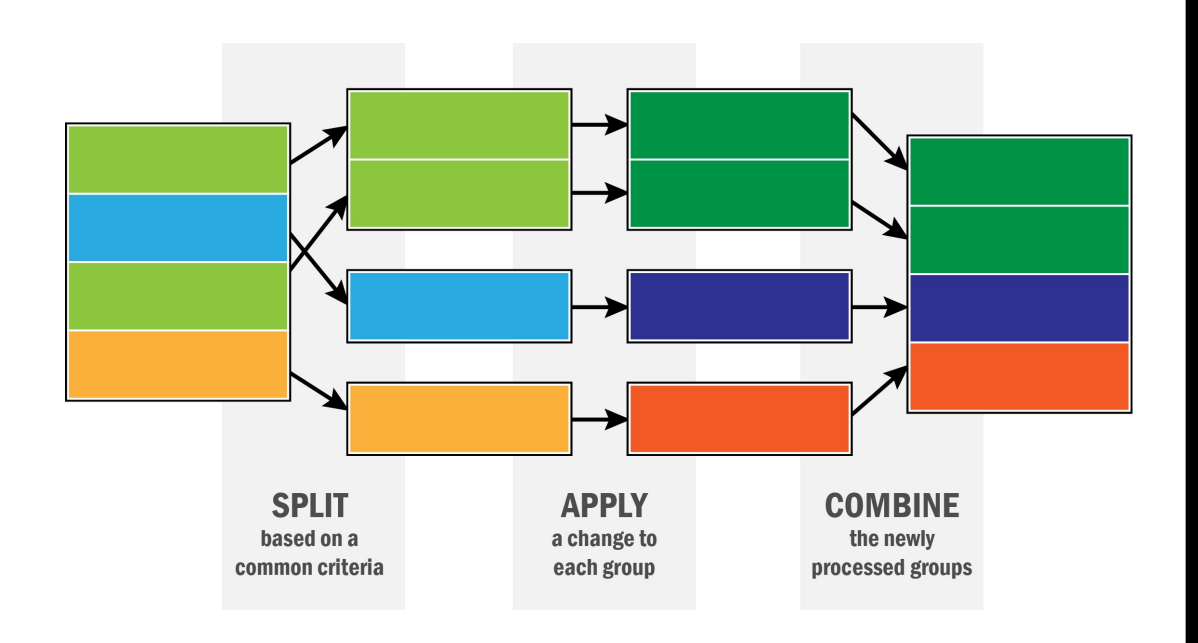
```
head(split_data[['NP_958782']],8)
```

```
Complete.TCGA.ID ER.Status variable value
#
         TCGA-A2-A0CM Negative NP_958782 0.6834035
         TCGA-A2-A0D2 Negative NP_958782 0.1074909
  5
  9
         TCGA-A2-A0EQ Negative NP_958782 -0.9126703
         TCGA-A2-A0EV Positive NP_958782 0.4529859
  13
         TCGA-A2-A0EX Positive NP_958782 1.1851082
  17
#
  21
         TCGA-A2-A0EY Positive NP_958782 1.1748810
#
         TCGA-A2-A0SW Positive NP_958782 -0.4877725
  25
#
  29
         TCGA-A2-A0SX
                      Negative NP_958782 -0.3985598
#
```

class(split_data[['NP_958782']])

```
# [1] "data.frame"
```

Applying a function to the split data



Applying a function to the split data

We're going to run t-tests to see if the expression for each protein differs by ER status

First we create a function that we'll run on every protein-specific data set

```
run_t_test <- function(d) {
   test <- t.test(value ~ ER.Status, data=d)
   pvalue <- test$p.value
   return(pvalue)
}</pre>
```

