IA1

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```
knitr::opts_chunk$set(message=FALSE, warning = FALSE)
```

Enriched diet affects zebrafish size

To study the effects of an enriched diet on mean zebrafish size, 200 freshly hatched zebrafish were taken from the same clutch and randomly assigned to an enriched or a control diet (unenriched). The diet was administered for two months at which point the standard length (mm) and mass (kg) of each fish was measured. Results were compiled in a tab-separated file containing the following columns: "Individual", "Diet", "SL", and "Weight."

The following are the hypothesis to be tested in this experiment:

 H_0 : Feeding zeebrafish an enriched diet will not significantly affect their mean size after two months

 H_A : Feeding zebrafish an enriched diet will significantly affect their mean size after two months

Exploratory Data Analysis

```
library(tidyverse)
library(magrittr)
library(ggpubr)
library(knitr)
library(lmodel2)
library(broom)
zfish_raw <- read_tsv('zfish_diet_IA.tsv')
head(zfish_raw)</pre>
```

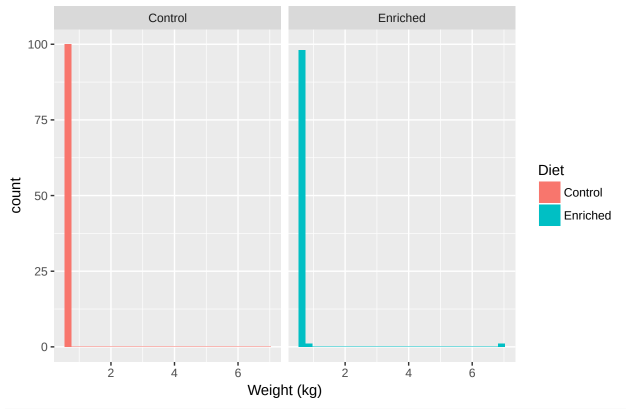
```
## # A tibble: 6 x 4
##
    Individual
                   Diet
                           SL Weight
##
         <int>
                  <chr> <dbl> <dbl>
## 1
               Control 3.58
                               0.554
## 2
             2 Enriched 4.66 0.699
## 3
             3 Enriched 4.50 0.644
## 4
             4 Control 3.71 0.624
## 5
             5
                Control 4.36 0.688
## 6
             6 Control 3.89 0.613
```

Histograms

Let's take a look at some of the ways the data are distributed. First we can look at histograms of continuous variables weight and standard length

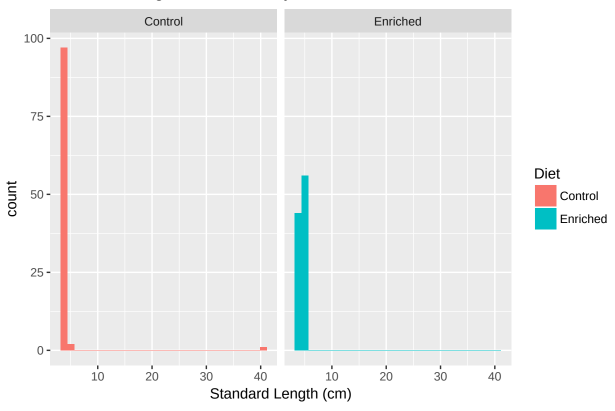
```
ggplot(zfish_raw, aes(fill = Diet))+
  geom_histogram(aes(Weight))+
  facet_wrap(~Diet)+
  xlab('Weight (kg)')+
  ggtitle('Weight Distribution by Diet')
```

Weight Distribution by Diet



```
ggplot(zfish_raw, aes(fill = Diet))+
geom_histogram(aes(SL))+
facet_wrap(~Diet)+
xlab('Standard Length (cm)') +
ggtitle('Standard Length Distribution by Diet')
```

Standard Length Distribution by Diet



There appear to be erroneous data that is obscuring the distribution of both SL, and Weight. I will remove the obvious errors from the dataset as shown below:

```
zfish_correct <- zfish_raw %>%
  filter(., Weight < 4 & SL < 20)</pre>
# total SL hist colored by diet
dt_n <- ggplot(zfish_correct, aes(SL, fill = Diet))+</pre>
            geom_histogram(binwidth = 0.05)
# SL hist faceted by diet
dt <- ggplot(zfish_correct, aes(SL, fill = Diet)) +</pre>
          geom_histogram(binwidth = 0.05)+
          facet_grid(Diet~.)+
          xlab('Standard Length (cm)') +
  theme(
  strip.background = element_blank(),
  strip.text.x = element_blank(),
  strip.text.y = element_blank()
# total weight hist colored by diet
wt_n <- ggplot(zfish_correct, aes(Weight, fill = Diet)) +</pre>
            geom_histogram(binwidth = 0.01)+
```

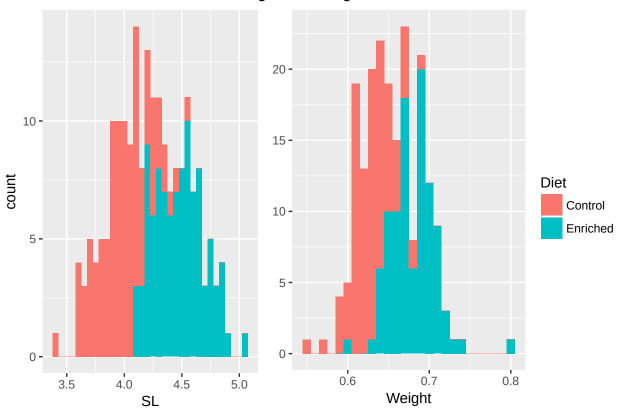
```
theme(axis.title.y = element_blank())

# weight hists faceted by diet

wt <- ggplot(zfish_correct, aes(Weight, fill = Diet)) +
    geom_histogram(aes(Weight, fill = Diet), binwidth = 0.007) +
    facet_grid(Diet~.)+
    xlab('Weight (kg)') +
    theme(
    strip.background = element_blank(),
    strip.text.x = element_blank(),
    strip.text.y = element_blank(),
    axis.title.y = element_blank()
)

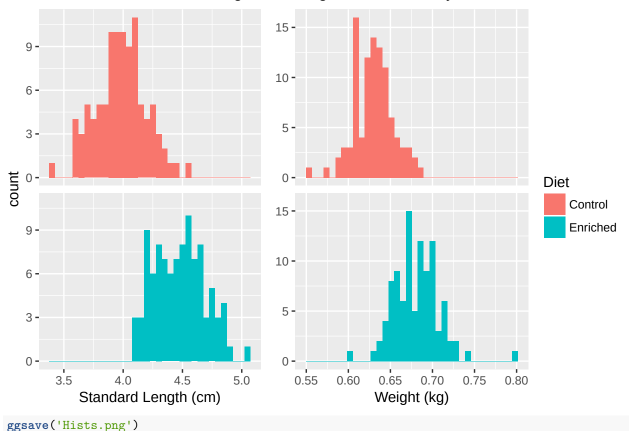
# group figures
ggarrange(dt_n, wt_n, common.legend = TRUE, legend = 'right') %>% annotate_figure(., top = text_grob("S
```

Standard Length and Weight Distributions



ggarrange(dt, wt, common.legend = TRUE, legend = 'right') %>% annotate_figure(., top = text_grob("Stand

Standard Length and Weight Distributions by Diet



Okay that certainly changed the histograms! These data seem to be fairly normally distributed. The histograms of continuous variables split by diet show a positive shift in both continuous variables for the enriched diet compared to the control, let's do some statistics to back this up!

Boxplots

First similar comparisons will be made using boxplots.

```
bp_d <- ggplot(zfish_correct, aes(Diet, SL, fill = Diet))+
  geom_boxplot() +
  ylab('Standard Length (cm)')+
  theme(
    axis.title.x = element_blank()
)

bp_w <- ggplot(zfish_correct, aes(Diet, Weight, fill = Diet))+
  geom_boxplot()+
  ylab('Weight (kg)')+
  theme(
    axis.title.x = element_blank()
)

ggarrange(bp_d, bp_w, common.legend = TRUE, legend = 'right') %>% annotate_figure(., top = text_grob("...))
```



Again we see the overall increase in both metrics with an enriched diet as compared to the control.

Summary statistics

And a summary table to compare relevant statistics:

```
Summ_stats <- zfish_correct %>%
  group_by(Diet) %>%
  summarise_at(vars(SL, Weight), c('mean', 'sd', 'var'))

kable(Summ_stats, align = 'c', caption = 'Summary statistics for zebra fish measuremnets by diet', digi
```

Table 1: Summary statistics for zebra fish measuremnets by diet

Diet	SL_mean	$Weight_mean$	SL_sd	$Weight_sd$	SL_var	$Weight_var$
Control Enriched	3.982 4.473	$0.630 \\ 0.681$	0.211 0.213	0.023 0.027	0.045 0.046	0.001 0.001

T Test

Because the data look to be normally distributed, appear to have equal variances (var[control] $\sim=$ var[diet]), and occur as independent observations, the distributional assumptions are met to perform a parametric T Test. The T Test will test a difference in the means of the Control and Enriched treatments for both weight, and standard length.

