Statistics 135 – Lab Project

April 28, 2015

1 Background

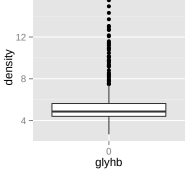
```
library("ggplot2")
library("grid")
library("gridExtra")

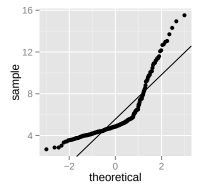
data.df <- na.omit(read.csv("diabetes.csv"))</pre>
```

2 Accessing Data, Visualization and Summarization



```
75 - 150 - 25 - 4 8 12 16 glyhb
```

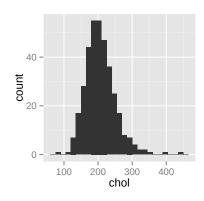


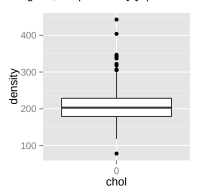


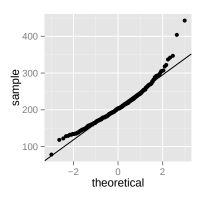
```
2. chol.histogram <- ggplot(data.df) +
    geom_histogram(aes(x=chol), binwidth=15)

chol.boxplot <- ggplot(data.df) +</pre>
```

Histogram, Boxplot and QQ-plot of chol



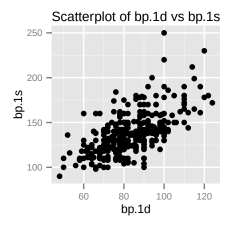


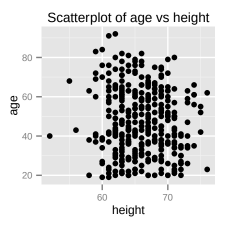


```
3. bp.1d.bp.1s.scatterplot <- ggplot(data.df) +
    geom_point(aes(x=bp.1d, y=bp.1s)) +
    labs(title="Scatterplot of bp.1d vs bp.1s") +
    theme(text=element_text(size=8.5))

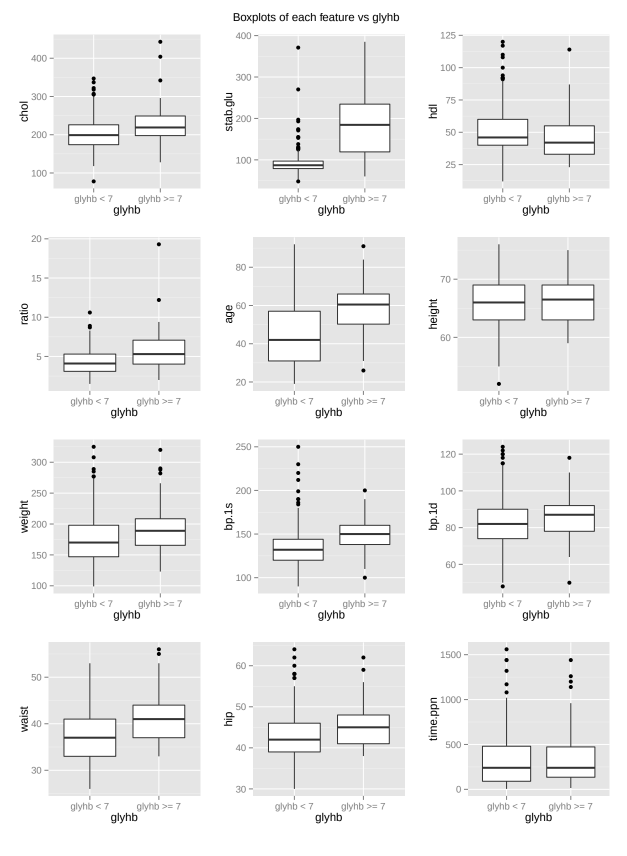
height.age.scatterplot <- ggplot(data.df) +
    geom_point(aes(x=height, y=age)) +
    labs(title="Scatterplot of age vs height") +
    theme(text=element_text(size=8.5))

grid.arrange(bp.1d.bp.1s.scatterplot, height.age.scatterplot, ncol=2)</pre>
```