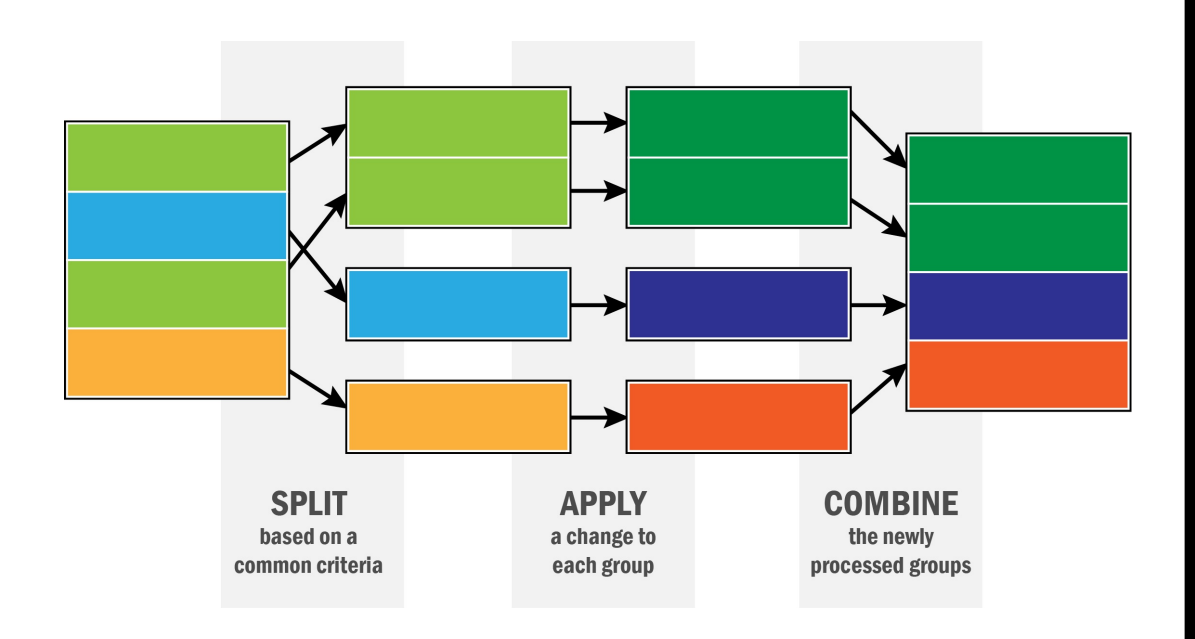
Applying a function to the split data

Now we can apply the same function to each element of the list of data frames using the lapply command

```
pvalues <- lapply(split_data, run_t_test)
pvalues</pre>
```

```
# $NP_958782
# [1] 0.5287476
#
# $NP_958785
# [1] 0.5243093
#
# $NP_958786
# [1] 0.519653
#
# $NP_000436
# [1] 0.5337173
```

Combining the data



Combining the data

```
pvalues_final <- unlist(pvalues)
pvalues_final</pre>
```

```
# NP_958782 NP_958785 NP_958786 NP_000436
# 0.5287476 0.5243093 0.5196530 0.5337173
```

Combining the data

pvalues_final <- plyr::ldply(pvalues) # Use ldply from the plyr package pvalues_final

```
# .id V1
# 1 NP_958782 0.5287476
# 2 NP_958785 0.5243093
# 3 NP_958786 0.5196530
# 4 NP_000436 0.5337173
```

The ldply function inputs a list and exports a data.frame with the elements concatenated if possible.

Plotting

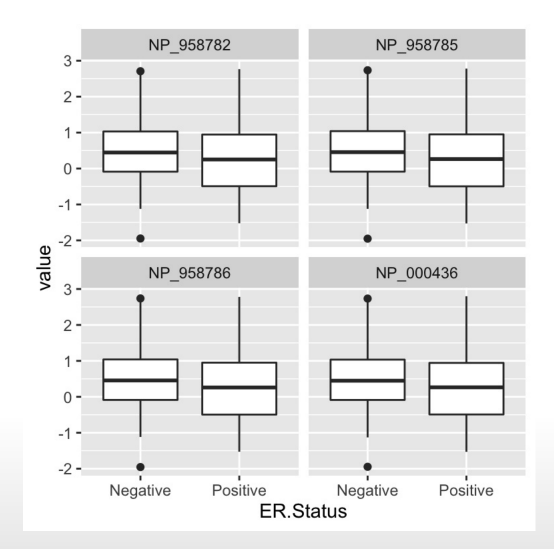
The data

head(reshaped_data, 5)

```
# Complete.TCGA.ID ER.Status variable value
# 1 TCGA-A2-A0CM Negative NP_958782 0.6834035
# 2 TCGA-A2-A0CM Negative NP_958785 0.6944241
# 3 TCGA-A2-A0CM Negative NP_958786 0.6980976
# 4 TCGA-A2-A0CM Negative NP_000436 0.6870771
# 5 TCGA-A2-A0D2 Negative NP_958782 0.1074909
```

