Environmental Statistics Homework 3

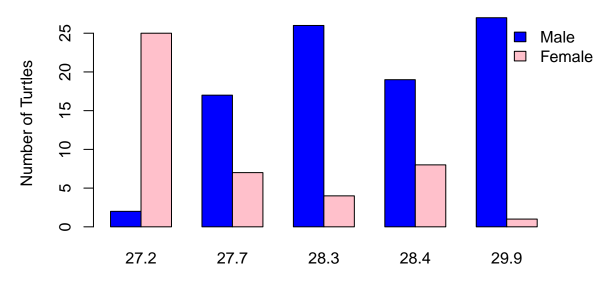
Environmental Statistics

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Exercise 1

```
library(faraway)
data("turtle")
attach(turtle)
agg_data <- aggregate(cbind(turtle$male, female) ~ temp, data = turtle, sum)</pre>
# Create a bar plot
par(mar = c(5, 4, 4, 2))
barplot(
  t(as.matrix(agg_data[, -1])),
  beside = TRUE,
  col = c("blue", "pink"),
  names.arg = agg_data$temp,
  xlab = "Incubation Temperature (°C)",
  ylab = "Number of Turtles",
  main = "Sex Determination in Turtles",
  legend.text = c("Male", "Female"),
  args.legend = list(x ='topright', inset=c(-0.1,0), bty = "n")
```

Sex Determination in Turtles



Incubation Temperature (°C)

```
# Models
  modmale <- glm(cbind(male, female) ~ temp, data = turtle, family = "binomial")</pre>
  summary(modmale)
Call:
glm(formula = cbind(male, female) ~ temp, family = "binomial",
    data = turtle)
Deviance Residuals:
    Min
              1Q
                 Median
                                3Q
                                        Max
-2.0721 -1.0292 -0.2714
                                     2.5550
                           0.8087
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -61.3183
                       12.0224 -5.100 3.39e-07 ***
temp
              2.2110
                        0.4309
                                5.132 2.87e-07 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 74.508 on 14 degrees of freedom
Residual deviance: 24.942 on 13 degrees of freedom
AIC: 53.836
Number of Fisher Scoring iterations: 5
```

```
modfem <- glm(cbind(female, male) ~ temp, data = turtle, family = "binomial")</pre>
  summary(modfem)
Call:
glm(formula = cbind(female, male) ~ temp, family = "binomial",
    data = turtle)
Deviance Residuals:
   Min
              10
                  Median
                                30
                                        Max
-2.5550 -0.8087
                  0.2714
                          1.0292
                                     2.0721
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) 61.3183
                        12.0224 5.100 3.39e-07 ***
             -2.2110
                         0.4309 -5.132 2.87e-07 ***
temp
___
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 74.508 on 14 degrees of freedom
Residual deviance: 24.942 on 13 degrees of freedom
AIC: 53.836
Number of Fisher Scoring iterations: 5
```

For the male model, the intercept is negative and the effect of temperature on the male gender is positive, i.e. with increasing temperature, the probability of male turtles hatching increases. For the female model it is the other way around, the intercept is positive and the effect of temperature on the female gender negative, so with increasing temperature, the probability of female turtles hatching decreases. In both cases, the effect of temperature was highly significant.

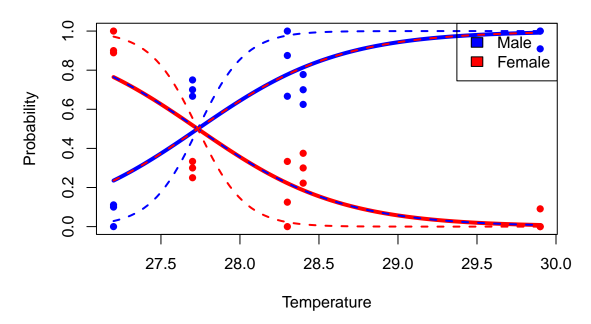
```
# Males
lines(new_data$temp, plogis(predicted_probsmale$fit), type = "l", lwd=4, col =
# Females
lines(new_data$temp, plogis(predicted_probsfemale$fit), type = "l",lwd=4, col =

    "red")

# 95% confidence
lines(new_data$temp, plogis(predicted_probsmale$fit + 2 *

→ predicted_probsmale$fit),
      type = "1", lty=2, lwd=2, col = "blue" )
lines(new_data$temp, plogis(predicted_probsmale$fit - 2 *
→ predicted_probsmale$fit),
      type = "1", lty=2, lwd=2, col = "blue" )
lines(new_data$temp, plogis(predicted_probsfemale$fit + 2*
→ predicted_probsfemale$fit),
      type = "1", lty= 2, lwd=2, col = "red")
lines(new_data$temp, plogis(predicted_probsfemale$fit - 2*
→ predicted_probsfemale$fit),
      type = "1", lty= 2, lwd=2, col = "red")
legend("topright", legend = c("Male", "Female"), fill = c("blue", "red"))
```

