Homework 2 Solution

Brian Lois 1/27/2017

```
library(ggplot2)
library(caret)
library(mlbench)
library(corrplot)
```

1.

```
RMSE <- function(y, y_hat){
  sq_er <- (y - y_hat)^2
  mean_sq_er <- mean(sq_er)
  rmse <- sqrt(mean_sq_er)
  return(rmse)
}</pre>
```

2.

```
data(Glass)
```

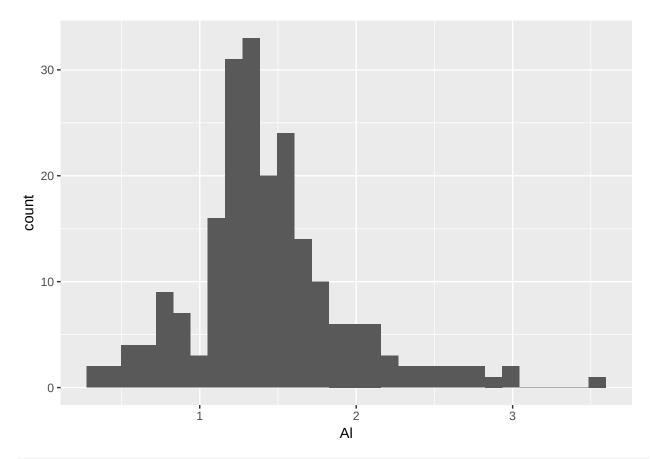
a)

Look at all histograms

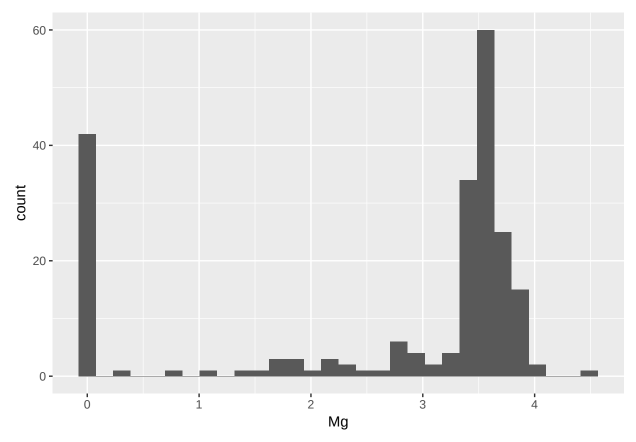
```
for (col in colnames(Glass)){
  print(ggplot(data = Glass) + geom_histogram(mapping = aes_string(x = col)))
}
```

Only plot 2 (any two interesting ones are correct).

```
ggplot(data = Glass) + geom_histogram(mapping = aes(x = Al))
```

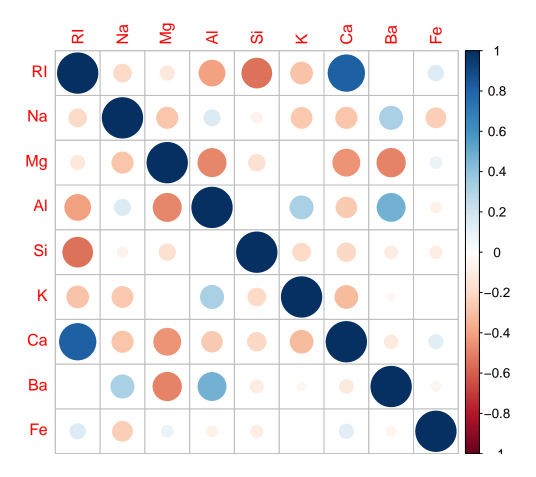


ggplot(data = Glass) + geom_histogram(mapping = aes(x = Mg))



Correlation matrix:

```
correlation_matrix <- cor(subset(Glass, select = -c(Type)))
corrplot(correlation_matrix)</pre>
```



b)

```
skewness_list <- apply(Glass[,1:9], 2, BoxCoxTrans)
for (col in colnames(Glass)[1:9]){
   print(paste(col, skewness_list[[col]]$skewness))
}
## [1] "RI 1.60271508274373"</pre>
```

