Chickweight Plotting Exercise

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I. Use the ChickWeight R dataset to examine if there is an effect of diet on chick growth:

```
chickweight=ChickWeight
chickweight <- within(chickweight,{
   Chick <- factor(Chick)
   #Time <- factor(Time)
   Diet <- factor(Diet)
})</pre>
```

1. Plot and examine if there is a statistical difference between the weights of chicks assigned to different diet treatments on day 0.

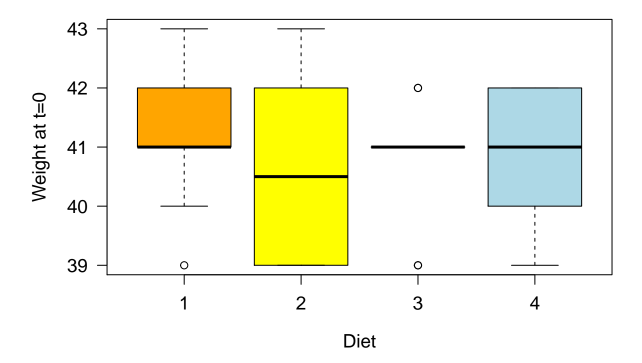
```
dayOdf <- chickweight[chickweight$Time==0,]
summary(dayOdf) #there are four diets</pre>
```

```
##
                                  Chick
                                          Diet
       weight
                       Time
##
         :39.00
                       :0
                                     : 1
                                          1:20
  1st Qu.:41.00
                  1st Qu.:0
                              16
                                     : 1
                                          2:10
## Median :41.00
                  Median :0
                              15
                                     : 1
                                          3:10
## Mean
         :41.06
                              13
                                          4:10
                 Mean :0
                                     : 1
## 3rd Qu.:42.00
                  3rd Qu.:0
                                     : 1
          :43.00
## Max.
                              20
                                     : 1
                  Max. :0
##
                              (Other):44
```

```
summary(anova1 <- aov(day0df$weight~day0df$Diet))</pre>
```

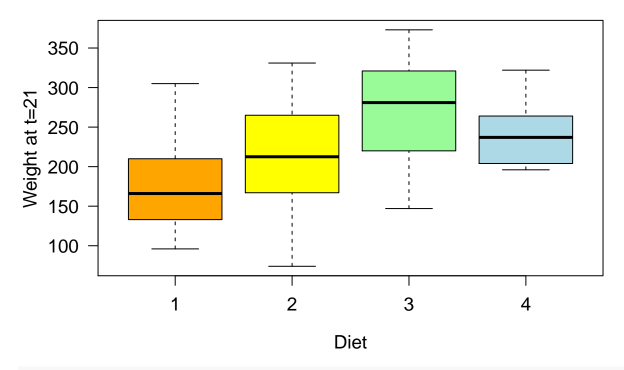
```
## Df Sum Sq Mean Sq F value Pr(>F)
## dayOdf$Diet 3 4.32 1.440 1.132 0.346
## Residuals 46 58.50 1.272
```

```
plot(day0df$Diet, day0df$weight, # data to plot
   ylab="Weight at t=0",xlab="Diet", # label axes
   col=c("orange","yellow","pale green","light blue"),las=1, cex.axis=1.2,cex.lab=1.2)
```



2. Plot and examine if there is a statistical difference between the weights of chicks assigned to different diet treatments at the end of the study (day 21).

```
day21df <- chickweight[chickweight$Time==21,]</pre>
summary(day21df) #there are four diets
##
        weight
                          Time
                                       Chick
                                                Diet
           : 74.0
##
   Min.
                     Min.
                            :21
                                   13
                                          : 1
                                                1:16
    1st Qu.:167.0
                     1st Qu.:21
                                   9
                                                2:10
##
                                          : 1
    Median :205.0
                     Median:21
                                          : 1
##
                                   20
                                                3:10
           :218.7
##
    Mean
                     Mean
                            :21
                                   10
                                          : 1
                                                4: 9
##
    3rd Qu.:266.0
                     3rd Qu.:21
                                   17
                                          : 1
##
    Max.
           :373.0
                     Max.
                            :21
                                   19
                                          : 1
                                   (Other):39
##
summary(anova1 <- aov(day21df$weight~day21df$Diet))</pre>
##
                Df Sum Sq Mean Sq F value Pr(>F)
                 3 57164
                             19055
                                      4.655 0.00686 **
## day21df$Diet
## Residuals
                 41 167839
                              4094
## ---
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
plot(day21df$Diet, day21df$weight, # data to plot
    ylab="Weight at t=21",xlab="Diet", # label axes
      col=c("orange","yellow","pale green","light blue"),las=1, cex.axis=1.2,cex.lab=1.2)
```



TukeyHSD(anova1)

```
Tukey multiple comparisons of means
##
##
       95% family-wise confidence level
##
## Fit: aov(formula = day21df$weight ~ day21df$Diet)
##
## $`day21df$Diet`
##
            diff
                        lwr
                                  upr
                                           p adj
## 2-1
       36.95000
                  -32.11064 106.01064 0.4868095
## 3-1
       92.55000
                   23.48936 161.61064 0.0046959
       60.80556
                  -10.57710 132.18821 0.1192661
## 4-1
## 3-2
       55.60000
                  -21.01591 132.21591 0.2263918
                  -54.85981 102.57092 0.8486781
## 4-2
       23.85556
## 4-3 -31.74444 -110.45981 46.97092 0.7036249
```