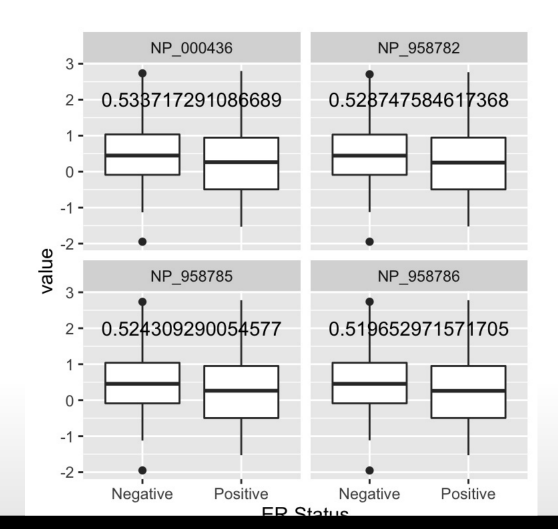
A panel of plots

```
library(ggplot2)
ggplot(reshaped_data, aes(x = ER.Status, y = value))+
  geom_boxplot()+
  facet_wrap(~variable, ncol=2)
```

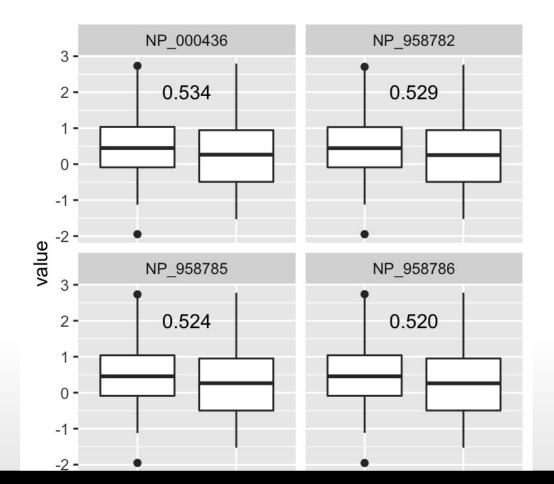


Adding p-values to the plot

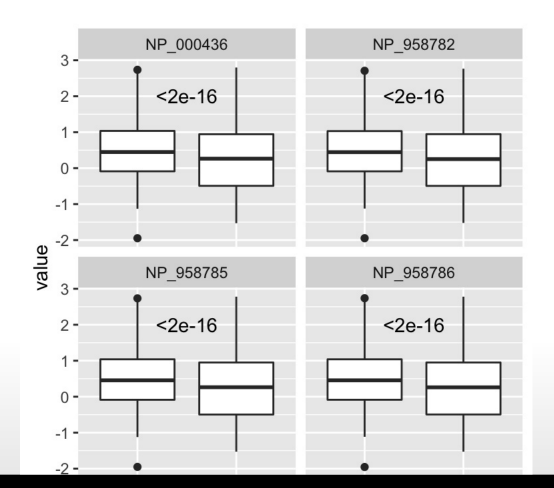
```
names(pvalues_final) <- c('variable','pvalue')
ggplot(reshaped_data, aes(x = ER.Status, y = value))+
  geom_boxplot()+
  facet_wrap(~variable, ncol=2)+
  geom_text(data=pvalues_final, aes(x=1.5, y=2, label=pvalue))</pre>
```



Formatting p-values



Formatting p-values



The dplyr package

Action words

dplyr basically gives you 6 actions to do on data.frame objects:

- 1. mutate: Change particular variables
- 2. select: Select (or deselect) variables
- 3. arrange: Order by some variables
- 4. filter: Select rows by some criteria
- 5. group_by: Group by some variable (so the split part of our exercise)
- 6. summarise: Summarise a variable using some function