```
# T tests for each
sl_T <- t.test(zfish_correct$SL ~ zfish_correct$Diet)</pre>
wt T <- t.test(zfish correct$Weight ~ zfish correct$Diet)
# format for reporting t test
t.report <- function(tt){</pre>
  tvalue <- tt$statistic %>% formatC(digits = 2, format = "f")
  pvalue <- tt$p.value %>% formatC(digits = 2, format = "E")
  if (round(tt$parameter, 0) == tt$parameter) {
    df <- tt$parameter</pre>
  } else {
    df <- formatC(tt$parameter, digits = 2, format = "f")</pre>
  if (tt$p.value < 0.0005) {
    pvalue <- " < 0.001"
  } else {
    if (tt$p.value < 0.005) {
      pvalue <- paste0(" = ",tt$p.value %>% formatC(., digits = 3, format = "f"))
      pvalue <- paste0(" = ",tt$p.value %>% formatC(., digits = 2, format = "f"))
    }
  paste0("*t*(",df,") = ",tvalue, ", *p*", pvalue)
```

Now the statement can be made that zebrafish have a significantly higher standard length when fed an enriched diet as compared to a normal diet, t(195.98) = -16.27, p < 0.001, and have a significantly higher weight when fed an enriched diet as compared to a normal diet, t(190.90) = -14.08, p < 0.001. The null hypothesis that there is no significant effect of an enriched diet on mean fish size can be rejected.

Non-parametric T test

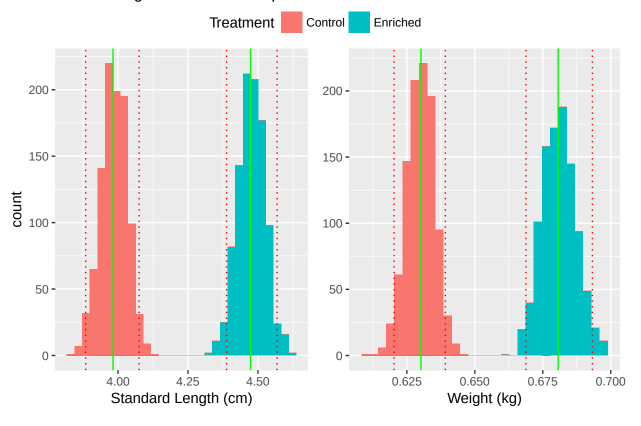
The hypothesis testing can be done using a non-parametric parameter estimation (calculated by resampling of the means) and then performing a T-test. This resampling will cooerce the distribution of the parameter estimate into normality (if the data are not already normally distributed) before performing hypothesis tests, such that our confidence interval about the hypothesis test is valid.

