# STATS 5620 Final Report

# Linking Diet and Reproductive Success in Northwest Atlantic Grey Seals

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Data Analysis STAT 4620/5620 Winter 24-25"

Link to GitHub Repository: https://github.com/MaxHA17/STAT5620.Project

Abstract: Understanding how top predators use different foraging strategies to navigate and adapt to patchy and unpredictable prey availability has important implications for individual fitness and population dynamics. Here we explore the relationship between diet and reproductive success in a capital breeding pinniped, the northwest Atlantic grey seal (*Halichoerus grypus*) through two research questions: 1. Does variation in diet influence maternal mass change over the foraging period leading up to parturition in Northwest Atlantic grey seals; and 2. Does female diet predict pup weaning mass in Northwest Atlantic grey seals? Using linear models (LMs), generalized linear models (GLMs), and generalized linear mixed effects models (GLMMs), we found that females who primarily fed on pollock experienced lower mass change and pup weaning masses than those who primarily fed on cod, sand lance, or redfish. While models for both questions were well fit, model predictive errors were relatively high, possibly indicating that missing covariates outside of diet are playing a significant role in explaining maternal mass change and pup weaning mass.

**Keywords:** Capital breeding strategy, Generalized Linear Models, Pinnipeds, Quantitative Fatty Acid Signature Analysis, Reproductive Ecology

# Introduction

For marine predators, prey distribution may vary unpredictably in time and space, undergoing seasonal, inter-annual, and longer-term changes in environmental conditions. Understanding how top predators use different foraging strategies to navigate and adapt to patchy and unpredictable prey availability has important implications for individual fitness and population dynamics, yet studies exploring the consequences of such decisions on fitness are relatively rare (Nathan et al., 2008).

Vertebrates fuel the costs of reproduction along a continuum of an income-capital breeding strategy, where animals that use an income breeding strategy rely on concurrent energy accumulation during the breeding period as opposed to a capital breeding strategy, where energy acquired and stored prior to the breeding period is used solely to finance major reproductive costs (Stephens et al., 2009; Stephens et al., 2014). A capital breeding strategy is thought to be advantageous for large animals capable of carrying large energy stores and offers energetic benefits in areas of patchy or unreliable food availability and allows parents to direct more resources (e.g. time and energy) towards offspring, as opposed to foraging, during the breeding and lactation period (Stephens et al., 2009; Stephens et

al.,2014). Given the importance of pre-breeding and lactation period foraging success and energy acquisition in capital breeders, these species offer interesting model systems to study the relationships between diet and reproductive success.

Phocid seals are wide-ranging, large-bodied, and long-lived marine predators, with individuals of many species foraging across a wide range of habitats over large temporal scales (Bowen and Jonsen, 2022). Grey seals (Halichoerus grypus) are a long-lived, sexually dimorphic, phocid species and are capital breeders. In the Sable Island population, pregnant females haul out in late December or early January, give birth to a single pup and nurse that pup for 16-18 days, relying solely on energy accumulated during the foraging period prior to parturition to support the costs of lactation (Iverson et al., 1993; Stephens et al., 2009). At or near the abrupt weaning of that pup, females are mated and return to sea to replenish spent body reserves and, after a several month period of delayed implantation, to support gestation and most importantly preparation for the subsequent December or early January lactation period. As in all mammals, the costs of lactation, and its contribution to reproductive success, far exceed the costs of gestation. Like most other phocids, grey seals are generalist predators (Bowen and Jonsen, 2022), but females tend to feed on a narrower range of energy rich species (predominantly sand lance, redfish, and other pelagic fishes) following the lactation period and expand their prey diversity in the several months leading up to the December-January breeding and lactation period (Becket al., 2007). Female grey seals reach sexual maturity at age 4-5 years and can continue to reproduce as late as into their early 40s (Bowen et al., 2006). During the brief 16–18-day lactation period, females utilize approximately 25-38% of parturition body mass and daily maternal mass loss is a significant predictor of pup growth rates during lactation, with heavier mothers at parturition weaning heavier pups (Iverson et al., 1993; Mellish et al., 1999). Pup survival is positively related to pup weaning mass up to the mean population weaning mass of 51.5 kg, as predicted by a bigger-is-better hypothesis for size selective mortality (Bowen et al., 2015). The phenomena of heavier females weaning larger offspring is evident even from primiparity where it is believed that heavier females have larger resource stores that they can mobilize during lactation to produce larger offspring at weaning (Iverson et al., 1993; Mellish et al., 1999; Bowen et al., 2006). Following the lactation period, parental care abruptly ends and the pup is left to fast and survive for several weeks on the energy acquired during lactation before undertaking its first foraging trip (Noren et al., 2008). Thus, answering questions relating to the diet of females prior to parturition will improve our understanding of the importance of habitat use and energy acquisition as they relate to life history characteristics, such as reproductive success, in these capital breeders.

Northwest Atlantic grey seals breeding on Sable Island, Nova Scotia, tend to be central place foragers and have high fidelity to Sable Island for breeding and as a haul-out site, making them excellent candidates for long-term studies (Bowen et al., 2015). Using diet estimations methods, such as quantitative fatty acid (FA) signature analysis (QFASA; Iverson et al., 2004, Beck et al., 2007), it becomes possible to explore the relationship between diet variation and reproductive success at the individual level. QFASA is used to estimate predator diet by comparing the FA signatures of predator adipose tissue to those of candidate prey species (Iverson, 2009; Karnovsky et al., 2012). Predator FA signatures alone have historically been used to provide a qualitative assessment of spatial and temporal patterns in diet diversity. However, since many FAs that are biosynthesized within unique prey species accumulate in predator adipose tissue over time, evaluating the amount of prey-specific FAs relative to the predator's overall FA stores can provide a quantitative estimate of diet, known as QFASA (Iverson

etal., 2004). For accurate diet estimation, species-specific calibration coefficients must be experimentally derived to account for the altered incorporation of prey FAs into the predator adipose tissue due to lipid metabolic processes within the predator (Iverson et al., 2004; Karnovsky et al., 2012). When calibration coefficients are derived and the FA signature of many or all potential prey species in an ecosystem is known, as in the case of the Northwest Atlantic grey seal range (Iverson et al., 2004; Beck et al., 2007), QFASA can provide a statistical approach to accurately depicting diet and prey diversity over ecologically relevant time scales (e.g., weeks to several months). This study will test the hypothesis that differences in diet will affect an individual's ability to store energy prior to parturition, and thus influence their reproductive success. Specifically, we will explore two research questions:

- 1. Does variation in diet influence maternal mass change over the foraging period leading up to parturition in Northwest Atlantic grey seals?
- 2. Does female diet predict pup weaning mass in Northwest Atlantic grey seals?

### **Data description**

Response variables (organized by research question):

- 1. Maternal mass change during the pre-breeding season foraging period: continuous variable ranging from -45 88 kg. Calculated by subtracting the maternal recovery mass from her deployment mass. Insert plot of mass change distribution here?
- 2. Pup weaning mass: continuous variable ranging from 34.5 63 kg. Denotes the mass of the pup after the mother has terminated the lactation period and departed the pup and/or Sable Island. Insert plot of pup weaning mass distribution here?

# Predictor or supporting variables

- 1. Year: discrete variable with values ranging from 1996-2015. Denotes the year in which each observation was collected.
- 2. Maternal ID: discrete variable with a unique identifier for each individual female.
- 3. Pup sex: categorical; male or female.
- 4. Maternal age: discrete variable with values ranging from 7-40 years old.
- 5. Maternal deployment mass: continuous variable with values ranging from 116 250 kg. Describes the female mass when she was located pregnant on Sable Island 3-6 months prior to parturition and prior to undergoing her pre-parturition foraging period.
- 6. Maternal mass at parturition: continuous variable ranging from ~ 137 250 kg. This is the mass of the female after returning to Sable Island and giving birth to her pup.

- 7. Maternal dominant prey species: discrete variable with 6 unique values. Represents the species making up the greatest proportion of the female diet.
- 8. Maternal dietary energy density: continuous variable ranging from 4.88 6.51 kJ/g. Energy density (kJ/g) of the female diet is calculated from the average energy content per gram of prey body tissue.
- 9. Maternal diet diversity: continuous variable 0.18 0.58, no specific units. Diversity is calculated using a standardized Shannon-Weaver diversity index to quantify individual diet diversity.

# Method

The initial step was data cleaning to ensure all readily apparent data errors were rectified prior to data analysis and model fitting. Most of the data errors discovered resulted from the absent of critical data (i.e. missing covariates) and these observations were removed from the data-set. As a result, the first question data-set resulted in 56 observations (n=56) and the second question resulted in 76 observations (n=76). It should also be noted that additional observations were filtered from the data-set throughout the analysis process due to the identification of outliers with high leverage. A description of outlier identification and removal can be found in the analysis section below. Thus, the final data-set used for question one had 72 observation and the final data-set for question two had 50 observations.

Once cleaned and uploaded to RStudio, initial data plots were made using the 'flexplot' and 'plot\_explore' R function we have included in our Git Hub. Description of 'plot\_explore' function created in this R package can be found within the Vignettes in the STAT5620. Project package. These visualizations facilitated the exploration of variable distributions, specifically the response variables, and the apparent relationships between covariates and each response variable. From this it was determined that the response variables for both questions, Maternal Mass Change and Pup Weaning Mass, were Gaussian distributed. This normality allowed the exploration of the data through General Linear Models as well as more complex Generalized Linear Models and Mixed Models. Although our original analytical approach proposed the use of Generalized Linear Mixed Models, it felt prudent to explore the efficiency, interpretability and computational simplify of Linear Models if the assumption were met and the data suggested linearity.

Both questions were analysed first as Linear Models (LM), then as Generalized Linear Models (GLM) with Gaussian distributions, and finally as Generalized Linear Mixed Models (GLMM) with random effects and fixed effects of covariates. Initially, Maternal ID was to be included as a random effect to account for repeated measures, individual heterogeneity, and allow for population level inferences to be made using these data. However, a lack of repeated measures (<10 per question) limited our ability to include Maternal ID as a random effect and Maternal ID was assumed to be fixed. For question one in particular, including a random slope for the effect of dominant prey species was explored due to the dominant prey species included in this data-set being a random draw from all possible prey species. However, the goal of this analysis was to explore the direct effect of each prey species on maternal mass gain, resulting in dominant prey species being included as a fixed effect in the final model. With each model type, we began with all the predictor variables prior to using the backward and forward

"Step" function to reduce the number of variables through elimination based on the p-values for Linear Models and AIC for GLM and GLMM.

Once we had established the best fit model for both questions using AIC, the performance of the best models were tested using Cross Validation, question one using 7 folds and question two using 11 folds. For question 1, all modelling and cross-validation was performed using functions included in base R.

# **Analysis**

#### **Research Question 1**

Here we are exploring whether variation in diet influence maternal mass change over the foraging period leading up to parturition in Northwest Atlantic grey seals?

The response variable of interest is maternal mass change. We will first look at the distribution of this variable:

```
#Load the data
library(readr)
Data_Q1<- read.csv("/Users/peterbraithwaite/Desktop/IDPhD Classes/Stat 5620_Updated/Final
#Ensure the data types are correctly assigned
Data_Q1$MomID <- as.factor(Data_Q1$MomID)
Data_Q1$Year <- factor(Data_Q1$Year, levels = sort(unique(Data_Q1$Year)), ordered = TRUE)
Data_Q1$`Dominant.prey.species`<- as.character(Data_Q1$`Dominant.prey.species`)
Data_Q1$`Diet.diversity` <- as.numeric(Data_Q1$`Diet.diversity`)
Data_Q1$`Dietary.energy.density` <- as.numeric(Data_Q1$`Dietary.energy.density`)
summary(Data_Q1)</pre>
```

```
Dietary.energy.density Diet.diversity
    MomID
                  Year
4269
       : 3
            2013
                    :13
                         Min.
                                 :4.881
                                                Min.
                                                        :0.1788
       : 2
                          1st Qu.:5.623
                                                 1st Qu.:0.3063
24
            2011
                    :11
146
       : 2
            2010
                    : 9
                         Median :5.753
                                                Median :0.3572
829
       : 2
             2012
                    : 9
                         Mean :5.736
                                                        :0.3625
                                                Mean
       : 2
3271
            2009
                    : 6
                         3rd Qu.:5.893
                                                 3rd Qu.:0.4181
     : 2
3616
             2015
                    : 5
                                :6.517
                                                Max.
                                                        :0.5782
                         Max.
(Other):63
             (Other):23
Dominant.prey.species Mass.change
Length:76
                      Min.
                             :-40.50
Class :character
                      1st Qu.: 6.25
Mode :character
                      Median : 22.75
                     Mean : 26.02
                      3rd Qu.: 45.00
```

Max. : 88.00

```
#rename columns
library(dplyr)

Attaching package: 'dplyr'

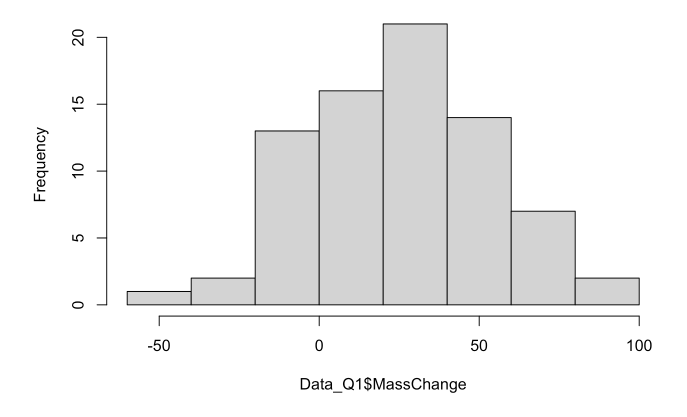
The following objects are masked from 'package:stats':
    filter, lag

The following objects are masked from 'package:base':
    intersect, setdiff, setequal, union

Data_01 <- Data_01 %>%
    rename(
    DietEngDen = `Dietary.energy.density`,
    DietDiv = `Diet.diversity`,
    DomSpp = `Dominant.prey.species`,
    MassChange = `Mass.change`)

hist(Data_01$MassChange)
```

## Histogram of Data\_Q1\$MassChange



There are a total of n=76 observations and mass change appears to follow a normal distribution a Gaussian family distribution is a good starting point for an initial model.

One of the covariates, dominant prey species, is a categorical variable. We will first check the number of observations in each category:

```
table(Data_Q1$DomSpp)
```

AtlanticCod	Capelin	${\tt NorthernSandlance}$	Pollock
6	3	29	5
Redfish	WhiteHake		
32	1		

With only one observation of white hake and 3 observations of capelin, there is likely not enough data to estimate the effect of these species on the response variable and we are unable to draw meaningful conclusions. These points also cause extreme residual values. Capelin and white hake observations will be removed from the data set:

```
Data_Q1 <- Data_Q1 %>%
filter(DomSpp != "WhiteHake")
```

```
Data_Q1 <- Data_Q1 %>%
filter(DomSpp != "Capelin")
```

The initial model seeks to model maternal mass change as a function of dominant prey species, diet diversity, dietary energy density, and deployment year. We will now explore the apparent relationships between each continuous and categorical covariate using the plot\_explore function built and contained in the STAT5620.Project R package:

Initially, it appears that diet diversity is negatively related to mass change, with energy density possibly having a slight negative relationship. It also appears that animals with cod, redfish or sand lance as the dominant prey species gain more mass during foraging than females who primarily prey on Pollock. There does not appear to be any clear trends in mass change over time.