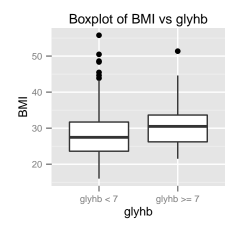


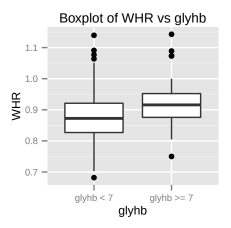
```
4. data.df$glyhb.cond <- NA data.df$glyhb>=7, ]$glyhb.cond <- "glyhb >= 7" data.df[data.df$glyhb<7, ]$glyhb.cond <- "glyhb < 7"
```



```
5. data.df$BMI <- 703*data.df$weight/data.df$height^2
data.df$WHR <- data.df$waist/data.df$hip

features <- c("BMI", "WHR")
feature.boxplots = list()
for (i in 1:length(features)) {
   feature <- features[i]</pre>
```





6.

3 Parametric Inferece

1.

$$X \sim Gamma(\alpha, \beta) = \frac{\beta^{\alpha}}{\Gamma(\alpha)} x^{\alpha - 1} e^{-\beta x}$$
$$E(X) = \frac{\alpha}{\beta}$$
$$E(X^{2}) = Var(X) + [E(X)]^{2}$$

$$E(X') = Var(X) + [E(X)]$$

$$= \frac{\alpha}{\beta^2} + \left(\frac{\alpha}{\beta}\right)^2$$

$$= \frac{\alpha(\alpha + 1)}{\beta^2}$$

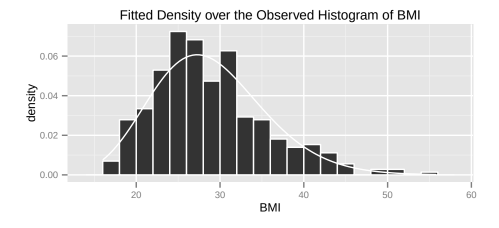
$$\begin{cases} E(X) = \frac{\alpha}{\beta} \\ E(X^2) = \frac{\alpha(\alpha+1)}{\beta^2} \end{cases} \implies \begin{cases} \alpha = \frac{[E(X)]^2}{Var(x)} \\ \beta = \frac{E(X)}{Var(x)} \end{cases} \implies \begin{cases} \hat{\alpha}_{MOM} = \frac{\overline{X}_n}{\frac{1}{n} \sum_{i=1}^n (X_i - \overline{X}_n)^2} \\ \hat{\beta}_{MOM} = \frac{\overline{X}_n^2}{\frac{1}{n} \sum_{i=1}^n (X_i - \overline{X}_n)^2} \end{cases}$$

```
alpha.mom <- mean(data.df$BMI)^2/var(data.df$BMI)
beta.mom <- mean(data.df$BMI)/var(data.df$BMI)

B <- 1000
BMI.bootstrap <- sapply(1:B, function(x) {
    BMI.sample <- sample(data.df$BMI, nrow(data.df), replace=TRUE)
    alpha.sample <- mean(BMI.sample)^2/var(BMI.sample)</pre>
```

```
beta.sample <- mean(BMI.sample)/var(BMI.sample)</pre>
 return(c(alpha.sample, beta.sample))
})
CI.BMI <- sapply(1:2, function(x) {</pre>
 CI <- quantile(BMI.bootstrap[x, ], probs=c(0.025, 0.975))</pre>
 return(CI)
})
colnames(CI.BMI) <- c("alpha", "beta")</pre>
CI.BMI
##
        alpha
                beta
## 2.5% 15.91447 0.5477096
## 97.5% 21.79548 0.7626395
```

```
ggplot(data.df) +
  geom_histogram(aes(x=BMI, y=..density..), binwidth=2, col="white") +
  stat_function(fun=function(x)
   dgamma(x, shape=alpha.mom, rate=beta.mom), col="white") +
  labs(title="Fitted Density over the Observed Histogram of BMI") +
  theme(text=element_text(size=8.5))
```



2.

3.

4 Testing

1.

2.

3.

4.

5.

6.

5 Regression

- 1.
- 2.
- 3.
- 4.
- 5.
- 6.
- 7.