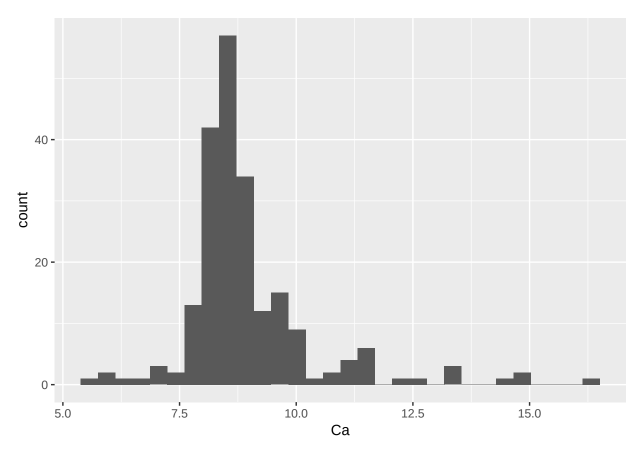
```
## [1] "Na 0.447834258917133"
## [1] "Mg -1.13645227846653"
## [1] "Al 0.89461041611312"
## [1] "Si -0.720239210805621"
## [1] "K 6.46008889572281"
## [1] "Ca 2.01844629445302"
## [1] "Ba 3.36867996880571"
```

[1] "Fe 1.7298107095598"

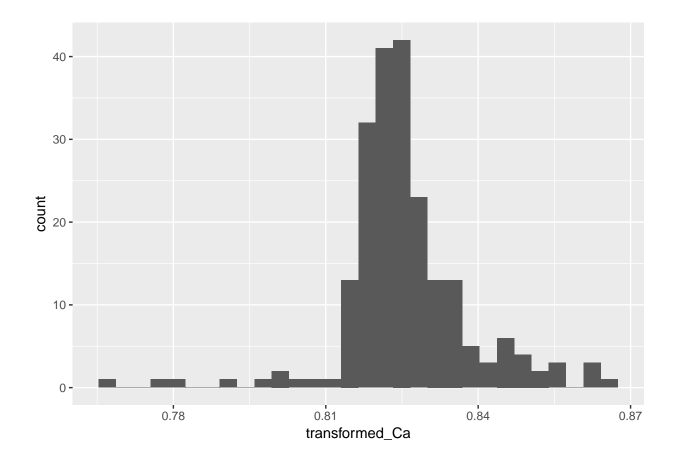
c)

We will do the Box-Cox transformation on Ca.

ggplot(data = Glass) + geom_histogram(mapping = aes(x = Ca))



Glass['transformed_Ca'] <- predict(skewness_list\$Ca, Glass\$Ca)
ggplot(data = Glass) + geom_histogram(mapping = aes(x = transformed_Ca))</pre>



3.

a)

```
data("Soybean")
P <- ncol(Soybean)
n <- nrow(Soybean)</pre>
degenerate <- function(dist){</pre>
  frac_unique <- length(dist) / sum(dist)</pre>
  ratio <- sort(dist, decreasing = TRUE)[1] / sort(dist, decreasing = TRUE)[2]</pre>
  if ((frac\_unique < .1) & (ratio > 20)){}
    return(TRUE)
  } else {
    return(FALSE)
  }
}
var_tables <- apply(Soybean, 2, table)</pre>
degenerate_features <- sapply(var_tables, degenerate)</pre>
Soybean <- Soybean[ ,!degenerate_features]</pre>
P <- ncol(Soybean)
```

b)

```
percent_missing <- function(x){
   p_m <- sum(is.na(x)) / length(x)
   return(p_m)
}</pre>
```

Percent missing by feature:

```
apply(Soybean, 2, percent_missing)
```

##	Class	date	plant.stand	precip
##	0.00000000	0.001464129	0.052708638	0.055636896
##	temp	hail	crop.hist	area.dam
##	0.043923865	0.177159590	0.023426061	0.001464129
##	sever	seed.tmt	germ	plant.growth
##	0.177159590	0.177159590	0.163982430	0.023426061
##	leaves	leaf.halo	leaf.marg	leaf.size
##	0.00000000	0.122986823	0.122986823	0.122986823
##	leaf.shread	leaf.malf	stem	lodging
##	0.146412884	0.122986823	0.023426061	0.177159590
##	stem.cankers	canker.lesion	fruiting.bodies	ext.decay
##	0.055636896	0.055636896	0.155197657	0.055636896
##	int.discolor	fruit.pods	fruit.spots	seed
##	0.055636896	0.122986823	0.155197657	0.134699854
##	mold.growth	seed.discolor	seed.size	shriveling
##	0.134699854	0.155197657	0.134699854	0.155197657
##	roots			
##	0.045387994			

Yes, some features are much more likely to be missing.