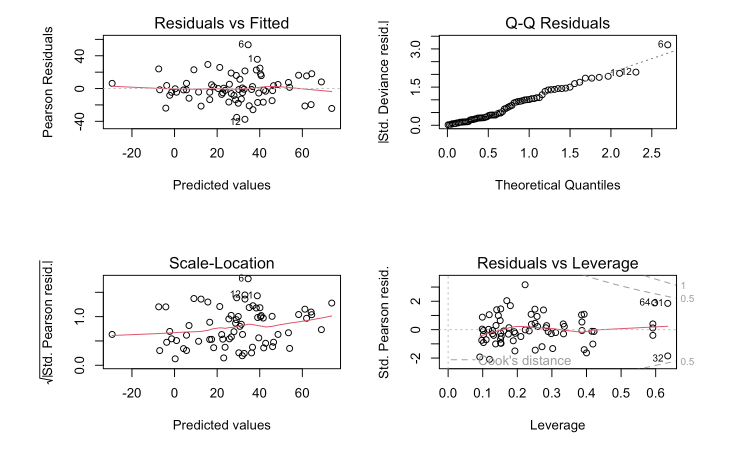
We will begin with an initial model with all covariates of interest. Based on the distribution of mass change, we will begin with a simple model with a gaussian distribution and an identity link function:

```
library(lme4)
```

Loading required package: Matrix

```
mom_mod <- glm(data=Data_Q1, family = gaussian(link="identity"), formula = MassChange ~ D
par(mfrow = c(2, 2))
plot(mom_mod)</pre>
```

Warning: not plotting observations with leverage one: 45



The residual plots from this initial model show good homoscedasticity and normality. There is one observation (observation 6) that appears to be an extreme value. This observation comes from a female that experiences an 88 kg increase in mass over the pre-breeding period foraging period, which is well above the mean mass change value of 28.5. However, none of the covariate values associated with this observation are abnormal and this point does not have high leverage. As a results, the decision is to retain this data point since it is not highly influential and knowledge of the system suggests it is a valid data point. Since the model residuals look good, there is no need to add complexity by using a different link function to generalize the model.

We can now use step selection to preform repeated hypothesis tests for variable selection to determine which covariates best explain maternal mass change while minimizing model complexity and colinearity between covariates.

```
step(mom_mod, direction = "both")
```

Start: AIC=647.55
MassChange ~ DietDiv + DietEngDen + Year + DomSpp

	Df	Deviance	AIC
- DietEngDen	1	19508	645.67
<none></none>		19478	647.55
<ul><li>DietDiv</li></ul>	1	20858	650.48
- DomSpp	3	23417	654.82

```
- Year 13 38977 671.50
```

Step: AIC=645.67

MassChange ~ DietDiv + Year + DomSpp

Df Deviance AIC <-none> 19508 645.67 + DietEngDen 1 19478 647.55 - DietDiv 1 20891 648.60 - DomSpp 3 23602 653.38 - Year 13 39769 670.95

Call:  $glm(formula = MassChange \sim DietDiv + Year + DomSpp, family = gaussian(link = "identity"),$ 

data = Data\_Q1)

#### Coefficients:

(Intercept)	DietDiv	Year.L
70.599	-63.415	28.367
Year.Q	Year.C	Year^4
12.387	18.307	15.184
Year^5	Year^6	Year^7
-15.549	3.305	23.361
Year^8	Year^9	Year^10
2.392	-23.277	-11.186
Year^11	Year^12	Year^13
13.261	21.328	-39.502
DomSppNorthernSandlance	DomSppPollock	${\tt DomSppRedfish}$
-18.212	-47.479	-20.303

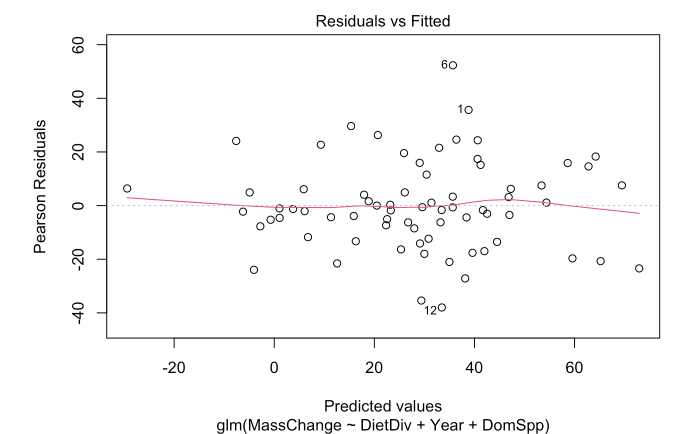
Degrees of Freedom: 71 Total (i.e. Null); 54 Residual

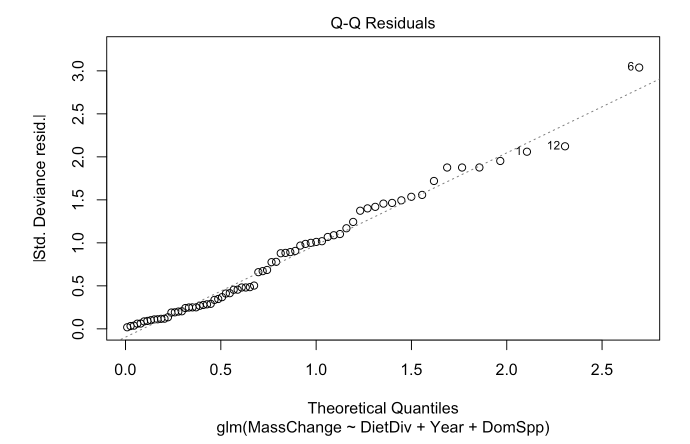
Null Deviance: 48570

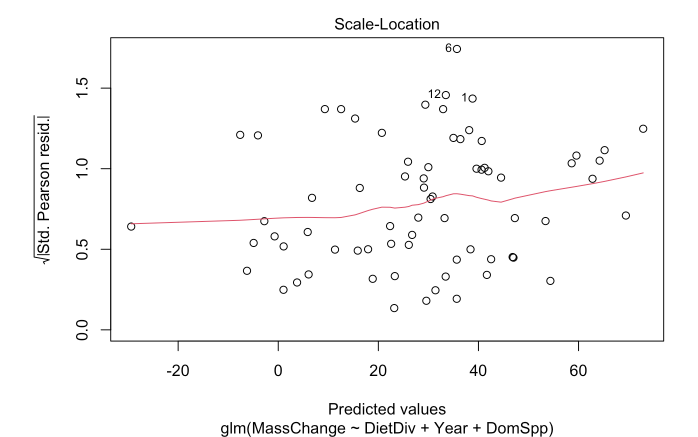
Residual Deviance: 19510 AIC: 645.7

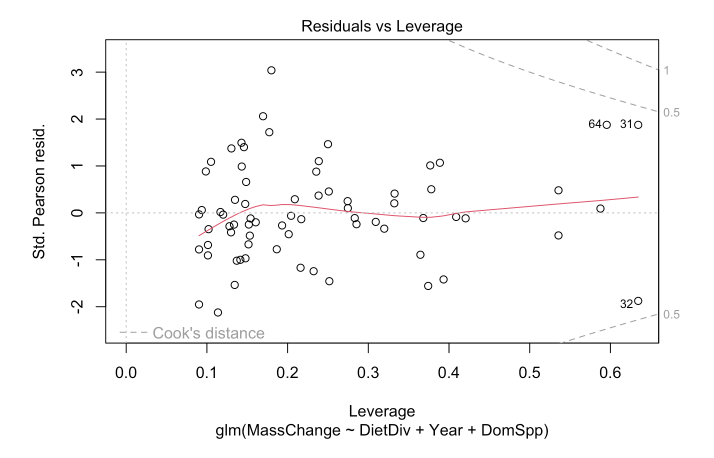
The lowest AIC model drops the dietary energy density covariate. We will update the initial model to drop the energy density covariate:

Warning: not plotting observations with leverage one:









All model residuals still look good. We can proceed to interpreting model output:

```
summary(mom_mod2)
```

#### 

#### Coefficients:

Estimate	Std. Error	t value	Pr(> t )	
70.599	16.241	4.347	6.15e-05	***
-63.415	32.417	-1.956	0.055621	•
28.367	15.657	1.812	0.075584	
12.387	12.620	0.982	0.330717	
18.307	13.274	1.379	0.173517	
15.184	14.787	1.027	0.309065	
-15.549	12.334	-1.261	0.212857	
3.305	10.924	0.303	0.763373	
23.361	10.525	2.219	0.030674	*
2.392	10.067	0.238	0.813123	
-23.277	10.057	-2.314	0.024475	*
-11.186	10.451	-1.070	0.289261	
13.261	9.842	1.347	0.183483	
	70.599 -63.415 28.367 12.387 18.307 15.184 -15.549 3.305 23.361 2.392 -23.277 -11.186	70.599 16.241 -63.415 32.417 28.367 15.657 12.387 12.620 18.307 13.274 15.184 14.787 -15.549 12.334 3.305 10.924 23.361 10.525 2.392 10.067 -23.277 10.057 -11.186 10.451	70.599       16.241       4.347         -63.415       32.417       -1.956         28.367       15.657       1.812         12.387       12.620       0.982         18.307       13.274       1.379         15.184       14.787       1.027         -15.549       12.334       -1.261         3.305       10.924       0.303         23.361       10.525       2.219         2.392       10.067       0.238         -23.277       10.057       -2.314         -11.186       10.451       -1.070	-63.415       32.417       -1.956       0.055621         28.367       15.657       1.812       0.075584         12.387       12.620       0.982       0.330717         18.307       13.274       1.379       0.173517         15.184       14.787       1.027       0.309065         -15.549       12.334       -1.261       0.212857         3.305       10.924       0.303       0.763373         23.361       10.525       2.219       0.030674         2.392       10.067       0.238       0.813123         -23.277       10.057       -2.314       0.024475         -11.186       10.451       -1.070       0.289261

```
Year^12
                        21.328
                                    8.874 2.403 0.019708 *
Year^13
                       -39.502
                                    9.864 -4.005 0.000191 ***
DomSppNorthernSandlance -18.212
                                   14.318 -1.272 0.208824
                                   16.434 -2.889 0.005550 **
DomSppPollock
                       -47.479
DomSppRedfish
                      -20.303
                                   13.938 -1.457 0.150992
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 361.2663)
   Null deviance: 48567 on 71 degrees of freedom
Residual deviance: 19508 on 54 degrees of freedom
AIC: 645.67
```

Number of Fisher Scoring iterations: 2

Cross validation We can perform cross-validation to to test this models predictions. In order to have each fold represent about 10% of the data, we will perform 7 fold cross validation:

```
set.seed(123)
#Number of folds
k <- 7
# Split the data into folds
DatCV <- Data_Q1 %>% mutate(fold = sample(rep(1:k, length.out = n())))
# Create a vector to store RMSE for each fold
rmse values <- numeric(k)</pre>
#Perform cross validation
for (i in 1:k) {
  train_data <- DatCV %>% filter(fold != i)
  test_data <- DatCV %>% filter(fold == i)
  # Fit the model on training data
modCV <- glm(MassChange ~ DietDiv + Year + DomSpp, family = gaussian(link = "identity"),</pre>
  # Predict on the test data
  predictions <- predict(modCV, newdata = test_data)</pre>
  # Compute RMSE for this fold
  actuals <- test_data$MassChange</pre>
  rmse_values[i] <- sqrt(mean((predictions - actuals)^2))</pre>
}
rmse_values
```

```
mean_rmse <- mean(rmse_values)
cat("Average RMSE across", k, "folds:", round(mean_rmse, 2), "\n")</pre>
```

Average RMSE across 7 folds: 15.98

The average root mean squared error is 15.98, indicating the model generally predicts values within 16 kg of the true value.

```
require(ggplot2)
```

Loading required package: ggplot2

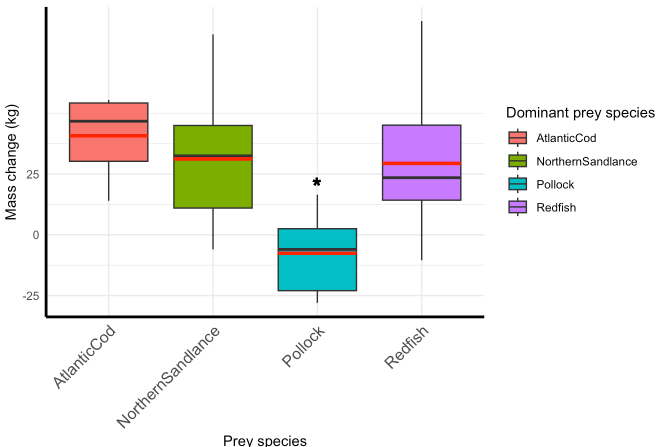
```
ggplot(Data_Q1, aes(x = DomSpp, y = MassChange, fill = DomSpp)) +
 geom boxplot() +
 stat_summary(
   fun = "mean", geom = "errorbar",
   aes(ymax = ..y.., ymin = ..y..),
   width = 0.75, color = "red", size = 1.2
  ) + geom_text(x = "Pollock", y = 20, label = "*",
                aes(x = x, y = y, label = label).
                color = "black", size = 6, fontface = "bold"
 ) +
 labs(
   title = "Figure 1. Median (black line) and mean (red line) mass change by dominant pr
   x = "Prey species",
   y = "Mass change (kg)", fill = "Dominant prey species"
  ) +
 theme minimal() + theme(
    axis.line = element_line(color = "black", size = 1),  # Add axis lines
   axis.text.x = element_text(size = 12, angle = 45, hjust = 1)
  ) +
 scale y continuous(
   breaks = seq(0, max(Data Q1$MassChange), by = 25) - 50
  )
```

```
Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.
i Please use `linewidth` instead.

Warning: The `size` argument of `element_line()` is deprecated as of ggplot2 3.4.0.
i Please use the `linewidth` argument instead.

Warning: The dot-dot notation (`..y..`) was deprecated in ggplot2 3.4.0.
i Please use `after_stat(y)` instead.
```

Figure 1. Median (black line) and mean (red line) mass change by dominant prey



It may also be worth noting that diet diversity is almost significant at p = 0.0556. While this p-value is not significant, the negative relationship between diet diversity and mass change may support the observed effects of dominant prey species, where individuals who primarily prey on cod, sand lance, or redfish and have a lower diet diversity due to this focus on a few profitable species are more successful in gaining mass than individuals who feed on a more diverse array of prey species which include less profitable species.