Probability Turtle Emergence vs. Temperature



```
xprob <- new_data$temp</pre>
  yprob <- plogis(predicted_probsfemale$fit)</pre>
  # Define the y-value for which you want to estimate the x-value
  desired_y <- 0.5</pre>
  # Use the `approx` function to estimate the x-value
  estimated_x <- approx(yprob, xprob, xout = desired_y)$y</pre>
  cat("Estimated x-value for y =", desired_y, "is x =", estimated_x, "\n")
Estimated x-value for y = 0.5 is x = 27.7329
  attach(turtle)
The following objects are masked from turtle (pos = 3):
    female, male, temp
  #Plot with line at 50/50 point
  par(mar = c(5, 4, 4, 2))
  plot(male / (female + male) ~ temp, pch = 16, col = "blue",
      xlab = "Temperature", ylab = "Probability",
      main = "Probability Turtle Emergence vs. Temperature")
  points(female / (female + male) ~ temp, pch = 16, col = "red")
  # Males
  lines(new_data$temp, plogis(predicted_probsmale$fit), type = "l", lwd=4, col =
   → "blue" )
  # Females
  lines(new_data$temp, plogis(predicted_probsfemale$fit), type = "l",lwd=4, col =

    "red")

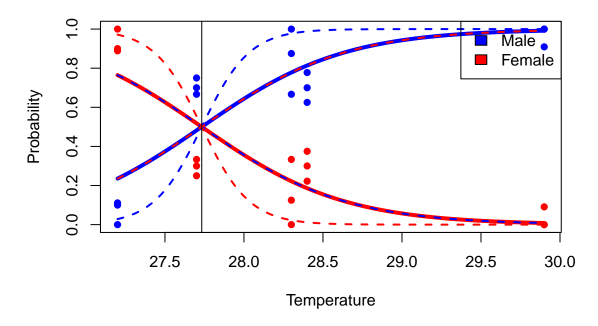
  # 95% confidence
  lines(new_data$temp, plogis(predicted_probsmale$fit + 2 *
   → predicted_probsmale$fit),
        type = "1", lty=2, lwd=2, col = "blue" )
  lines(new_data$temp, plogis(predicted_probsmale$fit - 2 *

→ predicted probsmale$fit),
        type = "1", lty=2, lwd=2, col = "blue" )
```

```
lines(new_data$temp, plogis(predicted_probsfemale$fit + 2*
    predicted_probsfemale$fit),
        type = "l", lty= 2,lwd=2, col = "red")
lines(new_data$temp, plogis(predicted_probsfemale$fit - 2*
    predicted_probsfemale$fit),
        type = "l", lty= 2, lwd=2, col = "red")

legend("topright", legend = c("Male", "Female"), fill = c("blue", "red"))
abline(v=27.7329)
```

Probability Turtle Emergence vs. Temperature



Instead of trying to find the intersect between the two lines, we tried to determine the temperature at which 50% of the hatched turtles are female, which would automatically be the 50/50 point between male and female.

Subexercise 1.5

The 50/50 point is very far to the left, i.e. the temperature is very low. This shows that over the course of the whole experiment, the probability of receiving a male turtle is higher than that of receiving a female turtle, because the regression line for male is above that for female for most of the time. At a temperature of 27,75 °C there is exactly a 50/50 chance that an egg will yield a female turtle or male turtle. The 50/50 chance of hatching can be seen at the intersect of the predicted male and female hatching curves. Regarding the plot it is remarkable to say that the 95% confidence interval of male hatching is the same as the regression line of female hatching after the intersection at 27.75 °C and viceversa.