3. Is there an effect of diet on chick growth? (HINT: you will need to use a repeated measures ANOVA)

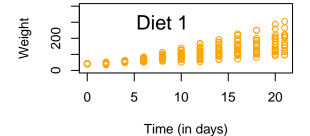
summary(aov(weight~Diet*Time+Error(Chick),data=chickweight))

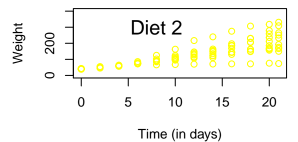
```
##
## Error: Chick
##
             Df Sum Sq Mean Sq F value
                                          Pr(>F)
## Diet
              3 155863
                          51954
                                  7.221 0.000482 ***
                          55175
                                  7.668 0.008197 **
## Time
                 55175
## Diet:Time
             1
                  2482
                           2482
                                  0.345 0.559984
## Residuals 44 316586
                           7195
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Error: Within
```

4. Plot the effects of diet on chick growth.

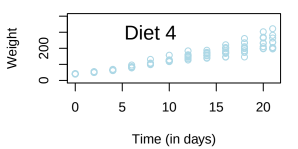
Instead of a simple answer to this question, I present to you a story of struggling to find a more elegant solution.

```
#First, I made separate plots for each diet.
par(mfrow=c(2,2))
chickcols <- c("orange","yellow","pale green","light blue")
for (i in 1:4){
   dietdf <- chickweight[chickweight$Diet==i,]
   plot(dietdf$Time,dietdf$weight,ylim=c(0,400),col=chickcols[i],xlab="Time (in days)",ylab="Weight")
text(8,300,paste('Diet', i),cex=1.5)
}</pre>
```

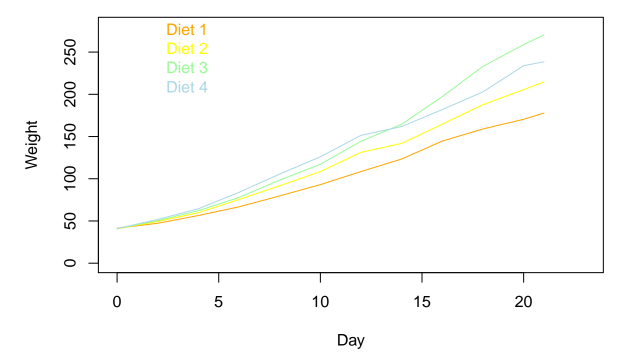








```
weights_for_this_day <- dietdf[dietdf$Time == day, 1] #take the first column (weights) from the die
    mean_weight_per_day[day+1] <- mean(weights_for_this_day, na.rm = T) #take the average of those weig
}
mean_weight_per_day <- mean_weight_per_day[which(!is.nan(mean_weight_per_day))] #remove the NaNs
#days <- c(seq(from = 0, to = max(chickweight$Time), by = 2), 21)
days <- unique(chickweight$Time) #Same as above
points(mean_weight_per_day~days, type = "l", col = chickcols[as.numeric(dietTypes)])
}
legend(1,300,paste("Diet",levels(chickweight$Diet)),bty='n',text.col=chickcols)</pre>
```

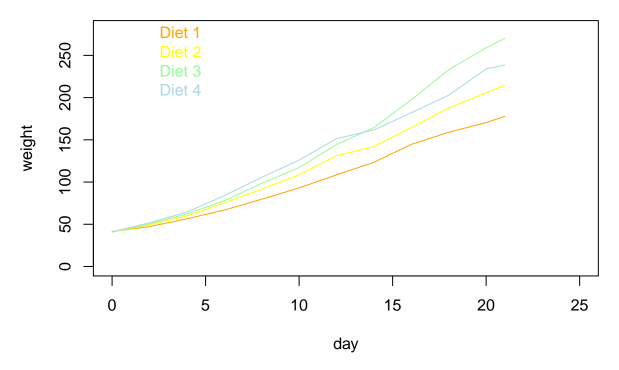


```
#Gaurav helped me come up with alternative and more elegant solutions.

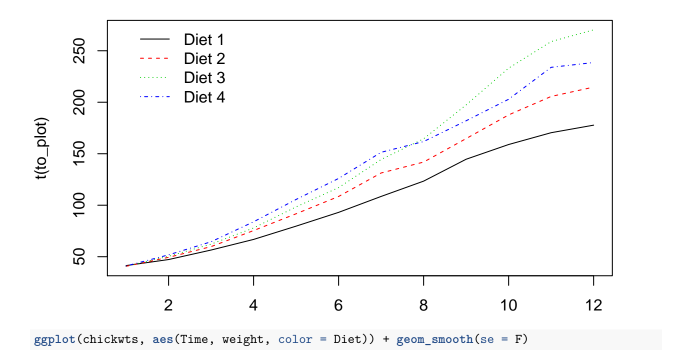
#The first uses dplyr:
library(dplyr)
```

```
##
## Attaching package: 'dplyr'
##
## The following objects are masked from 'package:stats':
##
##
       filter, lag
##
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
chickwts <- chickweight %>% group_by(Diet, Time) %>% summarise(weight = mean(weight))
plot(1, type = "n", xlim = c(0, 25), ylim = c(0, 280),
     xlab = "day", ylab = "weight")
for(dietTypes in unique(chickwts$Diet)) {
```