We can perform cross-validation to to test this models predictions. In order to have each fold represent about 10% of the data, we will perform 7 fold cross validation:

```
#Number of folds
k <- 7

# Split the data into folds
DatCV <- Data_Q1 %>% mutate(fold = sample(rep(1:k, length.out = n())))

# Create a vector to store RMSE for each fold
rmse_values <- numeric(k)

#Perform cross validation
for (i in 1:k) {</pre>
```

```
train_data <- DatCV %>% filter(fold != i)
  test_data <- DatCV %>% filter(fold == i)

# Fit the model on training data
modCV <- glm(MassChange ~ DietDiv + Year + DomSpp, family = gaussian(link = "identity"),

# Predict on the test data
predictions <- predict(modCV, newdata = test_data)

# Compute RMSE for this fold
actuals <- test_data$MassChange
rmse_values[i] <- sqrt(mean((predictions - actuals)^2))
}

# Summarize
rmse_values</pre>
```

[1] 22.44760 11.88442 18.05650 11.05129 18.98355 15.02906 14.37965

```
mean_rmse <- mean(rmse_values)
cat("Average RMSE across", k, "folds:", round(mean_rmse, 2), "\n")</pre>
```

Average RMSE across 7 folds: 15.98

The average root mean squared error is 15.98, indicating the model generally predicts values within 16 kg of the true value. Check the standard deviation to compare the average RMSE to:

```
sd(Data_Q1$MassChange)
```

#### [1] 26.15433

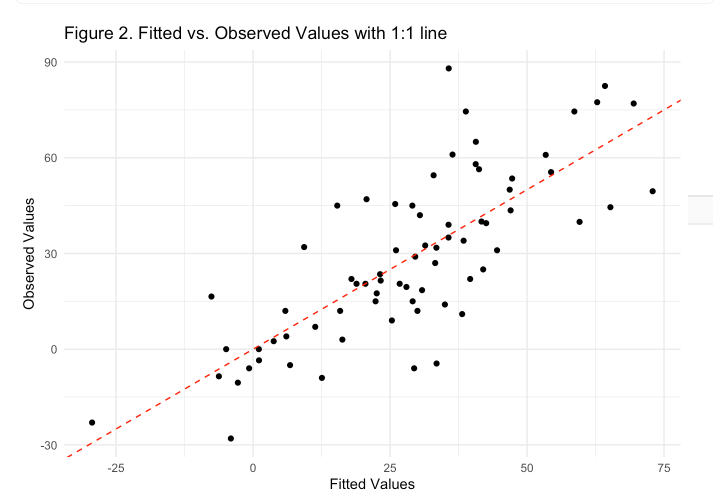
Compared to the standard deviation of mass change, the RMSE value is about 40% lower than the standard deviation, meaning the model predictions are much better than using the population mean to predict mass change. However, compared to the overall mean mass change within our dataset (28.5 kg), there is relatively large prediction error. We can also visualize prediction accuracy with a fitted vs observed value plot (figure 2):

```
fit_mom <- fitted(mom_mod2)

# Get the observed response variable (actual values)
obs_mom <- Data_Q1$MassChange

# Plot the fitted vs. observed values
ggplot(data = Data_Q1, aes(x = fit_mom, y = obs_mom)) +
geom_point(color = "black") +
geom_abline(slope = 1, intercept = 0, color = "red", linetype = "dashed") + # Add 1:1
labs(
    title = "Figure 2. Fitted vs. Observed Values with 1:1 line",</pre>
```

```
x = "Fitted Values",
y = "Observed Values"
) +
theme_minimal()
```



While figure 2 shows that the model predictions have consistent accuracy across the range of fitted values, as suggested by the average RMSE value obtained from cross-validation, there is reasonably large prediction error. Given that the model residuals suggest good model fit, the relatively large prediction error suggest the set of covariates used in this model are not sufficient in predicting maternal mass change. From this we can conclude that females who feed on redfish, sand lance, or cod experience greater mass change than individuals who feed on Pollock, but other factors outside of maternal diet, such as individual physiology or characteristics associated with individual foraging behavior, are responsible for explaining a large proportion of variation in maternal mass change.

## **Research Question 2**

Does variation in diet of maternal female predict pup weaning mass in Northwest Atlantic grey seals?

#### **Load Data**

require(tinytex) quarto install tinytex

Cleaned data file was uploaded. seal\_data = read.csv("Data\_Q2.csv", header = T)

```
seal_data <- read.csv("/Users/peterbraithwaite/Desktop/IDPhD Classes/Stat 5620_Updated/Fi
summary(seal_data)</pre>
```

```
Year
    MomID
                               Dietary.energy.density Diet.diversity
Min.
           19
                Min.
                       :1996
                               Min.
                                      :5.207
                                                      Min.
                                                              :0.1788
1st Qu.: 467
                1st Qu.:2002
                               1st Qu.:5.676
                                                      1st Qu.:0.3108
Median: 4688
                Median :2011
                               Median :5.784
                                                      Median :0.3564
     : 4999
Mean
                Mean
                       :2008
                               Mean
                                      :5.772
                                                      Mean
                                                             :0.3622
3rd Qu.: 9412
                3rd Qu.:2013
                               3rd Qu.:5.900
                                                      3rd Qu.:0.4181
Max.
       :10690
                Max.
                       :2015
                               Max.
                                      :6.517
                                                      Max.
                                                             :0.5290
                                                      Pup.Wean.Mass
Dominant.prey.species
                         Mom.Age
                                         Pup.sex
Length:56
                      Min.
                             : 9.00
                                      Min.
                                             :1.000
                                                      Min.
                                                             :34.50
                                                      1st 0u.:45.50
Class :character
                      1st Qu.:23.00
                                      1st Ou.:1.000
Mode :character
                      Median :25.00
                                      Median :1.000
                                                      Median :50.75
                      Mean
                             :23.46
                                      Mean
                                            :1.464
                                                      Mean
                                                             :50.31
                      3rd 0u.:26.25
                                      3rd Qu.:2.000
                                                      3rd Qu.:55.12
                             :31.00
                                      Max.
                                            :2.000
                      Max.
                                                      Max.
                                                             :63.00
```

# **Data Exploration**

The package "flexplot" was used to explore the predictor and response variables.

```
require(flexplot)

Loading required package: flexplot

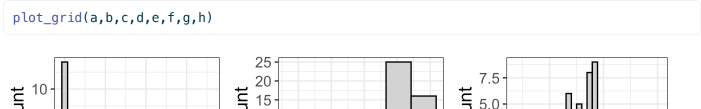
Attaching package: 'flexplot'

The following object is masked from 'package:ggplot2':
    flip_data
```

```
a = flexplot (MomID~1, data= seal_data)
b = flexplot (Year~1, data= seal_data)
c = flexplot (Dietary.energy.density~1, data= seal_data)
d = flexplot (Diet.diversity~1, data= seal_data)
e = flexplot (Dominant.prey.species~1, data= seal_data)
f = flexplot (Mom.Age~1, data= seal_data)
g = flexplot (Pup.sex~1, data= seal_data)
```

```
h = flexplot (Pup.Wean.Mass~1, data= seal_data)
require (cowplot)
```

Loading required package: cowplot



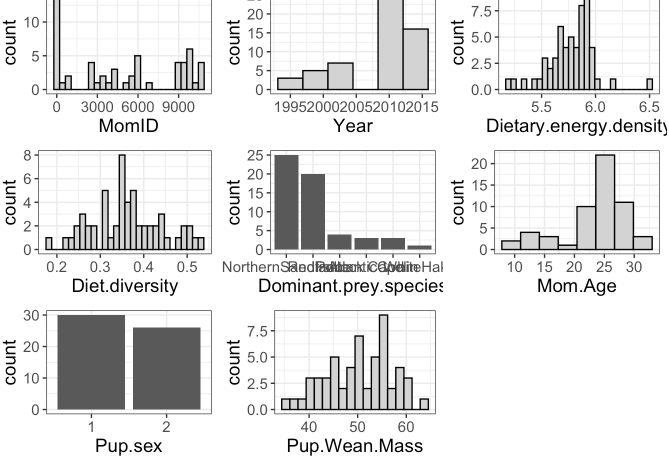


Figure 1 - Exploring Variable Distributions

From the plots above it seems that the response variable (Pup Wean Mass) has a Gaussian distribution.

To further explore the relationship between variable the package "ggplot2" was used plot individual variables in relation to the repsonse variables.

# **Pup Wean Mass and Dietary Engery Density**

```
require(ggplot2)
# Pup Wean Mass + Dietary.energy.density
```

 $geom_smooth()$  using formula = 'y  $\sim$  x'

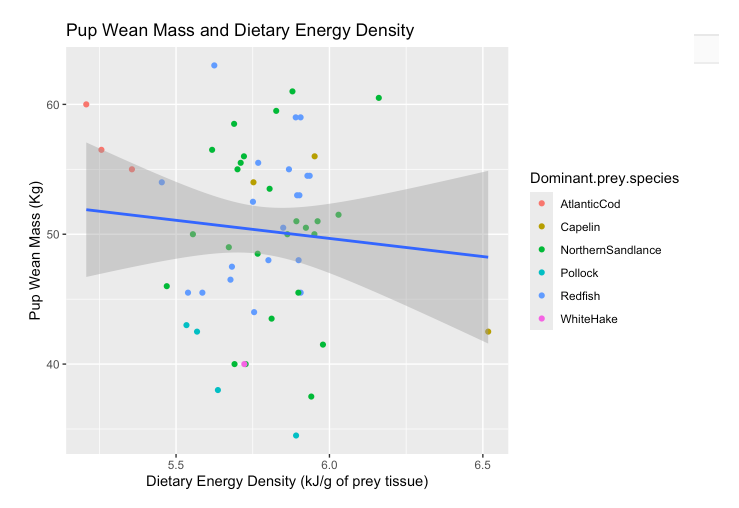


Figure 2 - Pup Wean Mass and Dietary Energy Density

From this plot we can see a negative or inverse relationship between Pup Wean Mass and Dietary Energy Density. This indicates that maternal mothers that feed on prey species resulting in greater dietary energy density tend to have pups with a lower wean mass. This is somewhat at odds with what might have been expected. We might have expected the greater the dietary energy density of the maternal mother would result in heavier and ultimately more fit pups.

We can also see that the Dominant Species Capelin has only three observation and one of these appears to be an outlier. It should also be noted that White Hake has only one observation and Atlantic Cod has only three. This lack of observations may be problematic while fitting a model and we may need to remove these Dominant Prey Species.

# **Pup Wean Mass and Dietary Engery Density**

```
# Pup Wean Mass + Diet Diveristy
ggplot(seal_data) + geom_point(aes(Diet.diversity, Pup.Wean.Mass, color = Dominant.prey.s
    labs(title = "Pup Wean Mass and Diet Diversity", x = "Diet Diveristy (no specific units
    geom_smooth(aes(Diet.diversity, Pup.Wean.Mass), method="lm", se=T)
```

#### Pup Wean Mass and Diet Diversity

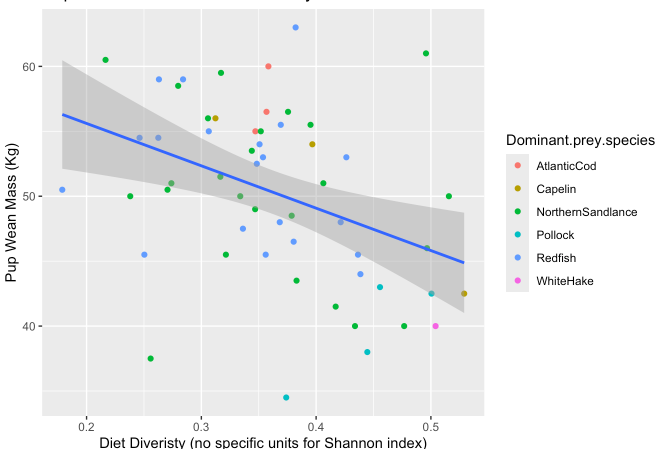


Figure 3 - Pup Wean Mass and Diet Diversity

From this plot we can see a negative or inverse relationship between Pup Wean Mass and Diet Diversity. This indicates that as the maternal mothers feed on more diverse prey species their pup wean mass decreases. This may indicates that mothers that reduce the diversity of their diet have healthier and more fit pups.

### Pup Wean Mass + Year

Exploring if specific years showed abnormally high or low Pup Wean Mass and if these were related to specific dominant prey species.

 $geom_smooth()$  using formula = 'y ~ x'

```
ggplot(seal_data) + geom_point(aes(Year, Pup.Wean.Mass, color = Dominant.prey.species)) +
labs(title = "Pup Wean Mass and Year", x = "Year", y = "Pup Wean Mass (Kg)") +
geom_smooth (aes(Year, Pup.Wean.Mass), method="lm", se=T)
```

`geom\_smooth()` using formula = 'y  $\sim$  x'

#### Pup Wean Mass and Year

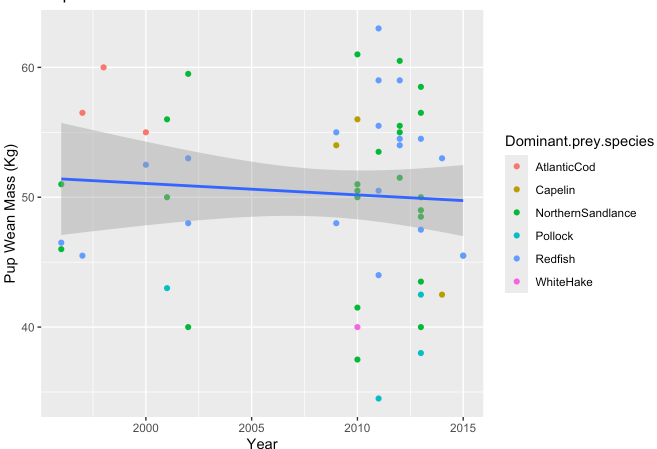


Figure 4 - Pup Wean Mass and Year

From this plot it seems Pup Wean Mass seems to be fairly evenly distributed over the observed years and it does not seem that a specific Dominant Prey Species had a disproportionate effect on the Pup Wean Mass.

### Pup Wean Mass + MomID

Exploring if Pup Wean Mass shows significant differences based on the maternal mother ID.

```
ggplot(seal_data) + geom_point(aes(MomID, Pup.Wean.Mass, color = Year )) +
  labs(title = "Pup Wean Mass and Mom ID", x = "Mom ID", y = "Pup Wean Mass (Kg)") +
  geom_smooth(aes(MomID, Pup.Wean.Mass), method="lm", se=T)
```

<sup>`</sup>geom\_smooth()` using formula = 'y  $\sim$  x'

#### Pup Wean Mass and Mom ID

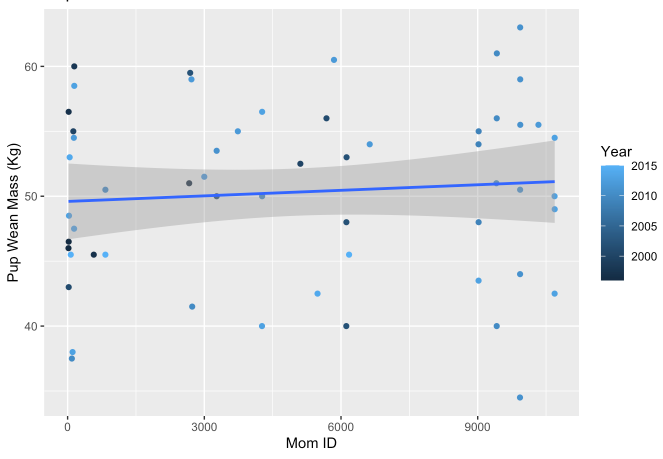


Figure 5 - Pup Wean Mass and Materianl Mother ID

With the vertical rows of the this plot representing unique maternal mothers, it does not seem any individual is creating disproportionately large or small pup weights. Rather, it seems most maternal mother are creating a fairly evenly distribution of pup weights dependent on the year.

# **Pup Wean Mass + Dominant Prey Species**

```
ggplot(seal_data) + geom_point(aes(Dominant.prey.species, Pup.Wean.Mass, color = Year ))
```

#### Pup Wean Mass and Dominant Prey Species

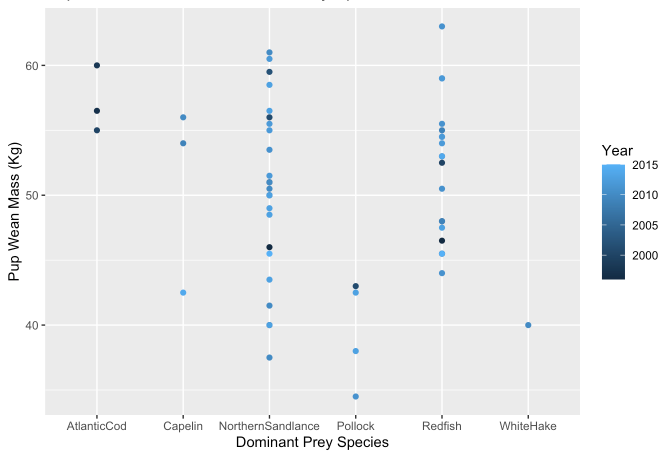


Figure 6 - Pup Wean Mass and Dominant Prey Species

A considerable amount of insight can be gained from this plot including: The Dominant Prey Species with the largest number of observations (Northern Sandiance and Redfish) show a variety of Pup Wean Masses and these occur over a wide variety of years; the Dominant Prey Species with the smallest number of observations tend to show either predominantly large (Atlantic Cod and Capelin) or very small (Pollock and White Hake) Pup Wean sizes and these tend to occur in a variety of years.