Exercise 2

```
library(faraway)
data("wcgs")
attach(wcgs)
```

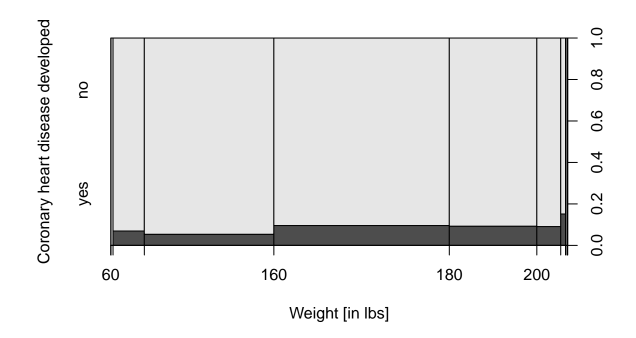
Subexercise 2.1

```
attach(wcgs)
```

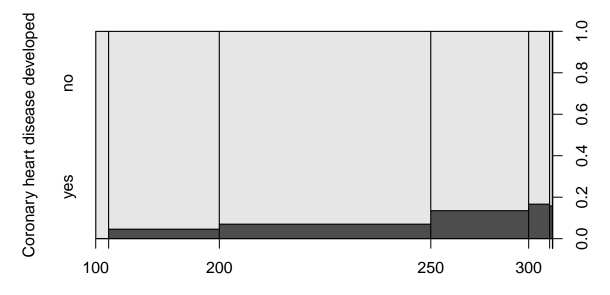
The following objects are masked from wcgs (pos = 3):

age, arcus, behave, chd, chol, cigs, dbp, dibep, height, sdp, timechd, typechd, weight

```
plot(chd ~ weight, xlab="Weight [in lbs]",
    ylab="Coronary heart disease developed")
```



plot(chd~chol, xlab="Fasting serum cholesterol [in mg/dl]",
 ylab= "Coronary heart disease developed")

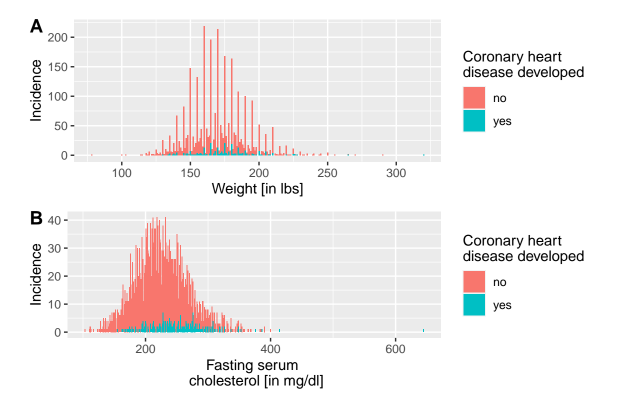


Fasting serum cholesterol [in mg/dl]

Incidence

Warning: Removed 12 rows containing non-finite values (`stat_count()`).

z



Subexercise 2.2

Deviance Residuals:

1Q

-1.0996 -0.4499 -0.3644 -0.2886

Median

```
attach(wcgs)
The following objects are masked from wcgs (pos = 5):
    age, arcus, behave, chd, chol, cigs, dbp, dibep, height, sdp,
    timechd, typechd, weight
The following objects are masked from wcgs (pos = 6):
    age, arcus, behave, chd, chol, cigs, dbp, dibep, height, sdp,
    timechd, typechd, weight

modchd <- glm(chd-weight + chol + weight*chol, family="binomial")
summary(modchd)
Call:
glm(formula = chd ~ weight + chol + weight * chol, family = "binomial")</pre>
```

3Q

Max

2.7814

Coefficients:

```
Estimate Std. Error z value Pr(>|z|)

(Intercept) -1.321e+01 2.580e+00 -5.121 3.03e-07 ***

weight 4.504e-02 1.449e-02 3.109 0.001876 **

chol 3.726e-02 1.038e-02 3.589 0.000333 ***

weight:chol -1.419e-04 5.864e-05 -2.420 0.015520 *

---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 1779.2 on 3141 degrees of freedom

Residual deviance: 1684.7 on 3138 degrees of freedom

(12 observations deleted due to missingness)

AIC: 1692.7
```

Weight and cholesterol separately have a "positive" effect, i.e. with increasing weight and increasing cholesterol levels, more people will suffer from a coronary heart disease. Both effects are significant. The effect of the interaction between weight and cholesterol levels on coronary heart diseases is estimated to be negative. This suggests that the interaction is associated with a decrease in coronary heart diseases, because the effect of e.g. weight on coronary heart diseases may be different depending on the level of cholesterol and this interaction has a negative impact on coronary heart diseases as compared to what we would expect based on the main effects alone. As in this case a negative effect indicates a decrease in coronary heart diseases it could be because if the cholesterol levels are lower, the effect of weight isn't as strong anymore and viceversa.

```
modchd$deviance/modchd$df.residual
```

