21BDS0340

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Exploratory Data Analysis Lab

Assignment - II

Experiment 5

Code:

library(dplyr)
library(missForest)
library(mice)
library(VIM)
library(ggplot2)
library(cowplot)

data = iris
View(data)

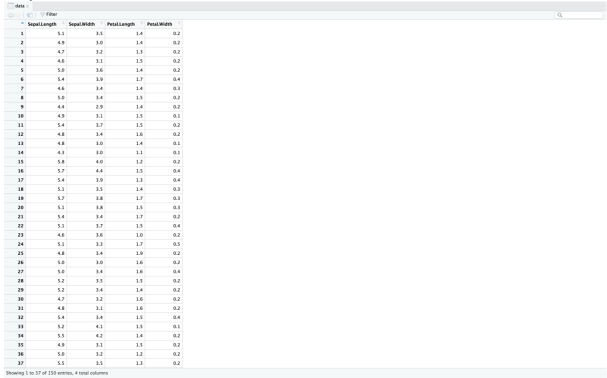
Output:

```
### Spallength | Spallwight | Petallwigh | Petallwigh | Species | Spallength | Spallwigh | Petallwigh | Species | Spallength | Spallwigh | Petallwigh | Species | Spallength | Spallwigh | Species | Spallength | Spallwigh | Species | Spallength | Spallwigh | Spallength | Spallen
```

Code:

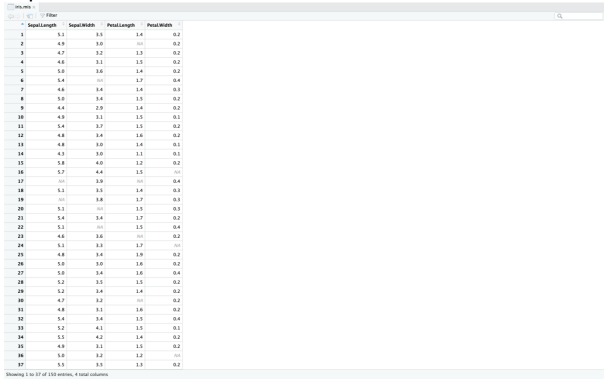
```
# dropping labels
data = data %>% select(-c("Species"))
View(data)
```

Output:



Code:

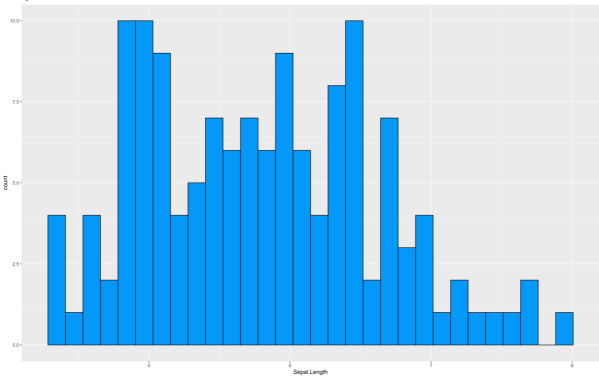
adding 10% random values
iris.mis <- prodNA(data, noNA = 0.1)
View(iris.mis)</pre>



Code:

```
ggplot(iris.mis, aes(x = Sepal.Length)) +
  geom_histogram(color="black", fill="#0099F8")
```

Output:

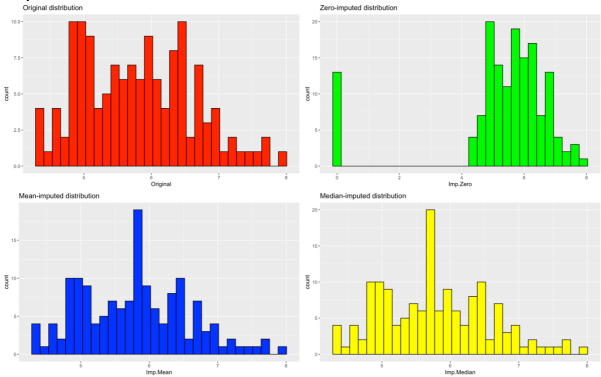


Code:

```
# simple imputations for Sepal.Length
imputed = data.frame(
 Original = iris.mis$Sepal.Length,
 Imp.Zero = replace(iris.mis$Sepal.Length, is.na(iris.mis$Sepal.Length), 0),
  Imp.Mean = replace(iris.mis$Sepal.Length, is.na(iris.mis$Sepal.Length),
mean(iris.mis$Sepal.Length, na.rm = TRUE)),
  Imp.Median = replace(iris.mis$Sepal.Length, is.na(iris.mis$Sepal.Length),
median(iris.mis$Sepal.Length, na.rm = TRUE))
)
# plotting the simple imputations
h1 = ggplot(imputed, aes(x=0riginal)) +
 geom_histogram(fill="red", color="black", position="identity") +
 ggtitle("Original distribution")
h2 = ggplot(imputed, aes(x=Imp.Zero)) +
 geom_histogram(fill="green", color="black", position="identity") +
  ggtitle("Zero-imputed distribution")
h3 = ggplot(imputed, aes(x=Imp.Mean)) +
 geom_histogram(fill="blue", color="black", position="identity") +
  ggtitle("Mean-imputed distribution")
h4 = ggplot(imputed, aes(x=Imp.Median)) +
 geom_histogram(fill="yellow", color="black", position="identity") +
  ggtitle("Median-imputed distribution")
```

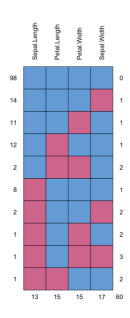
plot_grid(h1, h2, h3, h4, nrow=2, ncol=2)

Output:



Code:

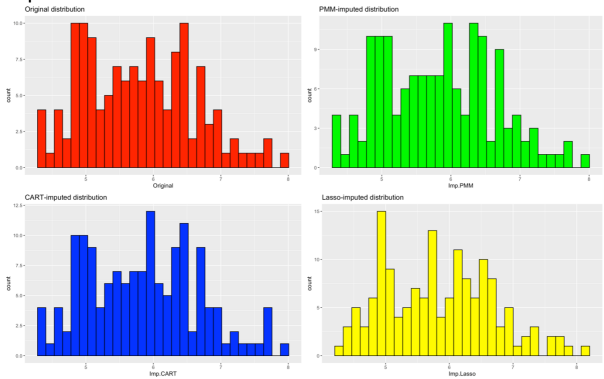
viewing missing values
md.pattern(iris.mis, rotate.names=TRUE)



Code:

```
# performing imputations with mice algorithms
mice_imputed = data.frame(
  Original = iris.mis$Sepal.Length,
  Imp.PMM = complete(mice(iris.mis, method="pmm"))$Sepal.Length,
  Imp.CART = complete(mice(iris.mis, method="cart"))$Sepal.Length.
  Imp.Lasso = complete(mice(iris.mis, method="lasso.norm"))$Sepal.Length
)
# plotting the mice imputations
h1 = ggplot(mice_imputed, aes(x=0riginal)) +
  geom_histogram(fill="red", color="black", position="identity") +
  ggtitle("Original distribution")
h2 = ggplot(mice_imputed, aes(x=Imp.PMM)) +
  geom_histogram(fill="green", color="black", position="identity") +
  ggtitle("PMM-imputed distribution")
h3 = ggplot(mice_imputed, aes(x=Imp.CART)) +
  geom_histogram(fill="blue", color="black", position="identity") +
  ggtitle("CART-imputed distribution")
h4 = ggplot(mice_imputed, aes(x=Imp.Lasso)) +
  geom_histogram(fill="yellow", color="black", position="identity") +
  ggtitle("Lasso-imputed distribution")
plot_grid(h1, h2, h3, h4, nrow=2, ncol=2)
```

Output:



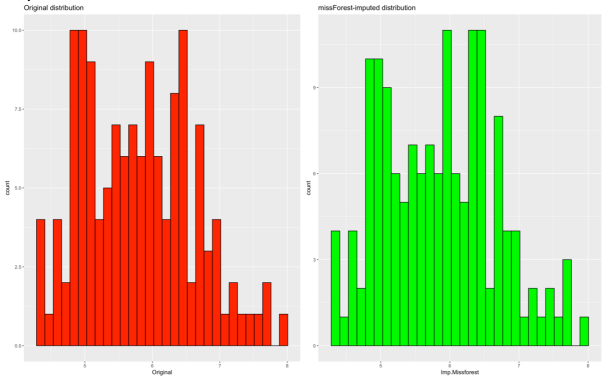
Code:

```
# imputations with missForest
missforest_imputed = data.frame(
    Original = iris.mis$Sepal.Length,
```

```
Imp.Missforest = missForest(iris.mis)$ximp$Sepal.Length
)

# plotting the missForest imputations
h1 = ggplot(missforest_imputed, aes(x=Original)) +
    geom_histogram(fill="red", color="black", position="identity") +
    ggtitle("Original distribution")
h2 = ggplot(missforest_imputed, aes(x=Imp.Missforest)) +
    geom_histogram(fill="green", color="black", position="identity") +
    ggtitle("missForest-imputed distribution")

plot_grid(h1, h2, nrow=1, ncol=2)
```



Experiment 6

Code:

```
X = scan(what = numeric())
Y = scan(what = numeric())

cov(X, Y)
cor(X, Y)
```

```
> X = scan(what = numeric())
1: 1 2 3 4 5 6 7 8 9 10
11:
Read 10 items
> Y = scan(what = numeric())
1: 9 3 4 1 5 6 2 7 10 9
11:
Read 10 items
> cov(X, Y)
[1] 3.777778
> cor(X, Y)
[1] 0.3981308
```

```
Experiment 7
```

```
Code:
```

```
# z score method
data = c(10, 10, 11, 11, 12, 12, 13, 12, 11, 14, 13, 15, 100)
mean.data = mean(data)
std.data = sd(data)
z.scores = (data - mean.data) / std.data
# outliers have -3 < z.score < 3</pre>
outliers = data[abs(z.scores) > 3]
outliers
Output:
> # z score method
> data = c(10, 10, 11, 11, 12, 12, 13, 12, 11, 14, 13, 15, 100)
> mean.data = mean(data)
> std.data = sd(data)
> z.scores = (data - mean.data) / std.data
> # outliers have -3 < z.score < 3</pre>
> outliers = data[abs(z.scores) > 3]
> outliers
[1] 100
Code:
# inter quartile range method
data = c(10, 10, 11, 11, 12, 12, 13, 12, 11, 14, 13, 15, 100)
q1 = quantile(data, 0.25)
q3 = quantile(data, 0.75)
iqr = q3 - q1
# outliers lie outside of the inter quartile range
outliers <- data[data < q1 | data > q3]
outliers
Output:
> # inter quartile range method
> data = c(10, 10, 11, 11, 12, 12, 13, 12, 11, 14, 13, 15, 100)
> q1 = quantile(data, 0.25)
> q3 = quantile(data, 0.75)
> iqr = q3 - q1
> # outliers lie outside of the inter quartile range
> outliers <- data[data < q1 | data > q3]
> outliers
```

```
[1] 10 10 14 15 100
```

Code:

```
# boxplot method (purely visualisation)
data = c(10, 10, 11, 11, 12, 12, 13, 12, 11, 14, 13, 15, 100)
boxplot(data)
```