# Pup Wean Mass + Pup Sex

```
ggplot(seal_data) + geom_point(aes(Pup.sex, Pup.Wean.Mass, color = Year )) +
labs(title = "Pup Wean Mass and Pup Sex", x = "Pup Sex", y = "Pup Wean Mass (Kg)") +
geom_smooth(aes(Pup.sex, Pup.Wean.Mass), method="lm", se=T)
```

 $<sup>\</sup>ensuremath{\text{`geom\_smooth()`}}\ using formula = 'y \sim x'$ 

#### Pup Wean Mass and Pup Sex

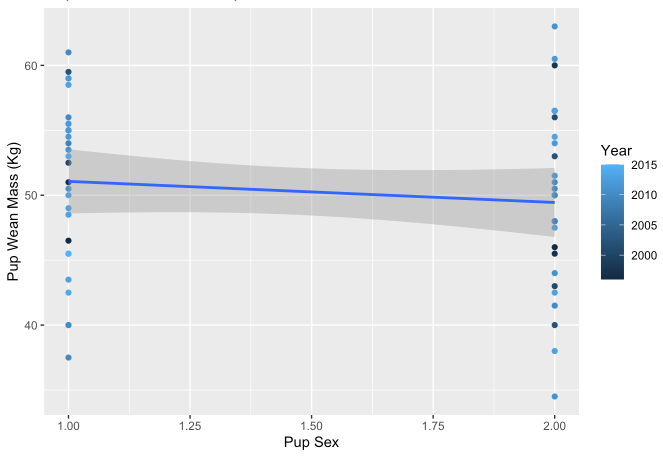


Figure 7 - Pup Wean Mass and Pup Sex

This plot shows that the Pup Sex seems to have little effect on the average Pup Mean Mass. Both Pup Sex seem to create a fairly even distribution of Pup Mean Mass over a variety of years. It is expected that this variable will have a small effect on the fit model.

# Pup Wean Mass + Maternal Age

```
ggplot(seal_data) + geom_point(aes(Mom.Age, Pup.Wean.Mass, colour = Dominant.prey.species
labs(title = "Pup Wean Mass and Maternal Age", x = "Maternal Age", y = "Pup Wean Mass (
```

<sup>`</sup>geom\_smooth()` using formula = 'y  $\sim$  x'

#### Pup Wean Mass and Maternal Age

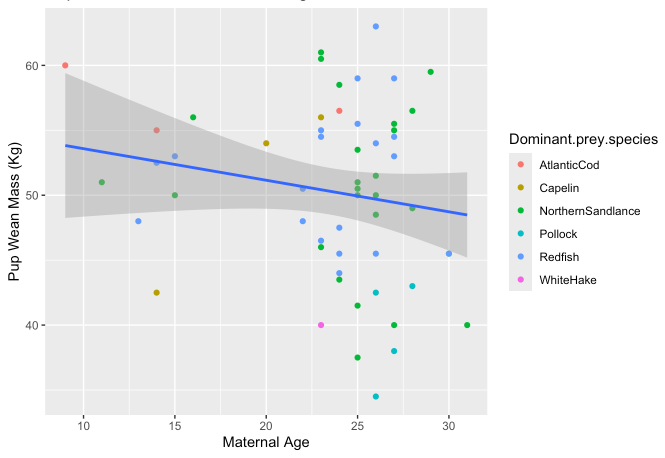


Figure 7 - Pup Wean Mass and Maternal Age

A considerable amount of insight can be gained from this plot. It seems younger maternal seals have larger pups but it is also noteworthy that the majority of older mothers prey on the most observed Dominant Prey Species including Redfish and Northern Sandlance. Alternatively it seems less frequent observed Dominant Prey Species were preyed on by young maternal mothers. It seems as though there might be a social aspects to why younger mothers prey on less frequently observed Dominant Prey Species.

# **Fit Linear Model**

We now explore fitting general linear models to further explore the variables, the residuals and their relationships.

First we convert the catagorical data to factors and determine the number of levels within each catagorical group.

```
# convert catagorical data to factors

seal_data$MomID = as.factor(seal_data$MomID)
seal_data$Year = as.factor(seal_data$Year)
```

```
seal_data$Dominant.prey.species = as.factor(seal_data$Dominant.prey.species)
seal_data$Pup.sex = as.factor(seal_data$Pup.sex)
seal_data$Pup.Mom.Age = as.factor(seal_data$Mom.Age)
nlevels(seal_data$MomID)
```

[1] 51

```
# 51
nlevels(seal_data$Year)
```

[1] 13

```
# 13
nlevels(seal_data$Dominant.prey.species)
```

[1] 6

```
# 5
nlevels(seal_data$Pup.sex)
```

[1] 2

```
# 2
nlevels(seal_data$Pup.Mom.Age)
```

[1] 17

# 17

# **Fit Linear Model**

#### **All Predictor Variables**

```
Q1 = lm(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density + Year + Mom.Age + Pup.sex + MomID, data = seal_data) summary(Q1)
```

```
Call:
```

```
lm(formula = Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity +
    Dietary.energy.density + Year + Mom.Age + Pup.sex + MomID,
    data = seal_data)
```

Residuals:

Coefficients: (16 not defined because of singularities)

coefficients: (10 not defined because (	_				
4-	Estimate	Std.			
(Intercept)	-127.5808		NaN	NaN	NaN
Dominant.prey.speciesCapelin	14.0800		NaN	NaN	NaN
Dominant.prey.speciesNorthernSandlance			NaN	NaN	NaN
Dominant.prey.speciesPollock	-19.7677		NaN	NaN	NaN
Dominant.prey.speciesRedfish	-16.6515		NaN	NaN	NaN
Dominant.prey.speciesWhiteHake	4.6941		NaN	NaN	NaN
Diet.diversity	8.3424		NaN	NaN	NaN
Dietary.energy.density	35.8485		NaN	NaN	NaN
Year1997	19.2395		NaN	NaN	NaN
Year1998	-1.3181		NaN	NaN	NaN
Year2000	14.1916		NaN	NaN	NaN
Year2001	-0.8124		NaN	NaN	NaN
Year2002	-3.0383		NaN	NaN	NaN
Year2009	29.8803		NaN	NaN	NaN
Year2010	-2.9017		NaN	NaN	NaN
Year2011	37.2376		NaN	NaN	NaN
Year2012	23.6996		NaN	NaN	NaN
Year2013	20.8844		NaN	NaN	NaN
Year2014	-55.5366		NaN	NaN	NaN
Year2015	16.4409		NaN	NaN	NaN
Mom.Age	-1.8940		NaN	NaN	NaN
Pup.sex2	14.4153		NaN	NaN	NaN
MomID23	4.4626		NaN	NaN	NaN
MomID24	27.5950		NaN	NaN	NaN
MomID30	-7 <b>.</b> 9254		NaN	NaN	NaN
MomID45	88.7703		NaN	NaN	NaN
MomID69	26.2285		NaN	NaN	NaN
MomID93	-2.2734		NaN	NaN	NaN
MomID109	-4.5789		NaN	NaN	NaN
MomID125	NA 2. 62.42		NA	NA	NA
MomID137	3.6242		NaN	NaN	NaN
MomID142	-4.6121		NaN	NaN	NaN
MomID146	1.8643		NaN	NaN	NaN
MomID574	NA 2. 1020		NA	NA	NA
MomID829	-3.1820		NaN	NaN	NaN
MomID2668	-19.0778		NaN	NaN	NaN
MomID2690	31.0252		NaN	NaN	NaN
MomID2718	16.5836		NaN	NaN	NaN
MomID2734	-15.3957		NaN	NaN	NaN
MomID2999	-31.0853		NaN	NaN	NaN
MomID3271	-22.2778		NaN	NaN	NaN
MomID3736	0.2347		NaN	NaN	NaN
MomID4266	-12.6464		NaN	NaN	NaN
MomID4269	-5.2029		NaN	NaN	NaN
MomID5108	NA NA		NA NA	NA NA	NA NA
MomID5485	NA 0. 0710		NA	NA NaN	NA
MomID5681	-9.8719		NaN	NaN	NaN

MomTDE946	21 6400	MaN	NaN	MaM
MomID5846	-31.6400	NaN	NaN	NaN
MomID6116	3.4943	NaN	NaN	NaN
MomID6118	-9.0816	NaN	NaN	NaN
MomID6122	NA	NA	NA	NA
MomID6177	8.7215	NaN	NaN	NaN
MomID6630	10.9350	NaN	NaN	NaN
MomID9018	-34.0206	NaN	NaN	NaN
MomID9019	-18.3770	NaN	NaN	NaN
MomID9020	-21.8694	NaN	NaN	NaN
MomID9021	NA	NA	NA	NA
MomID9410	-2.7034	NaN	NaN	NaN
MomID9417	NA	NA	NA	NA
MomID9418	NA	NA	NA	NA
MomID9420	17.6372	NaN	NaN	NaN
MomID9928	-34.8701	NaN	NaN	NaN
MomID9930	-27.9118	NaN	NaN	NaN
MomID9931	-12.0164	NaN	NaN	NaN
MomID9932	NA	NA	NA	NA
MomID9933	NA	NA	NA	NA
MomID9934	NA	NA	NA	NA
MomID10333	NA	NA	NA	NA
MomID10687	NA	NA	NA	NA
MomID10688	NA	NA	NA	NA
MomID10689	NA	NA	NA	NA
MomID10690	NA	NA	NA	NA

Residual standard error: NaN on 0 degrees of freedom Multiple R-squared: 1, Adjusted R-squared: NaN

F-statistic: NaN on 55 and 0 DF, p-value: NA

When this model is run we received an error suggesting we have more columns than rows. This suggest we have more variables than observations. As a result we must remove some of the categorical variables. The maternal mother ID seems to be adding a lot of categories (n -51) and did not seem very important in Figure 6 above. As a result Mom ID was removed and the model was fit again.

## All Predictor Variables Except Mom ID

```
Q2 = lm(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density + summary
```

```
standardGeneric for "summary" defined from package "base"
function (object, ...)
standardGeneric("summary")
<environment: 0x10d3d74b0>
Methods may be defined for arguments: object
Use showMethods(summary) for currently available ones.
```

```
# Adjusted R-squared: 0.209
# p-value: 0.08393
```

Although this model was not significant, it did provide a model that works as a starting point for the Step Function.

#### **Step Function (Forward)**

We now used the Step Function to determine the best fit model using AIC.

```
fwd.model = step (Q2, direction='forward')
```

Start: AIC=216.76

Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density + Year + Mom.Age + Pup.sex

The forward step retained all variables with AIC = 216.76

#### **Step Function (Backward)**

```
backward.model = step(Q2, direction='backward')
```

۸ T C

```
Start: AIC=216.76
Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
    Year + Mom.Age + Pup.sex
```

	υτ	Sum of Sq	RSS	AIC
– Year	12	347.70	1572.2	206.75
- Mom.Age	1	1.61	1226.1	214.83
- Pup.sex	1	17.14	1241.6	215.53
<none></none>			1224.5	216.76
<ul><li>Dietary.energy.density</li></ul>	1	47.89	1272.4	216.91
<ul><li>Diet.diversity</li></ul>	1	69.12	1293.6	217.83
<ul><li>Dominant.prey.species</li></ul>	5	329.04	1553.5	220.08

D4 C.... a4 Ca

Step: AIC=206.75

```
Df Sum of Sq
                                      RSS
                                            AIC
- Mom.Age
                        1
                              1.51 1573.7 204.81
                        1
                             3.13 1575.3 204.87
- Pup.sex
Dietary.energy.density 1
                            50.68 1622.9 206.53
<none>
                                   1572.2 206.75
- Dominant.prey.species 5 364.88 1937.1 208.44
                        1
                            162.65 1734.8 210.27
Diet.diversity
```

Step: AIC=204.81

Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density +

```
Df Sum of Sq
                                        RSS
                                               ATC
- Pup.sex
                         1
                                2.57 1576.3 202.90
- Dietary.energy.density 1
                               49.35 1623.0 204.54
                                     1573.7 204.81
<none>
                         5
                              415.52 1989.2 207.93
Dominant.prey.species
Diet.diversity
                              164.37 1738.1 208.37
                         1
Step: AIC=202.9
Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density
                        Df Sum of Sq
                                        RSS
                                               AIC
Dietary.energy.density 1
                               49.16 1625.4 202.62
<none>
                                     1576.3 202.90
Diet.diversity
                         1 166.48 1742.8 206.52
Dominant.prey.species
                         5
                              456.31 2032.6 207.14
Step: AIC=202.62
Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity
                       Df Sum of Sq
                                       RSS
                                              ATC
<none>
                                    1625.4 202.62
Diet.diversity
                        1
                             124.06 1749.5 204.74
- Dominant.prey.species 5
                             493.24 2118.7 207.46
The backward step retained only the Dominant Prey Species and Diet Diversity variables with AIC
202.62.
Explore The Resulting Linear Model From Backward Step Function
 Q1_Reduced = lm(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity,
                data = seal data)
 summary(Q1_Reduced)
Call:
lm(formula = Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity,
    data = seal_data)
Residuals:
     Min
               10
                   Median
                                30
                                        Max
                   0.0968 3.2694 13.3653
-15.0508 -4.1950
Coefficients:
                                      Estimate Std. Error t value Pr(>|t|)
                                        64.422
                                                    5.013 12.850 < 2e-16 ***
(Intercept)
Dominant.prey.speciesCapelin
                                        -5.133
                                                    4.743 -1.082 0.284509
Dominant.prey.speciesNorthernSandlance
                                        -6.630
                                                    3.519 -1.884 0.065539 .
```

-15.831

Dominant.prey.speciesPollock

4.500 -3.518 0.000949 \*\*\*

```
Dominant.prey.speciesRedfish -5.797 3.570 -1.624 0.110838
Dominant.prey.speciesWhiteHake -14.093 6.838 -2.061 0.044627 *
Diet.diversity -20.491 10.596 -1.934 0.058916 .
```

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

Residual standard error: 5.76 on 49 degrees of freedom Multiple R-squared: 0.3509, Adjusted R-squared: 0.2715 F-statistic: 4.416 on 6 and 49 DF, p-value: 0.00121

```
# Adjusted R-squared: 0.2715
# p-value: 0.00121
```

This found that the reduced Linear Model is significant with a p-value = 0.00121 and an Adjusted R-Squared = 0.2715 meaning the model accounts for about 27% of the variation of the Pup Wean Mass. It should also be noted that the most significant variables are Pollock with a p-value of 0.000949 and White Hake with a p-value = 0.044627.

We will now look at the residuals for the this model.

# **Explore Residuals for Reduced Linear Model**

```
library(ggfortify)
autoplot(Q1_Reduced)
```

Warning: Removed 5 rows containing missing values or values outside the scale range (`geom line()`).

Warning: Removed 1 row containing missing values or values outside the scale range (`geom\_point()`).

Warning: Removed 1 row containing missing values or values outside the scale range (`geom\_line()`).