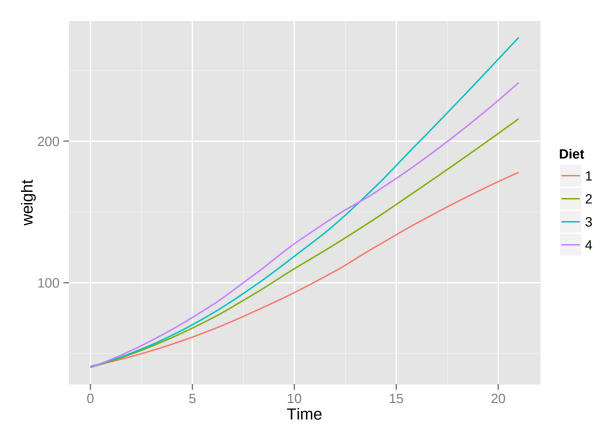
```
current_diet <- chickwts %>% filter(Diet == dietTypes)
lines(current_diet$weight~current_diet$Time,col=chickcols[as.numeric(dietTypes)])
}
legend(1,300,paste("Diet",levels(chickweight$Diet)),bty='n',text.col=chickcols)
```



```
#The second uses aggregate:
xx <- aggregate(chickwts$weight, by = list(chickwts$Diet, chickwts$Time), FUN = mean)
to_plot <- matrix(xx$x, nrow = 4)
matplot(t(to_plot), type = "l")
legend(1,280,paste("Diet",levels(chickweight$Diet)),bty='n',lty=1:4,col=1:4)
#I knew the lty and col of the lines because the default in matplot is lty = 1:5 and col = 1:6
#The third uses ggplot2:
library(ggplot2)</pre>
```

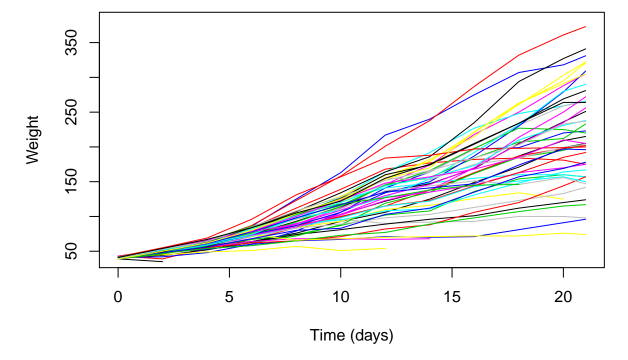


geom_smooth: method="auto" and size of largest group is <1000, so using loess. Use 'method = x' to classical description of the size of largest group is <1000, so using loess.



5. Using a 'for' loop, plot the growth of each chick in a different color, all on the same plot.

```
par(mfrow=c(1,1))
chickweight$Time <- as.numeric(chickweight$Time)
chicki <- chickweight[chickweight$Chick=1,]
plot(chicki$Time,chicki$weight,type='l',ylim=c(40,380),xlab="Time (days)",ylab="Weight")
chicki <- NA
chickcols <- 1
for (i in 2:length(levels(chickweight$Chick))) {
   chicki <- chickweight[chickweight$Chick=i,]
   lines(chicki$Time,chicki$weight,col=chickcols)
   chickcols <- chickcols + 1
}</pre>
```



II. If we made it to the population growth part, using the model we wrote in class plot the population dynamics of 5 different populations, each with a different growth rate, all on the same plot.

We didn't get to it in class but I worked on it anyway:

```
simDiscLog <- function(NO,RR,ttMax,color) {
   NN <- matrix(NA, nrow=1, ncol=ttMax+1)
   NN[1] <- NO # set first value to initial condition
   # use a loop to iterate the model the desired number of times
   for (tt in 1:ttMax) {
        NN[tt+1] <- RR*NN[tt]
   }
   points(1:(ttMax+1), NN, xlab="Time", ylab="Population size (N)", col=color, pch=16, las=1)
   # add a line:
   lines(1:(ttMax+1), NN, xlab="Time", ylab="Population size (N)", col=color, las=1)
}

plot(1, type = "n", xlim = c(0, 13), ylim = c(0, 300),
        xlab = "Time", ylab = "Population size (N)")</pre>
```

```
simDiscLog(100,1.08,10,"red")
simDiscLog(100,1.0,10,"orange")
simDiscLog(100,0.80,10,"yellow")
simDiscLog(100,0.5,10,"green")
simDiscLog(100,0.23,10,"blue")
legend(-1,300,bty='n',c("Rate = 1.08","Rate = 0.8","Rate = 0.5","Rate = 0.23"),text.col=c(
```