```
Control <- zfish_correct %>% filter(., Diet == 'Control')
Enriched <- zfish_correct %>% filter(., Diet == 'Enriched')
# Standard length stats and distribtion
sl_bootC_st <- boot(Control$SL)[1]</pre>
sl_bootC_dist <- boot(Control$SL)[2]</pre>
sl bootE st <- boot(Enriched$SL)[1]</pre>
sl_bootE_dist <- boot(Enriched$SL)[2]</pre>
# combine
sl_bootC <- tbl_df(data.frame(sl_bootC_dist, rep('Control', 1000)))</pre>
colnames(sl_bootC) <- c('boot_means_sl', 'Treatment')</pre>
sl_bootE <- tbl_df(data.frame(sl_bootE_dist, rep('Enriched', 1000)))</pre>
colnames(sl_bootE) <- c('boot_means_sl', 'Treatment')</pre>
# stack
sl_boot <- rbind(sl_bootC, sl_bootE)</pre>
# Weight stats and distribution
wt bootC st <- boot(Control$Weight)[1]</pre>
wt_bootC_dist <- boot(Control$Weight)[2]</pre>
wt bootE st <- boot(Enriched$Weight)[1]</pre>
wt_bootE_dist <- boot(Enriched$Weight)[2]</pre>
#combine
wt_bootC <- tbl_df(data.frame(wt_bootC_dist, rep('Control', 1000)))</pre>
colnames(wt_bootC) <- c('boot_means_wt', 'Treatment')</pre>
wt_bootE <- tbl_df(data.frame(wt_bootE_dist, rep('Enriched', 1000)))</pre>
colnames(wt_bootE) <- c('boot_means_wt', 'Treatment')</pre>
# stack
wt_boot <- rbind(wt_bootC, wt_bootE)</pre>
# plot resamples means for standard length
resampled_sl <- ggplot(sl_boot, aes(boot_means_sl, fill = Treatment))+
  geom histogram()+
 xlab('Standard Length (cm)')+
  geom_vline(xintercept = sl_bootE_st[[1]]$bm.lower.ci, color = 'red', linetype="dotted") +
  geom_vline(xintercept = sl_bootE_st[[1]]$bm.upper.ci, color = 'red', linetype="dotted") +
  geom_vline(xintercept = sl_bootC_st[[1]]$bm.lower.ci, color = 'red', linetype="dotted") +
  geom_vline(xintercept = sl_bootC_st[[1]]$bm.upper.ci, color = 'red', linetype="dotted") +
  geom_vline(xintercept = Summ_stats$SL_mean, color = 'green')
resampled_wt <- ggplot(wt_boot, aes(boot_means_wt, fill = Treatment))+
```

```
geom_histogram()+
    xlab('Weight (kg)')+
    geom_vline(xintercept = wt_bootE_st[[1]]$bm.lower.ci, color = 'red', linetype="dotted") +
    geom_vline(xintercept = wt_bootE_st[[1]]$bm.upper.ci, color = 'red', linetype="dotted") +
    geom_vline(xintercept = wt_bootC_st[[1]]$bm.lower.ci, color = 'red', linetype="dotted") +
    geom_vline(xintercept = wt_bootC_st[[1]]$bm.upper.ci, color = 'red', linetype="dotted") +
    geom_vline(xintercept = Summ_stats$Weight_mean, color = 'green') +
    theme(
        axis.title.y = element_blank()
)

ggarrange(resampled_sl, resampled_wt, common.legend = TRUE) %>% annotate_figure(., top = text_grob("His
```

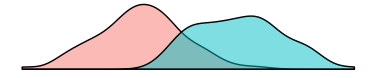
Histograms of the resampled means with confidence intervals

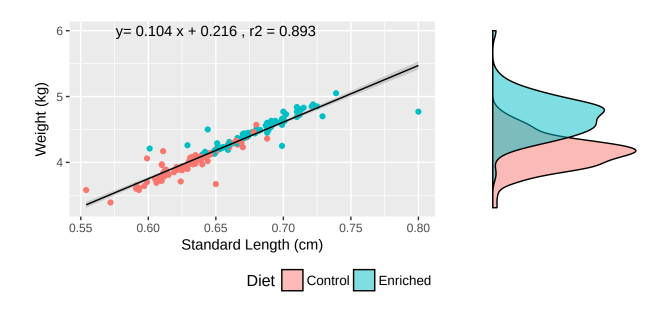


SL AND WEIGHT