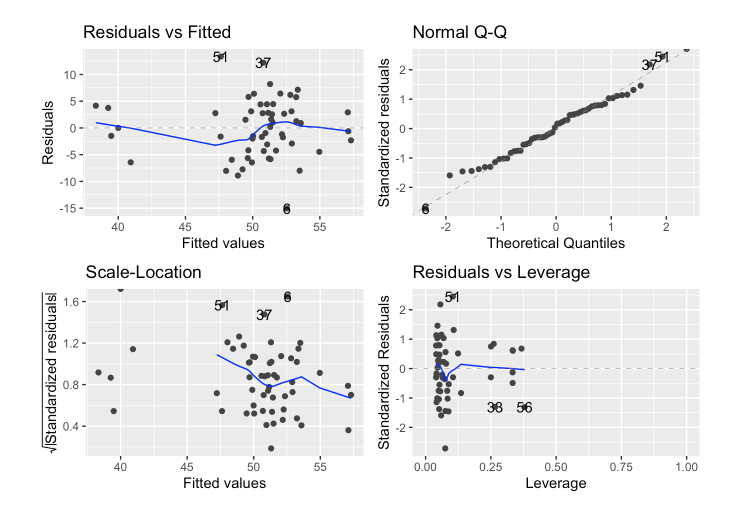


Figure 8 - Residual for Reduced Variables Linear Model

```
#residuals
res = resid(Q1_Reduced)
plot(fitted(Q1_Reduced), res)
abline(0,0)
```

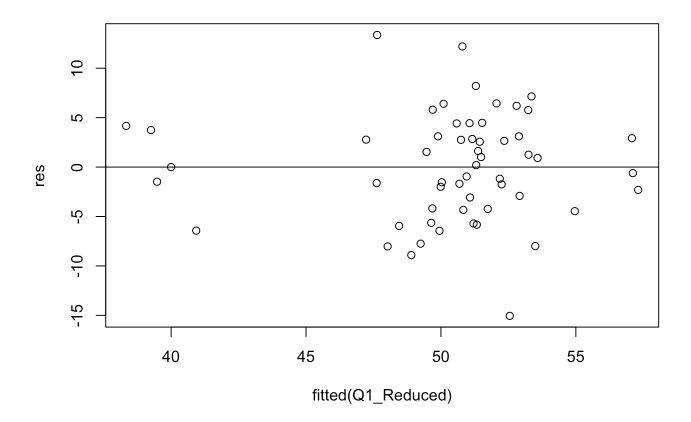


Figure 9 - Residual for Reduced Variables Linear Model

```
ggplot(Q1_Reduced, aes(x = .fitted, y = .resid, colour = Dominant.prey.species )) + geom_
```

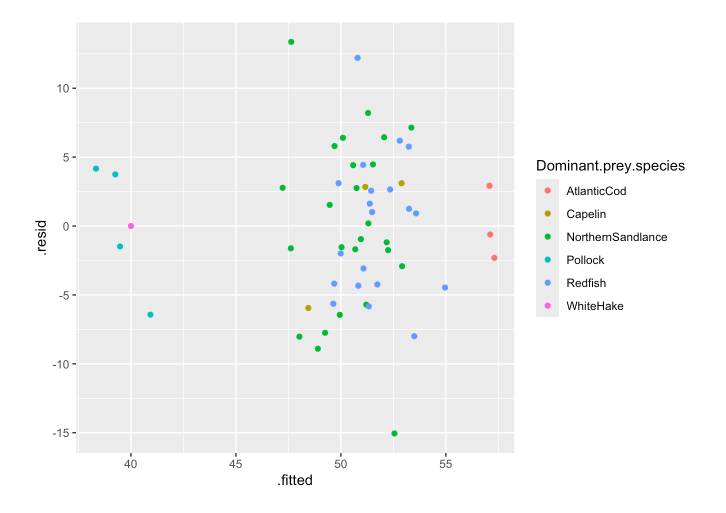


Figure 10 - Residual for Reduced Variables Linear Model

From these residual visualization it seems clear that a number observations seem to be outliers and they should be removed or we should try robust regression strategies. We can see from Figure 10 that it is the observation with White Hake and Pollock as the Dominant Prey Species are clearly outliers.

We will now try to fit a robust regression first.

#### Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                                         66.326
                                                     4.925 13.467 < 2e-16 ***
Dominant.prey.speciesCapelin
                                         -4.698
                                                     2.646 -1.776 0.081995 .
                                                     1.825 -3.540 0.000889 ***
Dominant.prey.speciesNorthernSandlance
                                         -6.460
                                                     3.194 -4.761 1.74e-05 ***
                                        -15.208
Dominant.prey.speciesPollock
Dominant.prey.speciesRedfish
                                         -6.031
                                                     1.719 -3.508 0.000977 ***
                                                     2.405 -5.518 1.28e-06 ***
Dominant.prey.speciesWhiteHake
                                        -13.271
                                                    13.446 -1.926 0.059903 .
                                        -25.898
Diet.diversity
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Robust residual standard error: 5.928
Multiple R-squared: 0.3699,
                                Adjusted R-squared: 0.2928
Convergence in 12 IRWLS iterations
Robustness weights:
 4 weights are \sim= 1. The remaining 52 ones are summarized as
   Min. 1st Qu. Median
                           Mean 3rd Qu.
                                           Max.
 0.4607 0.9074 0.9590 0.9238 0.9847 0.9974
Algorithmic parameters:
```

tuning.chi		bb	tu	ning.psi		refine.tol
1.548e+00	5.00	00e-01	4	∙685e+00		1.000e-07
rel.tol	sca	le.tol	S	olve.tol		zero.tol
1.000e-07	1.00	00e-10	1	.000e−07		1.000e-10
eps.outlier		eps.x	warn.limi	t.reject v	warn.l	.imit.meanrw
1.786e-03	1.81	19e-12	5	.000e-01		5.000e-01
nResample	max.it		best.r.s	k.fa	ast.s	k.max
500	50		2		1	200
maxit.scale	trace.lev		mts	compu	te.rd	fast.s.large.n
200	0		1000		0	2000
	psi	subs	sampling			cov
"bisqua	re"	"nons	ingular"	".\	vcov.a	avar1"
compute.outlier.st	ats					
11	SM''					

seed : int(0)

The robust regression returned a better Adjusted R-Squared = 2928 (appoximately 29% of variation) than the regular regression with that showed approximately 27%. Interestingly all the Dominant Prey Species are now significant coefficients except Capelin. We will now look at the residuals for the robust regression model.

```
#residuals
res = resid(Q1_Reduced_Robust)
plot(fitted(Q1_Reduced_Robust), res)
abline(0,0)
```

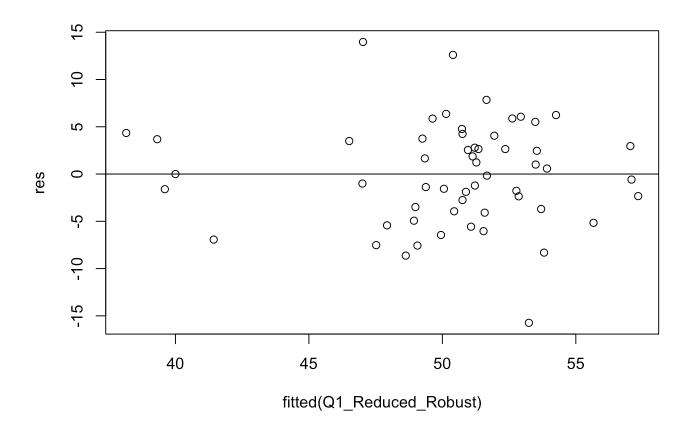


Figure 11 - Residual for Reduced Variables with Robust Linear Model

The residuals look a bit better but not a huge improvement over the linear model above. We will try removing the outlier observation. A new dataset was uploaded below with Pollock and White Hake Dominant Prey Species Removed.

```
seal_data_2 <- read.csv("/Users/peterbraithwaite/Desktop/IDPhD Classes/Stat 5620_Updated/
summary(seal_data_2)</pre>
```

```
MomID
                        Year
                                   Dietary.energy.density Diet.diversity
Min.
       :
           19.0
                   Min.
                          :1996
                                   Min.
                                          :5.207
                                                           Min.
                                                                   :0.1788
                   1st Qu.:2002
                                   1st Qu.:5.686
1st Qu.: 701.5
                                                           1st Qu.:0.3061
Median: 4269.0
                   Median :2011
                                   Median :5.805
                                                           Median :0.3518
Mean
       : 4897.8
                          :2008
                                          :5.782
                                                                   :0.3531
                   Mean
                                   Mean
                                                           Mean
3rd Ou.: 9020.5
                   3rd Qu.:2013
                                   3rd Qu.:5.904
                                                           3rd 0u.:0.3959
                                                                   :0.5290
Max.
       :10690.0
                   Max.
                          :2015
                                          :6.517
                                                           Max.
                                   Max.
Dominant.prey.species
                          Mom.Age
                                           Pup.sex
                                                         Pup.Wean.Mass
Length:51
                       Min.
                               : 9.00
                                               :1.000
                                                         Min.
                                                                :37.50
                                        Min.
                       1st Qu.:23.00
Class :character
                                        1st Qu.:1.000
                                                         1st Qu.:47.00
Mode :character
                       Median :25.00
                                        Median :1.000
                                                         Median :51.50
                       Mean
                              :23.22
                                               :1.431
                                                         Mean
                                                                :51.36
                                        Mean
```

3rd Qu.:26.00 3rd Qu.:2.000 3rd Qu.:55.50 Max. :31.00 Max. :2.000 Max. :63.00

#### Re-fit Linear Model with Reduced Data-set

Removed Pollock and White Hake from Data-set

```
Q3_Full= lm(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.densi summary(Q3_Full)
```

```
Call:
```

```
lm(formula = Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity +
    Dietary.energy.density + Year + Mom.Age + Pup.sex, data = seal_data_2)
```

#### Residuals:

```
Min 10 Median 30 Max
-14.9432 -3.2854 0.9013 3.1288 14.2253
```

#### Coefficients:

	Estimate	Std. Error	t value	Pr(> t )	
(Intercept)	-258.8154	396.4474	-0.653	0.5174	
Dominant.prey.speciesCapelin	-0.8433	6.9966	-0.121	0.9046	
Dominant.prey.speciesNorthernSandlance	-3.2757	5.2773	-0.621	0.5381	
Dominant.prey.speciesRedfish	-2.9541	5.0552	-0.584	0.5621	
Diet.diversity	-26.8081	11.8105	-2.270	0.0284	*
Dietary.energy.density	-7.2831	6.0669	-1.200	0.2367	
Year	0.1835	0.2045	0.897	0.3747	
Mom.Age	-0.1580	0.2413	-0.655	0.5160	
Pup.sex	-0.1694	1.7915	-0.095	0.9251	

\_\_\_

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

```
Residual standard error: 5.95 on 42 degrees of freedom
```

Multiple R-squared: 0.1856, Adjusted R-squared: 0.03048

F-statistic: 1.197 on 8 and 42 DF, p-value: 0.3243

```
# p-value: 0.3243
# Adjusted R-squared: 0.03048
```

## **Step Function (Forward)**

We now used the Step Function to determine the best fit model using AIC.

```
fwd.model = step (Q3_Full, direction='forward')
```

Start: AIC=190

Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
 Year + Mom.Age + Pup.sex

The forward step retained all variables with AIC = 190. It should be noted that this AIC is better than the best AIC 202.62 for the simple linear regression above.

### **Step Function (Backward)**

```
backward.model = step(Q3_Full, direction='backward')
Start: AIC=190
Pup.Wean.Mass ∼ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
   Year + Mom.Age + Pup.sex
                        Df Sum of Sq
                                        RSS
                                               ATC
Dominant.prey.species
                         3
                              27.752 1514.5 184.94
- Pup.sex
                         1
                               0.316 1487.0 188.01
                         1 15.188 1501.9 188.52
- Mom.Age
                         1 28.503 1515.2 188.97
- Year
- Dietary.energy.density 1 51.011 1537.7 189.72
                                     1486.7 190.00
<none>
                         1 182,378 1669,1 193,90
Diet.diversity
Step: AIC=184.94
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age + Pup.sex
                        Df Sum of Sq
                                        RSS
                                               ATC
- Pup.sex
                         1
                               1.573 1516.0 182.99
- Year
                         1
                              36.520 1551.0 184.16
                         1
- Mom.Age
                              52.015 1566.5 184.66
                                     1514.5 184.94
<none>
Dietary.energy.density 1 134.905 1649.4 187.29
Diet.diversity
                         1 195.457 1709.9 189.13
Step: AIC=182.99
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age
                        Df Sum of Sq
                                        RSS
                                               AIC
                              41.970 1558.0 182.39
- Year
                         1
- Mom.Age
                         1
                              52,962 1569,0 182,75
<none>
                                     1516.0 182.99
Dietary.energy.density 1 134.114 1650.2 185.32
                         1 195.334 1711.4 187.17
- Diet.diversity
Step: AIC=182.39
```

Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Mom.Age

```
Df Sum of Sq
                                        RSS
                                               AIC
- Mom.Age
                         1
                              17.150 1575.2 180.94
<none>
                                      1558.0 182.39
- Dietary.energy.density 1 93.118 1651.1 183.35
- Diet.diversity
                         1 187,904 1745,9 186,19
Step: AIC=180.94
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density
                        Df Sum of Sq
                                        RSS
                                               AIC
                                      1575.2 180.94
<none>
                              93.233 1668.4 181.88
Dietary.energy.density 1
- Diet.diversity
                          1
                             197.610 1772.8 184.97
```

The backward step retained the Diet Diversity and Dietary.energy.density variables with AIC 180.94. It should be noted that this AIC is better than the best AIC 202.62 for the simple linear regression above.

### **Explore Backward Step Linear Model**

```
Call:
```

```
lm(formula = Pup.Wean.Mass ~ Dietary.energy.density + Diet.diversity,
    data = seal_data_2)
```

#### Residuals:

```
Min 1Q Median 3Q Max -15.3764 -4.3276 0.7659 3.9409 13.9590
```

#### Coefficients:

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

(Intercept) 97.382 22.844 4.263 9.39e-05 ***

Dietary.energy.density -6.374 3.781 -1.686 0.0984 .

Diet.diversity -25.954 10.576 -2.454 0.0178 *

---

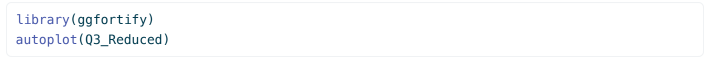
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 5.728 on 48 degrees of freedom Multiple R-squared: 0.1372, Adjusted R-squared: 0.1012

F-statistic: 3.815 on 2 and 48 DF, p-value: 0.029

```
# p-value: 0.029
# Adjusted R-squared: 0.1012
```

# **Explore Residuals For Reduced DataSet Romoving White Hake and Pollock**



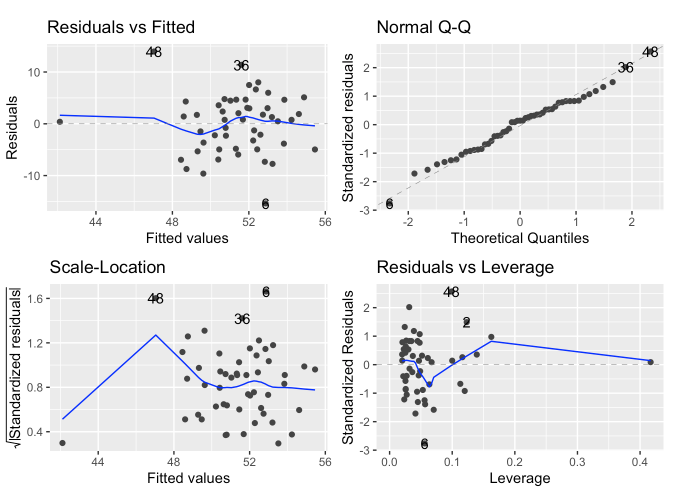


Figure 12 - Residual for Reduced Dataset (White Hake and Pollock removed).