Percent missing by class and feature: (Output is suppressed to save space)

```
missing_mat <- list()
for (level in levels(Soybean$Class)){
  for (col in colnames(Soybean)[2:P]){
    vec <- Soybean[[col]][Soybean$Class == level]
    missing_mat[[level]][[col]] <- percent_missing(vec)
  }
}
print(missing_mat)</pre>
```

Many classes have no missing values, and some classes have a lot of missing values. Some classes may even have all of the features missing.

c)

Possible strategies could include any combination of:

• Removing whole classes such as herbicide-injury and cyst-nematode.

- Mean, median, or knn imputation
- I don't see any columns that I would entirely remove based on missingness, but I won't take off points for removing features with more than 10% missing.
- Adding missing indicator columns.

The best approach here is unclear without knowing the ultimate goal of the project.

4.

a)

```
data(BloodBrain)
```

b)

Reuse our function from before

```
bbb_var_tables <- apply(bbbDescr, 2, table)
degenerate_vec <- sapply(bbb_var_tables, degenerate)
sum(degenerate_vec)</pre>
```

```
## [1] 7
```

```
degenerate_vec[degenerate_vec]
```

```
## negative peoe_vsa.2.1 peoe_vsa.3.1 a_acid vsa_acid
## TRUE TRUE TRUE TRUE TRUE
## frac.anion7. alert
## TRUE TRUE
```

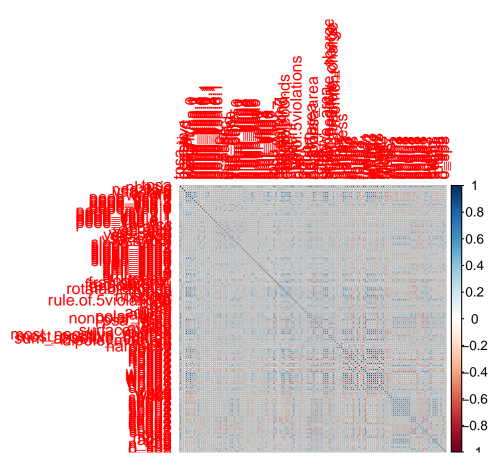
```
var_vec <- apply(bbbDescr, 2, var)
head(sort(var_vec), 10)</pre>
```

```
## chaa3 chdh3 tcsa tcnp fpsa3
## 7.512226e-07 8.830284e-07 9.064777e-06 4.618282e-05 3.793002e-04
## scdh3 scaa3 fnsa3 sadh3 rpcg
## 4.298434e-04 8.640507e-04 1.341075e-03 2.342703e-03 2.958051e-03
```

Seven are degenerate as defined in the chapter, and some others have \mathbf{very} near-zero variance.

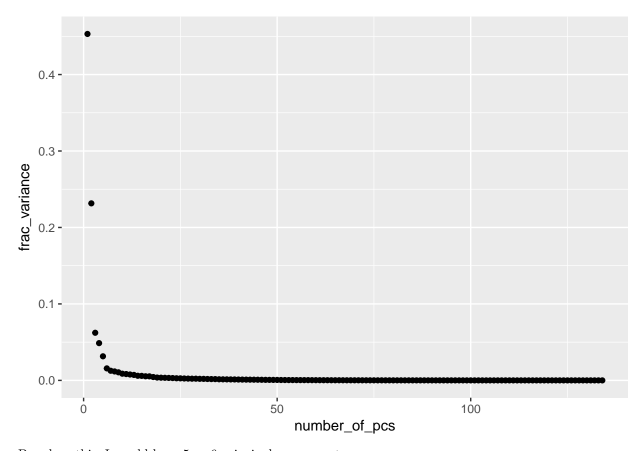
c)

```
cor_mat <- cor(bbbDescr)
corrplot(cor_mat)</pre>
```



Yes, there do appear to be strong relationships (dark colors in the corrplot).

```
pc <- prcomp(bbbDescr)
scaled_stdev <- as.data.frame(cbind(pc$sdev / sum(pc$sdev), 1:length(pc$sdev)))
colnames(scaled_stdev) <- c('frac_variance', 'number_of_pcs')
ggplot(data=scaled_stdev) + geom_point(mapping=aes(x=number_of_pcs, y=frac_variance))</pre>
```



Based on this, I would keep 5 or 6 principal components.