```
BIC
                           deviance df.residual
## 1 -1196.251 -1186.386 0.02674031
r2 <- format(data.frame(glance(zfish_lm)\squared)[[1]], digits = 3)
eq <- paste("y=", m, "x +", b,',', 'r2', "=", r2)
# plot the relationship between standard lenght and weight
corr <- ggplot(zfish_correct, aes(zfish_correct$Weight,</pre>
                          zfish_correct$SL,
                          color = Diet)) +
  geom_point() +
  geom_smooth(method = 'lm', color = 'black', size = 0.5) +
  xlab('Standard Length (cm)') +
  ylab('Weight (kg)')+
  annotate('text',x = 0.65, y = 6, label = eq)
sl_dens <- ggdensity(zfish_correct, 'SL', fill = 'Diet') + clean_theme()</pre>
wt_dens <- ggdensity(zfish_correct, 'Weight', fill = 'Diet') + rotate() + clean_theme()
# Arranging the plot
ggarrange(sl_dens, NULL, corr, wt_dens,
          ncol = 2, nrow = 2, align = "hv",
          widths = c(2, 1), heights = c(1, 2),
          common.legend = TRUE, legend = 'bottom') %>% annotate_figure(top = text_grob('The relationshi
```

The relationship between Weight and Standard length over a change in diet

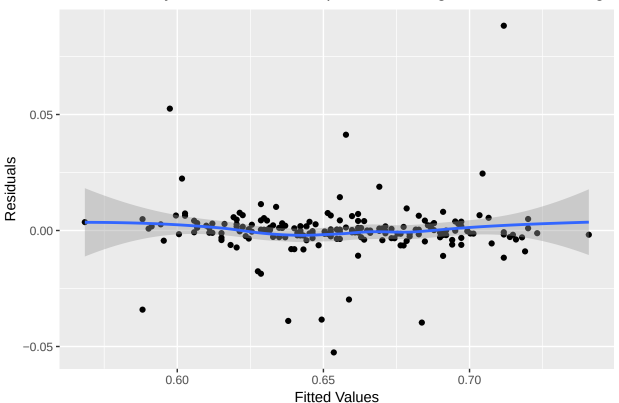




```
# combine residuals vs. fitted values
res <- tbl_df(data.frame(cbind(zfish_lm$residuals, zfish_lm$fitted.values)))

ggplot(res, aes(X2,X1)) +
   geom_point() +
   geom_smooth(mode = 'lm') +
   xlab('Fitted Values')+
   ylab('Residuals')+
   ggtitle('Residual analysis of the relationship between weight and standard length')</pre>
```

Residual analysis of the relationship between weight and standard length

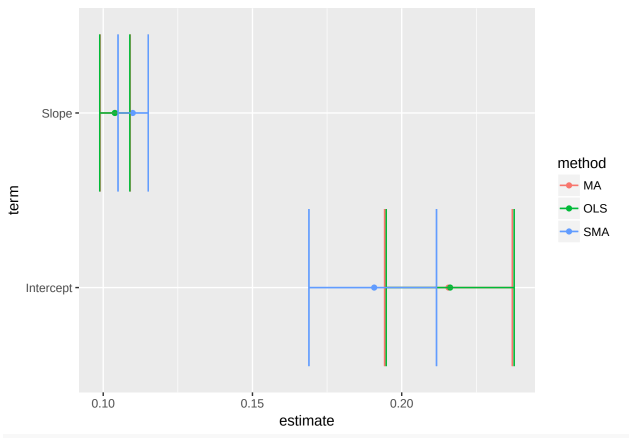


```
# type II model fit
zfish_lm2 <- lmodel2(zfish_correct$Weight~zfish_correct$SL)
tidy(zfish_lm2)</pre>
```

```
##
    method
                 term estimate
                                  conf.low conf.high
## 1
        MA Intercept 0.2156807 0.19427461 0.2370645
## 2
                Slope 0.1040081 0.09895024 0.1090711
## 3
       OLS Intercept 0.2162407 0.19481062 0.2376707
## 4
                Slope 0.1038756 0.09882167 0.1089295
       OLS
## 5
        SMA Intercept 0.1907835 0.16892498 0.2116598
## 6
       SMA
                Slope 0.1098969 0.10495909 0.1150670
glance(zfish_lm2)
```

```
## r.squared theta p.value H
## 1 0.8934216 0.701477 3.097717e-97 2.50623e-05
```

```
ggplot(tidy(zfish_lm2), aes(estimate, term, color = method)) +
geom_point() +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high)) +
geom_errorbarh(aes(xmin = conf.low, xmax = conf.high))
```



tidy(zfish_lm2)

```
##
     method
                 term estimate
                                   conf.low conf.high
## 1
         MA Intercept 0.2156807 0.19427461 0.2370645
## 2
                Slope 0.1040081 0.09895024 0.1090711
         MA
## 3
        OLS Intercept 0.2162407 0.19481062 0.2376707
## 4
        OLS
                Slope 0.1038756 0.09882167 0.1089295
## 5
        SMA Intercept 0.1907835 0.16892498 0.2116598
## 6
        {\tt SMA}
                Slope 0.1098969 0.10495909 0.1150670
```

There is a significant relationship between Weight(kg) and Standard Length (cm) F1,196 = 1643, p < 0.001