```
ggplot(Q3_Reduced, aes(x = .fitted, y = .resid, color = seal_data_2$Dominant.prey.species
```

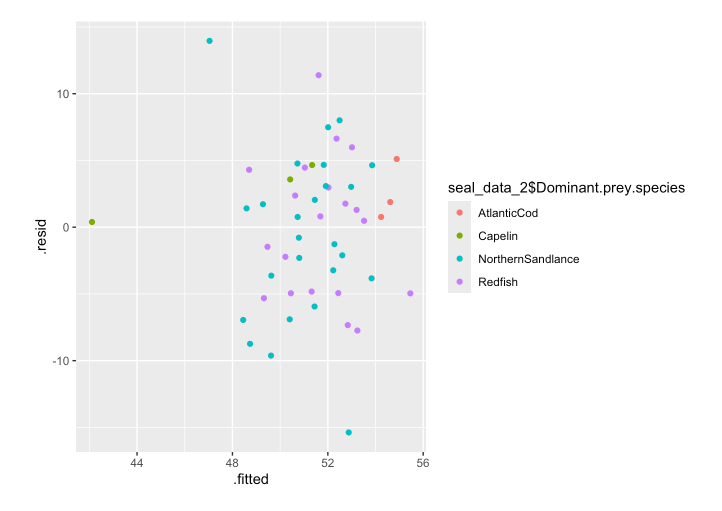


Figure 13 - Residual for Reduced Dataset (White Hake and Pollock removed).

The lowest value of Caplin is now an outlier and has very high leverage. It is most likely best to remove the lowest value of Capelin or remove the Capelin coefficient all together. Below we load a new dataset with the lowest value of Caplelin removed.

```
seal_data_3 <- read.csv("/Users/peterbraithwaite/Desktop/IDPhD Classes/Stat 5620_Updated/
summary(seal_data_2)</pre>
```

```
MomID
                        Year
                                  Dietary.energy.density Diet.diversity
Min.
       :
           19.0
                  Min.
                          :1996
                                  Min.
                                          :5.207
                                                          Min.
                                                                  :0.1788
                                  1st Qu.:5.686
1st Qu.: 701.5
                   1st Qu.:2002
                                                          1st Qu.:0.3061
Median: 4269.0
                  Median :2011
                                  Median :5.805
                                                          Median :0.3518
       : 4897.8
                          :2008
                                          :5.782
                                                                  :0.3531
Mean
                  Mean
                                  Mean
                                                          Mean
3rd Ou.: 9020.5
                   3rd Qu.:2013
                                  3rd Qu.:5.904
                                                          3rd 0u.:0.3959
Max.
       :10690.0
                  Max.
                          :2015
                                          :6.517
                                                          Max.
                                                                  :0.5290
                                  Max.
Dominant.prey.species
                          Mom.Age
                                          Pup.sex
                                                        Pup.Wean.Mass
Length:51
                       Min.
                              : 9.00
                                                        Min.
                                                               :37.50
                                       Min.
                                               :1.000
Class :character
                       1st Qu.:23.00
                                       1st Qu.:1.000
                                                        1st Qu.:47.00
Mode :character
                       Median :25.00
                                       Median :1.000
                                                        Median :51.50
                       Mean
                              :23.22
                                               :1.431
                                                               :51.36
                                       Mean
                                                        Mean
```

3rd Qu.:26.00 3rd Qu.:2.000 3rd Qu.:55.50 Max. :31.00 Max. :2.000 Max. :63.00

# Fit Full Model with Reduced Data Set

Removed Lowest Value of Capelin species from data.

```
Q4_Full= lm(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.densi summary(Q4_Full)
```

#### Call:

```
lm(formula = Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity +
    Dietary.energy.density + Year + Mom.Age + Pup.sex, data = seal_data_3)
```

#### Residuals:

```
Min 10 Median 30 Max -14.7728 -3.0865 0.7264 3.0387 12.9970
```

#### Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	-311.0480	401.3625	-0.775	0.443
Dominant.prey.speciesCapelin	-1.3014	7.0288	-0.185	0.854
Dominant.prey.speciesNorthernSandlance	-5.1433	5.6712	-0.907	0.370
Dominant.prey.speciesRedfish	-4.6264	5.3875	-0.859	0.395
Diet.diversity	-20.3147	13.8136	-1.471	0.149
Dietary.energy.density	-3.6124	7.2924	-0.495	0.623
Year	0.1990	0.2056	0.968	0.339
Mom.Age	-0.1911	0.2445	-0.782	0.439
Pup.sex	-0.1885	1.7953	-0.105	0.917

```
Residual standard error: 5.962 on 41 degrees of freedom Multiple R-squared: 0.1651, Adjusted R-squared: 0.002236 F-statistic: 1.014 on 8 and 41 DF, p-value: 0.441
```

```
# p-value: 0.002236
# Adjusted R-squared:0.004332
```

# **Step Function (Forward)**

We now used the Step Function to determine the best fit model using AIC.

```
fwd.model = step (Q4_Full, direction='forward')
```

Start: AIC=186.61
Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
 Year + Mom.Age + Pup.sex

The forward step retained all variables with AIC = 186.61.

## **Step Function (Backward)**

```
backward.model = step(Q4_Full, direction='backward')
Start: AIC=186.61
Pup.Wean.Mass ∼ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
   Year + Mom.Age + Pup.sex
                        Df Sum of Sq
                                        RSS
                                               ATC
Dominant.prey.species
                         3
                              52.481 1509.7 182.38
- Pup.sex
                         1
                               0.392 1457.6 184.62
Dietary.energy.density 1
                              8.721 1465.9 184.91
                              21.729 1478.9 185.35
Mom.Age
                         1
                              33.316 1490.5 185.74
Year
<none>
                                     1457.2 186.61
                         1 76.867 1534.1 187.18
Diet.diversity
Step: AIC=182.38
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age + Pup.sex
                        Df Sum of Sq
                                        RSS
                                               AIC
- Pup.sex
                               1.992 1511.7 180.45
                         1
                              39,426 1549,1 181,67
- Year
- Mom.Age
                              56.342 1566.0 182.21
                                     1509.7 182.38
<none>
- Dietary.energy.density 1 77.795 1587.5 182.89
- Diet.diversity
                         1
                             116.741 1626.4 184.10
Step: AIC=180.45
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age
                        Df Sum of Sq
                                        RSS
                                               AIC
- Year
                         1
                              45.154 1556.8 179.92
- Mom.Age
                         1
                              56,672 1568,3 180,29
                                     1511.7 180.45
<none>
Dietary.energy.density 1
                             78,067 1589,7 180,97
                         1 117.958 1629.6 182.20
- Diet.diversity
Step: AIC=179.92
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Mom.Age
```

Df Sum of Sa

RSS

AIC

```
Mom.Age
                         1 18.078 1574.9 178.50
- Dietary.energy.density 1 48.992 1605.8 179.47
                                     1556.8 179.92
- Diet.diversity 1 124.319 1681.1 181.76
Step: AIC=178.5
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density
                        Df Sum of Sa
                                        RSS
                                               AIC
                                     1574.9 178.50
<none>
                              66.647 1641.5 178.57
Dietary.energy.density 1
                             159,201 1734,1 181,31
Diet.diversity
                         1
The backward step retained the Diet Diversity and Dietary, energy, density variables with AIC=178.5. It
should be noted that this the best AIC value so far.
Explore Best Backward Step Linear Model
```

```
Q4_Reduced = lm(Pup.Wean.Mass ~ Dietary.energy.density + Diet.diversity,
                data = seal_data_3)
summary(Q4_Reduced)
```

```
Call:
```

```
lm(formula = Pup.Wean.Mass ~ Dietary.energy.density + Diet.diversity,
   data = seal_data_3)
```

#### Residuals:

```
Min
             1Q
                 Median
                            30
                                   Max
-15.3747 -4.5794
                 0.8175 4.1879 14.0677
```

#### Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                     98.933
                                29.029 3.408 0.00135 **
Dietary.energy.density -6.614
                                4.689 -1.410 0.16503
                     -26.461
Diet.diversity
                                12.140 -2.180 0.03432 *
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 5.789 on 47 degrees of freedom Multiple R-squared: 0.0977, Adjusted R-squared: 0.0593 F-statistic: 2.545 on 2 and 47 DF, p-value: 0.08928

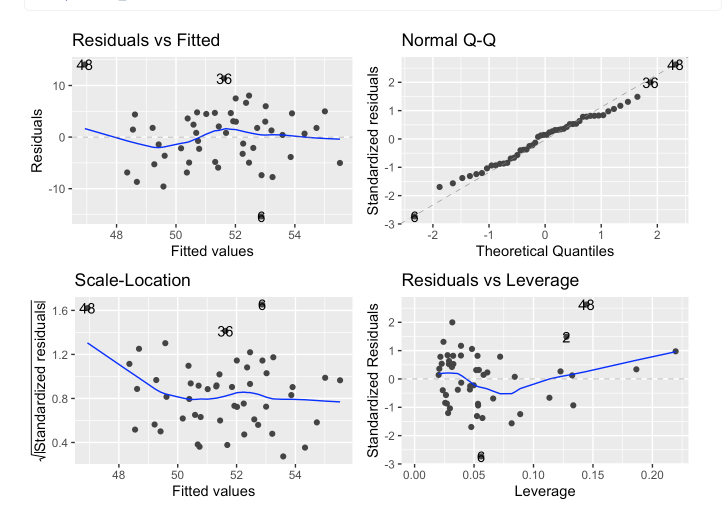
#### extractAIC(Q4\_Reduced)

#### [1] 3.000 178.496

```
# p-value: 0.08928
# Adjusted R-squared: 0.0593
```

# **Explore Residuals**

library(ggfortify)
autoplot(Q4\_Reduced)



ggplot(Q4\_Reduced, aes(x = .fitted, y = .resid, color = seal\_data\_3\$Dominant.prey.species

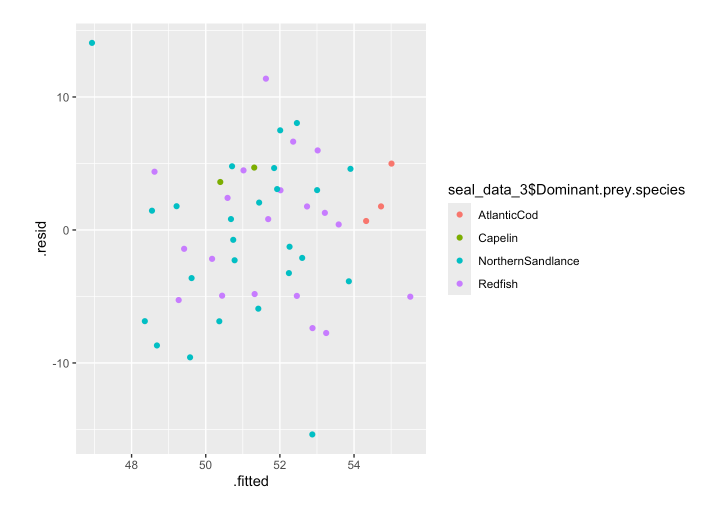


Figure 16 - Residual for Reduced Dataset (White Hake and Pollock removed completely and lowest value of Capeline).

From this figure it seems we have a fairly balanced residuals now. Although Northerbn Sandlance is show some extreme values (both large and small) for risk of overfitting the model we will stop removing variables and observations.

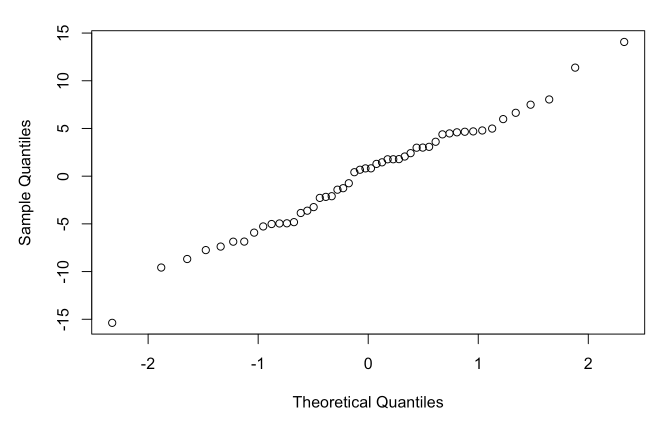
Now, since this is a General Linear Model and not a Generalized Linear Model we will need to check for the assumptions including linearity, independence of errors, homoscedasticity, and normality.

# **Testing Assumptions**

# **Checking for Normality**

qqnorm(Q4\_Reduced\$residuals)

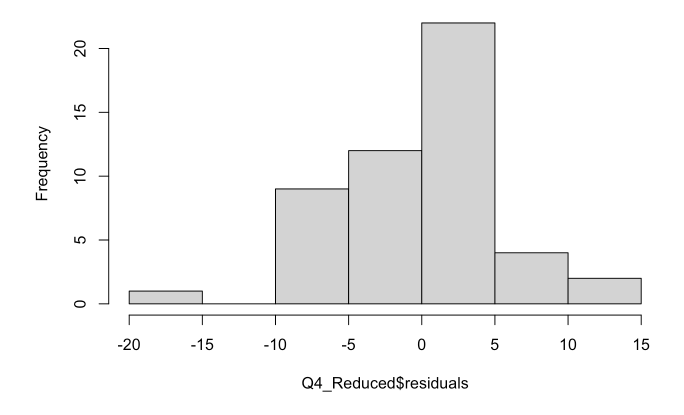
# **Normal Q-Q Plot**



Q-Q plot appears to show normality

hist(Q4\_Reduced\$residuals)

### Histogram of Q4\_Reduced\$residuals



Histogram appears to show normality

# Shapiro-Wilk's Test

```
shapiro.test(Q4_Reduced$residuals)
```

Shapiro-Wilk normality test

data: Q4\_Reduced\$residuals
W = 0.98681, p-value = 0.8456

From this W = 0.98681 and p-value = 0.8456. Given the p-value is not significant their is not a variation of the assumption of normality.

From these 3 explorations we are now confident in normality of the data.

# **Checking for Homogeneity of Variance**

Using Breusch\_Pagan Test

```
require (lmtest)
```

```
Loading required package: lmtest

Loading required package: zoo

Attaching package: 'zoo'

The following objects are masked from 'package:base':

as.Date, as.Date.numeric

bptest(Q4_Reduced)

studentized Breusch-Pagan test
```

```
data: Q4_Reduced
BP = 2.0797, df = 2, p-value = 0.3535
```

```
# BP = 2.0797
# df = 2
# p-value = 0.3535
```

Thus the p-value is greater then 0.05 so we accept the null hypothesis and assume we have homoscedasticity.

## **Checking for Independence of Predictor Variables**

Using the Dublin-Watson

```
dwtest(Q4_Reduced)
```

Durbin-Watson test

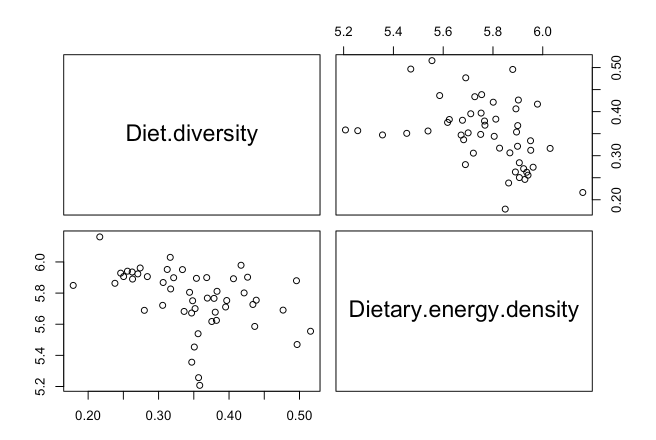
```
data: Q4_Reduced
DW = 2.2815, p-value = 0.792
alternative hypothesis: true autocorrelation is greater than 0
```

```
# DW = 2.2815
# p-value = 0.792
```

This p-value suggests very little to no autocorrelation.

# **Checking for Correlation of the Variables**

**Graphically Explore for Correlation** 



Data points look random and not correlated.

#### **Numerically Explore for Correlation**