Number of Fisher Scoring iterations: 5

[1] 0.5368713

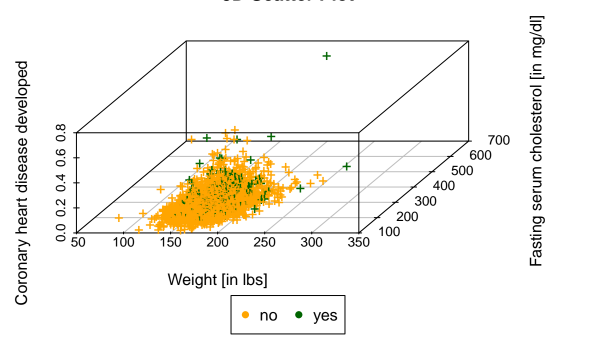
The dispersion parameter is lower than 1, indicating underdispersion. This means that the observed variance in the response variable (i.e. coronary heart diseases) is less than expected based on the model. This suggests that the model is overly conservative in its estimates of variability.

```
library(faraway)
data("wcgs")
library(scatterplot3d)
#install.packages("scatterplot3d")

# count cells with na values in the chol column sum(is.na(wcgs$chol))
```

```
#remove na's to avoid errors
wcgs <- wcgs[!is.na(wcgs$chol),]</pre>
# Generate a linear model
mod <- glm(chd~weight + chol + weight*chol, family="binomial", data = wcgs)</pre>
# Get predicted values of chd from chol and weight
wcgs$pred_chd_1 <- predict(mod, type = "response")</pre>
#2 Farben:
colors <- c("orange", "darkgreen")</pre>
colors <- colors[as.numeric(wcgs$chd)]</pre>
scatterplot3d(wcgs$weight, wcgs$chol, wcgs$pred_chd_1, type="p", xlab="Weight [in
    lbs]",
              ylab="Fasting serum cholesterol [in mg/dl]",
              angle=55,
              zlab= "Coronary heart disease developed",
              pch = "+", color = colors,
              main = "3D Scatter Plot")
legend("bottom", legend = levels(wcgs$chd),
       col = c("orange", "darkgreen"), pch = 16,
       inset = -0.45, xpd = TRUE, horiz = TRUE)
```

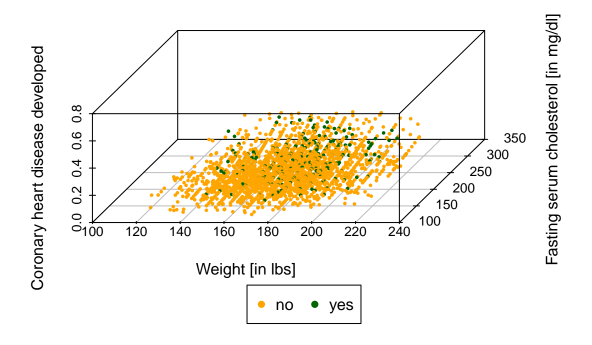
3D Scatter Plot



[1] 12

```
#remove na's to avoid errors
wcgs <- wcgs[!is.na(wcgs$chol),]</pre>
# Generate a linear model
mod <- glm(chd~weight + chol + weight*chol, family="binomial", data = wcgs)</pre>
# Get predicted values of chd from chol and weight
wcgs$pred_chd_2 <- predict(mod, type = "response")</pre>
# Define a function to remove outliers
remove_outliers <- function(x, na.rm = TRUE) {</pre>
  qnt <- quantile(x, probs=c(.25, .75), na.rm = na.rm)</pre>
  H \leftarrow 1.5 * IQR(x, na.rm = na.rm)
  x[x < (qnt[1] - H)] <- NA
  x[x > (qnt[2] + H)] \leftarrow NA
}
colors <- c("orange", "darkgreen")</pre>
colors <- colors[as.numeric(wcgs$chd)]</pre>
shapes = c(17, 18)
shapes <- shapes[as.numeric(wcgs$chd)]</pre>
# Remove outliers for weight and chol
wcgs$weight <- remove_outliers(wcgs$weight)</pre>
wcgs$chol <- remove_outliers(wcgs$chol)</pre>
scatterplot3d(wcgs$weight, wcgs$chol, wcgs$pred_chd_2,
               pch = 16, angle=55, type = "p", cex.symbols = 0.5,
               xlab="Weight [in lbs]",
               ylab="Fasting serum cholesterol [in mg/dl]",
```

3D Scatter Plot



Subexercise 2.4

All coefficients are statistically significant. Weight as predictor has a positive coefficient in the glm. That means that with increasing weight, the probability of suffering from a coronary heart disease increases (holding other variables constant). Similarly, weight also has a positive coefficient in the glm, meaning that higher cholesterol levels are associated with a higher probability of suffering from a coronary heart disease (holding other variables constant). As the coefficient for the interaction term is negative, the effect of weight on the probability of suffering from a coronary heart disease is weaker as cholesterol levels increase. In other words, the effect of weight is moderated by the level of cholesterol. As cholesterol levels increase, the effect of weight on the probability of coronary heart disease becomes weaker. This can be seen in the 3D-plots.