```
cor.test( ~Diet.diversity + Dietary.energy.density, data = seal_data_3)
```

Pearson's product-moment correlation

```
data: Diet.diversity and Dietary.energy.density
t = -3.0499, df = 48, p-value = 0.00372
alternative hypothesis: true correlation is not equal to 0
95 percent confidence interval:
    -0.6125589 -0.1402952
sample estimates:
        cor
-0.402909
```

```
# t = -0.35989
# df = 74
```

```
\# p-value = 0.72
```

From this we can conclude that the variables are not correlated.

Thus, all our assumptions are met for the Linear Model and our findings can be accepted.

# **Generalized Linear Model**

Given we will be fitting a Gaussian (normal) distribution Generalized Linear Model it seeem safe to assume that the model will demonstrate the same data issues as the Linear Model with regard to residual outliers. For this reason we start with the reduced dataset that eliminated the the Dominant Prey Species White Hake, Pollock, and the lowest value of Capelin.

# Fitting Full Generalized Linear Model (Gaussian Family)

#### **All Predictor Variables**

```
GLM_1 = glm (Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity + Dietary.energy.dens
summary (GLM 1)
```

```
glm(formula = Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity +
    Dietary.energy.density + Year + Mom.Age + Pup.sex + MomID,
    family = gaussian, data = seal_data_3)
```

```
Coefficients:
```

Call:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                                      -1.076e+02 4.092e+02 -0.263
                                                                     0.7940
                                      -4.464e+00 7.103e+00 -0.629
Dominant.prey.speciesCapelin
                                                                     0.5332
Dominant.prey.speciesNorthernSandlance -6.486e+00 5.592e+00 -1.160
                                                                     0.2530
Dominant.prey.speciesRedfish
                                      -6.424e+00 5.363e+00 -1.198
                                                                     0.2380
                                      -2.408e+01 1.366e+01 -1.762
Diet.diversity
                                                                     0.0856 .
                                      -3.922e+00 7.123e+00 -0.551
                                                                     0.5850
Dietary.energy.density
Year
                                       9.835e-02 2.090e-01
                                                             0.471
                                                                     0.6405
                                      -1.175e-01 2.425e-01 -0.484
                                                                     0.6307
Mom.Age
Pup.sex
                                      -4.899e-01 1.762e+00 -0.278
                                                                     0.7824
MomID
                                       4.208e-04 2.432e-04 1.730
                                                                     0.0913 .
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 33.89238)
```

Null deviance: 1745.4 on 49 degrees of freedom Residual deviance: 1355.7 on 40 degrees of freedom

ATC: 328.9

```
# AIC = 328.9
```

From here we will use the GLM\_1 model for the Step Funtion to reduce the number of predictor variables and increase the accuracy and efficiency of the model.

#### **Step Function (Backward)**

```
backward.model = step(GLM_1, direction='backward')
Start: AIC=328.9
Pup.Wean.Mass ∼ Dominant.prey.species + Diet.diversity + Dietary.energy.density +
   Year + Mom.Age + Pup.sex + MomID
                       Df Deviance
                                      ATC
Dominant.prey.species
                        3 1412.7 324.96
Pup.sex
                        1 1358.3 326.99
                        1 1363.2 327.17
- Year
Mom.Age
                        1 1363.7 327.19

    Dietary.energy.density 1 1366.0 327.27

                            1355.7 328.90
<none>
- MomID
                        1 1457.2 330.51
Diet.diversity
                        1 1461.0 330.64
Step: AIC=324.96
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age + Pup.sex + MomID
                       Df Deviance AIC
                        1 1416.7 323.10
Pup.sex
- Year
                        1 1422.6 323.30
                        1 1444.5 324.07
Mom.Age
                           1412.7 324.96
<none>
- MomID
                        1 1509.7 326.28
- Dietary.energy.density 1 1521.1 326.65
                        1 1571.7 328.29
- Diet.diversity
Step: AIC=323.1
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Year +
   Mom.Age + MomID
                       Df Deviance
                                      AIC
- Year
                        1 1430.0 321.57
                        1 1449.0 322.23
Mom.Age
<none>
                            1416.7 323.10
- MomID
                        1 1511.7 324.34
- Dietary.energy.density 1 1525.0 324.78
Diet.diversity
                        1 1576.8 326.45
```

```
Step: AIC=321.57
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + Mom.Age +
   MomID
                        Df Deviance
                                      ATC
                         1 1449.0 320.23
- Mom.Age
                            1430.0 321.57
<none>
- Dietary.energy.density 1 1525.9 322.81
- MomID
                         1 1556.8 323.81
Diet.diversity
                         1 1601.1 325.21
Step: AIC=320.23
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + MomID
                        Df Deviance
                                      ATC
<none>
                             1449.0 320.23

    Dietary.energy.density 1 1569.6 322.22
```

1 1574.9 322.39

1 1661.7 325.07

The backward step retained the Diet Diversity and Dietary.energy.density variables with AIC AIC = 320.23. It should be noted this is the same predictor variables that were selected in the Step Function as the Linear Model but the AIC is much worse then the reduced Linear Model with AIC = 178.5. This is because the Linear Model is a less complex model and thus has a lower AIC.

## **Explore Backward Step Linear Model**

MomID

AIC: 322.39

Diet.diversity

```
Call:
glm(formula = Pup.Wean.Mass ~ Dietary.energy.density + Diet.diversity,
   family = gaussian, data = seal_data_3)
Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
                        98.933
(Intercept)
                                  29.029 3.408 0.00135 **
Dietary.energy.density -6.614
                                  4.689 -1.410 0.16503
Diet.diversity
                       -26.461
                                 12.140 -2.180 0.03432 *
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for gaussian family taken to be 33.50838)
   Null deviance: 1745.4 on 49
                                degrees of freedom
Residual deviance: 1574.9 on 47
                                degrees of freedom
```

```
# AIC = 322.39
```

# **Explore Residuals**

library(ggfortify)
autoplot(GLM\_2)

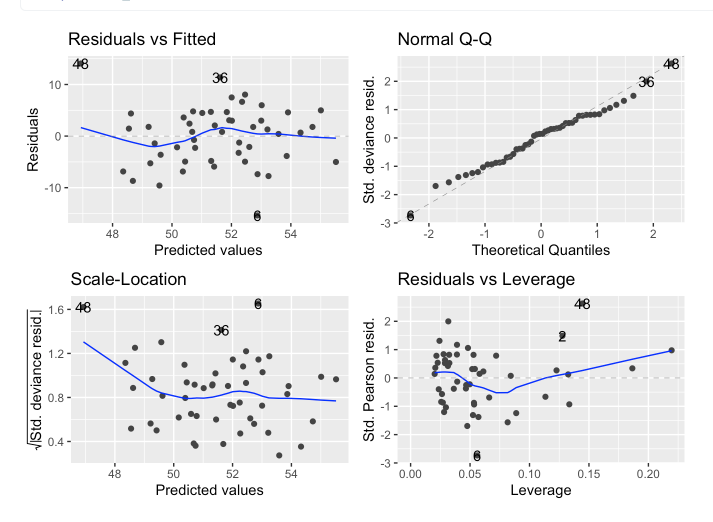


Figure 17 - Residual for Generalized Linear Model With Reduced Dataset

As we might expect the residuals for this Gaussian Generalized Linear Model are very similar to the residuals for the Linear Model above.

```
ggplot(GLM_2, aes(x = .fitted, y = .resid, color = seal_data_3$Dominant.prey.species )) +
```

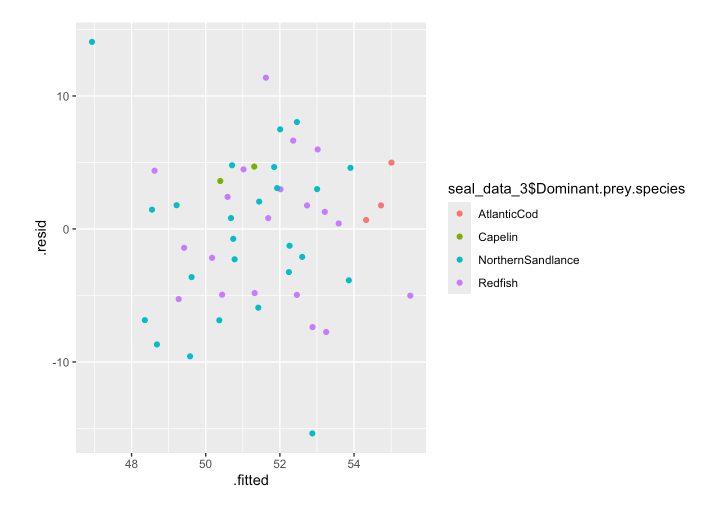


Figure 18 - Residual for Generalized Linear Model With Reduced Dataset

Again, as we might expect the residuals for this Gaussian Generalized Linear Model are very similar to the residuals for the Linear Model above. Given the Generalized Linear Model is creating similar residuals with the a same predictor variables as the Linear Model but with a less desirable AIC (GLM = 322.39 vs. LM = 178.5), it

seems the more simple Linear Model should be used.

# **Fit Linear Mixed Model**

The Linear Model may benefit from Random Effects on the Dominant Prey Species variable given the different prey species would have an effect on the other two remaining variables, Dietary Energy Density and Diet Diversity.

# Linear Mixed Model With Both Diet Diversity and Dietary Energy Loss Random Effects Based on Cluster Variable Dominant Prey Species

require(lme4)

```
GLMM_1 = lmer(Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Dietary.energy)
boundary (singular) fit: see help('isSingular')
 summary(GLMM_1)
Linear mixed model fit by REML ['lmerMod']
Formula:
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Dietary.energy.density |
    Dominant.prey.species) + (Diet.diversity | Dominant.prey.species)
   Data: seal_data_3
REML criterion at convergence: 301.5
Scaled residuals:
    Min
             10 Median
                             30
                                   Max
-2.6560 -0.7911 0.1412 0.7235 2.4302
Random effects:
                                               Variance Std.Dev. Corr
 Groups
                        Name
 Dominant.prey.species (Intercept)
                                               1.228e-02 0.11081
                         Dietary.energy.density 3.838e-04 0.01959 -1.00
 Dominant.prey.species.1 (Intercept)
                                               0.000e+00 0.00000
                         Diet.diversity
                                               2.394e-04 0.01547
                                                                   NaN
 Residual
                                               3.351e+01 5.78864
Number of obs: 50, groups: Dominant.prey.species, 4
Fixed effects:
                       Estimate Std. Error t value
(Intercept)
                        98.933
                                   29.029
                                            3.408
Diet.diversity
                       -26.461
                                   12.140 -2.180
Dietary.energy.density -6.613
                                   4.689 -1.410
Correlation of Fixed Effects:
            (Intr) Dt.dvr
Diet.dvrsty -0.522
Dtry.nrgy.d -0.991 0.403
optimizer (nloptwrap) convergence code: 0 (OK)
boundary (singular) fit: see help('isSingular')
 extractAIC(GLMM 1)
```

#### [1] 10.0000 334.3899

```
# AIC = 337.7947
```

# Linear Mixed Model With Dietary Energy Loss Random Effects Based on Cluster Variable Dominant Prey Species

```
GLMM 2 = lmer(Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Dietary.energy
boundary (singular) fit: see help('isSingular')
 summary(GLMM_2)
Linear mixed model fit by REML ['lmerMod']
Formula:
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Dietary.energy.density |
    Dominant.prey.species)
   Data: seal_data_3
REML criterion at convergence: 301.5
Scaled residuals:
             10 Median
    Min
                             30
                                    Max
-2.6560 -0.7911 0.1412 0.7235 2.4302
Random effects:
                                              Variance Std.Dev. Corr
 Groups
                       Name
 Dominant.prey.species (Intercept)
                                              1.420e-03 0.037686
                       Dietary.energy.density 4.228e-05 0.006502 -1.00
 Residual
                                              3.351e+01 5.788642
Number of obs: 50, groups: Dominant.prey.species, 4
Fixed effects:
                       Estimate Std. Error t value
                                             3.408
(Intercept)
                         98.933
                                    29.029
Diet.diversity
                        -26.461
                                    12.140 -2.180
Dietary.energy.density -6.614
                                     4.689 -1.410
Correlation of Fixed Effects:
            (Intr) Dt.dvr
Diet.dvrsty -0.522
Dtry.nrgy.d -0.991 0.403
optimizer (nloptwrap) convergence code: 0 (OK)
boundary (singular) fit: see help('isSingular')
 extractAIC(GLMM_2)
```

[1]

7.0000 328.3899

# AIC = 328.3899

# Linear Mixed Model With Diet Diversity Random Effects Based on Cluster Variable Dominant Prey Species

```
GLMM 3 = lmer(Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Diet.diversity
boundary (singular) fit: see help('isSingular')
 summary(GLMM 3)
Linear mixed model fit by REML ['lmerMod']
Formula:
Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (Diet.diversity |
    Dominant.prey.species)
   Data: seal_data_3
REML criterion at convergence: 301.5
Scaled residuals:
             10 Median
    Min
                             30
                                    Max
-2.6560 -0.7911 0.1412 0.7235 2.4302
Random effects:
                                      Variance Std.Dev. Corr
 Groups
                       Name
 Dominant.prey.species (Intercept)
                                      0.000e+00 0.000e+00
                       Diet.diversity 2.911e-17 5.395e-09 NaN
 Residual
                                      3.351e+01 5.789e+00
Number of obs: 50, groups: Dominant.prey.species, 4
Fixed effects:
                       Estimate Std. Error t value
(Intercept)
                         98.933
                                    29.029
                                             3.408
Diet.diversity
                        -26.461
                                    12.140 -2.180
Dietary.energy.density -6.614
                                     4.689 -1.410
Correlation of Fixed Effects:
            (Intr) Dt.dvr
Diet.dvrsty -0.522
Dtry.nrgy.d -0.991 0.403
optimizer (nloptwrap) convergence code: 0 (OK)
boundary (singular) fit: see help('isSingular')
 extractAIC(GLMM_3)
```

#### [1] 7.0000 328.3899

```
# AIC = 328.3899
```

# Linear Mixed Model With Intercept Random Effects Based on Cluster Variable Dominant Prey Species

```
GLMM 4 = lmer(Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (1 | Dominant.pr
boundary (singular) fit: see help('isSingular')
 summary(GLMM_4)
Linear mixed model fit by REML ['lmerMod']
Formula: Pup.Wean.Mass ~ Diet.diversity + Dietary.energy.density + (1 |
    Dominant.prey.species)
  Data: seal_data_3
REML criterion at convergence: 301.5
Scaled residuals:
    Min
             1Q Median
                             30
                                    Max
-2.6560 -0.7911 0.1412 0.7235 2.4302
Random effects:
 Groups
                       Name
                                   Variance Std.Dev.
 Dominant.prey.species (Intercept) 0.00
                                            0.000
                                   33.51
                                            5.789
Number of obs: 50, groups: Dominant.prey.species, 4
Fixed effects:
                       Estimate Std. Error t value
(Intercept)
                        98.933
                                    29.029 3.408
Diet.diversity
                       -26.461
                                    12.140 -2.180
Dietary energy density -6.614
                                     4.689 -1.410
Correlation of Fixed Effects:
            (Intr) Dt.dvr
Diet.dvrsty -0.522
Dtry.nrgy.d -0.991 0.403
optimizer (nloptwrap) convergence code: 0 (OK)
boundary (singular) fit: see help('isSingular')
extractAIC(GLMM 4)
Warning in optwrap(optimizer, devfun, x@theta, lower = x@lower, calc.derivs =
```

TRUE, : convergence code 3 from bobyqa: bobyqa -- a trust region step failed to

[1] 5.0000 324.3899

reduce q

```
# AIC = 324.3899
```

So, in summary, none of the Linear Mixed Models with Random Effect on the cluster variavble Dominant Prey Species resulted in a more desirable AIC. The best Mixed Model model had an AIC = 324.3899 but the best Linear Model had an AIC=178.496. Thus the less complex Linear Model is the most efficient and most accurate model.

# **Cross Validation LM**

It has been determined that a simple Linear Model using a maternal mother seal's Dietary Energy Density and Diet Diversity are the best predictor variables for Pup Wean Mass. The model Im(Pup.Wean.Mass ~ Dietary.energy.density + Diet.diversity, data = seal\_data\_3) has shown an AIC = 178.496.

We will now use Cross Validation to assess how well we expect this model to preform when predicting future Pup Wean Mass changes.

```
require(modelr)
```

Loading required package: modelr

```
require(caret)
```

Loading required package: caret

Loading required package: lattice

```
require(readr)

# Data
attach(seal_data_3)

# Set Random Seed
set.seed(1980)

# Create Index Matrix (80% train data and 20% test data )
index = createDataPartition(seal_data_3$Pup.Wean.Mass, p = .8, list = FALSE, times = 1)
summary(seal_data_3$Pup.Wean.Mass)
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 37.50 47.62 52.00 51.54 55.50 63.00
```

```
print(seal_data_3$Pup.Wean.Mass)
```

```
[1] 50.5 60.5 50.0 54.5 45.5 37.5 54.5 59.0 50.5 51.0 58.5 59.0 56.0 55.0 56.0 [16] 51.5 59.5 45.5 50.0 47.5 53.5 49.0 55.0 52.5 54.0 55.0 53.0 45.5 56.5 60.0
```

```
[31] 48.0 55.5 56.5 48.5 46.5 63.0 43.5 55.5 54.0 51.0 41.5 48.0 53.0 40.0 45.5 [46] 44.0 40.0 61.0 46.0 50.0
```

```
# Convert data frame
data_frame = as.data.frame(seal_data_3)

# Create a data frame for the train data that is 80%
train_data_frame = data_frame[index,]

# Create a data frame for the test data that is 20% ("-" before index mean everything but test_data_frame = data_frame[-index,]

# Convert response variable in both train and test data frame to a factor train_data_frame$Pup.Wean.Mass = as.factor (train_data_frame$Pup.Wean.Mass)
test_data_frame$Pup.Wean.Mass = as.factor (test_data_frame$Pup.Wean.Mass)

# Ensure the response variables classes are factors class (train_data_frame$Pup.Wean.Mass)
```

#### [1] "factor"

```
# "factor"
class (test_data_frame$Pup.Wean.Mass)
```

#### [1] "factor"

```
# "factor"

# Specify type of training method used and the number of folds
control_specs = trainControl(method = "cv", number = 11 , savePredictions = "all")

# Set Random Seed
set.seed(1980)

require(randomForest)
```

```
Loading required package: randomForest
randomForest 4.7-1.2

Type rfNews() to see new features/changes/bug fixes.

Attaching package: 'randomForest'

The following object is masked from 'package:ggplot2':
```

```
margin
The following object is masked from 'package:dplyr':
```

#### combine

#### Linear Regression

```
50 samples
2 predictor

No pre-processing
Resampling: Cross-Validated (11 fold)
Summary of sample sizes: 45, 46, 46, 44, 46, 46, ...
Resampling results:
```

RMSE Rsquared MAE 5.434521 0.2021304 4.563434

Tuning parameter 'intercept' was held constant at a value of TRUE

```
# 50 samples
# 2 predictor

# RMSE    Rsquared    MAE
# 5.434521    0.20213048    4.563434

# We now apply the model to the test_data_frame we created from 20%
# of the data that the new model created from the 11 folds has not yet seen

# Predict outcome using model from train_data_frame applied to test_data_frame
Predict_Data = predict (model_cross, newdata = test_data_frame)

# predictions results

#. 52.74225 51.24721 50.67234 57.30204 51.49974 49.31590 50.08707 48.16223

test_data_frame$Pup.Wean.Mass
```

```
[1] 59 59.5 49 55 52.5 41.5 48 40 Levels: 40 41.5 48 49 52.5 55 59 59.5
```

```
# 59 59.5 49 55 52.5 41.5 48 40

#Creates vectors having data points
predicted_value <- (c(52.74225, 51.24721, 50.67234, 57.30204, 51.49974, 49.31590, 50.0870
expected_value <- (c(59, 59.5, 49, 55, 52.5, 41.5, 48, 40))

Cross_Validation_Corrolation = cor (predicted_value, expected_value)

### 0.6596914

R2 = (Cross_Validation_Corrolation^2 )

### 0.4351927 (43.52% of variance)
```

From this cross validation procedure we can see that the Linear Model has a validation score of about 0.66 which mean the Linear Model would do a fair job at predicting Pup Mean Mass based on the maternal mother seal's Dietary Energy Density and Diet Diversity. The estimated R-squared value suggests the model would account for about 43.5% of the variance of Pup Mean Mass.

### **Cross Validation GLM**

Cross Validation was explored for the best fit Generalized Linear Model.

```
require(modelr)
require(caret)
require(readr)

# Data
attach(seal_data_3)
```

The following objects are masked from seal\_data\_3 (pos = 4):

Diet.diversity, Dietary.energy.density, Dominant.prey.species,
Mom.Age, MomID, Pup.sex, Pup.Wean.Mass, Year

```
# Set Random Seed
set.seed(1984)

# Create Index Matrix (80% train data and 20% test data )
index = createDataPartition(seal_data_3$Pup.Wean.Mass, p = .8, list = FALSE, times = 1)
summary(seal_data_3$Pup.Wean.Mass)
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 37.50 47.62 52.00 51.54 55.50 63.00
```

```
print(seal data 3$Pup.Wean.Mass)
[1] 50.5 60.5 50.0 54.5 45.5 37.5 54.5 59.0 50.5 51.0 58.5 59.0 56.0 55.0 56.0
[16] 51.5 59.5 45.5 50.0 47.5 53.5 49.0 55.0 52.5 54.0 55.0 53.0 45.5 56.5 60.0
[31] 48.0 55.5 56.5 48.5 46.5 63.0 43.5 55.5 54.0 51.0 41.5 48.0 53.0 40.0 45.5
[46] 44.0 40.0 61.0 46.0 50.0
# Convert data frame
data frame = as.data.frame(seal data 3)
# Create a data frame for the train data that is 80%
train data frame = data frame[index,]
# Create a data frame for the test data that is 20% ("-" before index mean everything but
test data frame = data frame[-index,]
# Convert response variable in both train and test data frame to a factor
train data frame$Pup.Wean.Mass = as.factor (train data frame$Pup.Wean.Mass)
test_data_frame$Pup.Wean.Mass = as.factor (test_data_frame$Pup.Wean.Mass)
# Ensure the response variables classes are factors
class (train data frame$Pup.Wean.Mass)
[1] "factor"
# "factor"
class (test_data_frame$Pup.Wean.Mass)
[1] "factor"
# "factor"
# Specify type of training method used and the number of folds
control_specs = trainControl(method = "cv", number = 11 , savePredictions = "all")
# Set Random Seed
set.seed(1984)
require(randomForest)
# now we train the model with the best fit GLM above
```

# GLM\_Final = glm (seal\_data\$Pup.Wean.Mass ~ seal\_data\$Dominant.prey.species + seal\_data\$

```
model_cross_2 = train(Pup.Wean.Mass ~ Dominant.prey.species + Diet.diversity,
                     method = "glm", family = gaussian, data = seal_data_3, trControl = co
 print (model_cross_2)
Generalized Linear Model
50 samples
 2 predictor
No pre-processing
Resampling: Cross-Validated (11 fold)
Summary of sample sizes: 46, 47, 45, 45, 44, 45, ...
Resampling results:
  RMSE
            Rsquared
                       MAE
  5.342752 0.3885031 4.388397
 # 50 samples
 # 2 predictor
 # RMSE
             Rsquared
                         MAE
 # 5.816687 0.4291658 4.852387
 # We now apply the model to the test_data_frame we created from 20%
 # of the data that the new model created from the 11 folds has not yet seen
```

```
# Predict outcome using model from train_data_frame applied to test_data_frame
Predict_Data_2 = predict (model_cross_2, newdata = test_data_frame)

# predictions results

#. 52.74225 51.24721 50.67234 57.30204 51.49974 49.31590 50.08707 48.16223

test_data_frame$Pup.Wean.Mass
```

[1] 37.5 50.5 55 51.5 60 56.5 43.5 53 Levels: 37.5 43.5 50.5 51.5 53 55 56.5 60

```
# 59 59.5 49 55 52.5 41.5 48 40

#Creates vectors having data points
predicted_value <- (c(52.43245, 52.14797, 52.30886, 51.25847, 57.08296, 50.11807, 49.9749
expected_value <- (c(59, 59.5, 49, 55, 52.5, 41.5, 48, 40))
```

```
Cross_Validation_Corrolation = cor (predicted_value, expected_value)
### 0.4533892

R2 = (Cross_Validation_Corrolation^2)
### 0.2055618 (20.6% of variance)
```

From this cross validation procedure we can see that the Generalized Linear Model has a validation score of about 0.45 which mean the best fit GLM has a lower predictive accuracy then the Linear Model.

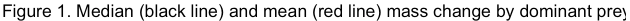
### Results

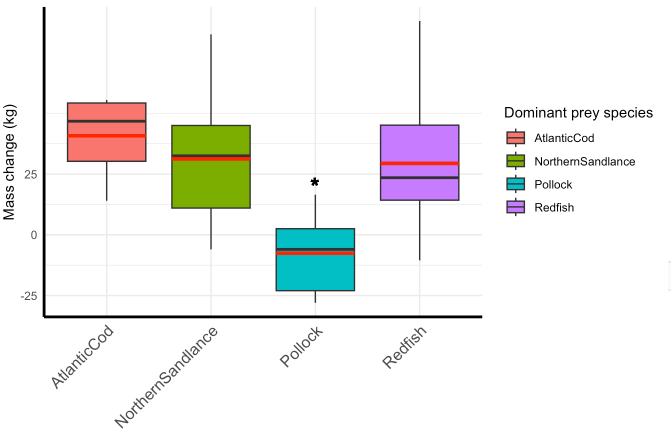
# Question 1 - Does variation in diet influence maternal mass change over the foraging period leading up to parturition in Northwest Atlantic grey seals?

After using step selection methods to determine the best model using diet to explain variation in maternal mass change during the foraging period leading up to parturition, the lowest AIC model retains diet diversity, year, and dominant prey species as significant covariates. Looking at the summary output, even though several years have mass change values that significantly differ from the expected mass change value, the effect of time is not of direct interest to our research question and is only present to account for changes in prey dynamics over time. As a result, the interpretation of this output is not relevant. However, we see that females who feed primarily on pollock experience significantly lower (p<0.05) mass change than the reference level, with the reference level in this case being females who primarily feed on cod. Being an identity link function, the model is essentially a simple linear model and there is no transformations applied to the response variable. As such, coefficient estimates can be interpreted directly. Individuals who feed on pollock experienced 47.5% lower mass change than individuals who primarily feed on cod, while females who fed primarily on redfish or sand lance experienced similar mass change to females who fed on cod. This effect is presented in figure 1.

```
ggplot(Data_Q1, aes(x = DomSpp, y = MassChange, fill = DomSpp)) +
  geom_boxplot() +
  stat_summary(
    fun = "mean", geom = "errorbar",
      aes(ymax = ..y.., ymin = ..y..),
    width = 0.75, color = "red", size = 1.2
) + geom_text(x = "Pollock", y = 20, label = "*",
            aes(x = x, y = y, label = label),
            color = "black", size = 6, fontface = "bold"
) +
  labs(
  title = "Figure 1. Median (black line) and mean (red line) mass change by dominant pr
  x = "Prey species",
  y = "Mass change (kg)", fill = "Dominant prey species"
) +
  theme_minimal() + theme(
```

```
axis.line = element_line(color = "black", size = 1), # Add axis lines
axis.text.x = element_text(size = 12, angle = 45, hjust = 1)
) +
scale_y_continuous(
breaks = seq(0, max(Data_Q1$MassChange), by = 25) - 50
)
```





It may also be worth noting that diet diversity is almost significant at p = 0.0556. While this p-value is not significant, the negative relationship between diet diversity and mass change may support the observed effects of dominant prey species, where individuals who primarily prey on cod, sand lance, or redfish are focusing efforts on these few profitable species, as reflected by more successful mass gain than individuals who feed on a more diverse array of prey species which include less profitable species. As a result, reduced diet diversity may reflect focus on highly profitable prey species and increased diet diversity may reflect a greater proportion of less profitable prey species.

Prey species

Cross-validation was used to varify the predictive power of the final model. The average root mean squared error (RMSE) after 7-fold cross-validation is 15.98, indicating the model generally predicts values within 16 kg of the true value. We can check the standard deviation to compare the average RMSE to:

#### [1] 26.15433

Compared to the standard deviation of mass change, the RMSE value is about 40% lower than the standard deviation, meaning the model predictions are better than using the population mean alone to predict mass change. However, compared to the overall mean mass change within our dataset (28.5 kg), there is relatively large prediction error. We can also visualize prediction accuracy with a fitted vs observed value plot (figure 2):

```
fit_mom <- fitted(mom_mod2)

# Get the observed response variable (actual values)
obs_mom <- Data_Q1$MassChange

# Plot the fitted vs. observed values
ggplot(data = Data_Q1, aes(x = fit_mom, y = obs_mom)) +
    geom_point(color = "black") +
    geom_abline(slope = 1, intercept = 0, color = "red", linetype = "dashed") + # Add 1:1
labs(
    title = "Figure 2. Fitted vs. Observed Values with 1:1 line",
    x = "Fitted Values",
    y = "Observed Values"
) +
    theme_minimal()</pre>
```

90
60
0
0
1-30
-25
0
Fitted Values

Figure 2. Fitted vs. Observed Values with 1:1 line

While figure 2 shows that the model predictions have consistent accuracy across the range of fitted values, as suggested by the average RMSE value obtained from cross-validation, there is reasonably large prediction error. Given that the model residuals suggest good model fit, the relatively large prediction error suggest the set of covariates used in this model are not sufficient in predicting maternal mass change. From this we can conclude that females who feed on redfish, sand lance, or cod experience greater mass change than individuals who feed on Pollock, but other factors outside of maternal diet, such as individual physiology or characteristics associated with individual foraging behavior, are likely responsible for explaining a large proportion of variation in maternal mass change.

# Question 2 - Does female diet predict pup weaning mass in Northwest Atlantic grey seals?

Although it was proposed that many predictor variables (2 continuous + 3 categorical for question one and 2 continuous + 5 categorical for question two) would allow greater computational potential to establish a well fit model, the added complexity of these variables was observed to significantly increase the AIC and potentially overfit the model. It was found, particularly in question two, that the simplicity of a Linear Model with a Least Square approach resulted in a simpler linear model with a significantly lower AIC.

In question 2 it was found that best fit Linear Model for Pup Wean Mass included the predictor variables Dietary Energy Density and Diet Diversity with an AIC = 178.496. When a Generalized Linear Model was explored with a Gaussian Family and Identity Link Function it was also found that the best fit model

included Dietary Energy Density and Diet Diversity predictor variable but the AIC was much less preferable at 322.39. The reasons for this decreased predicted fit is likely based on the more complex Maximum Likelihood Estimation used in Generalized Linear Models rather than the Linear Least Square method used in the Linear Models. Generalized Linear Mixed Models (GLMM) were also explored using the Dominant Prey Species as the Cluster Variable because it was assumed that this variable would likely have a effect on the prey energy density as well as the diet diversity based on feeding location ecosystems. The best fit GLMM was found to include Diet Diversity and Dietary Energy Density as fixed effects predictor variables and Dominant Prey Species as an intercept random effect variable which had a AIC of 324.3899. As a result it was determined that the best fit model to determine Pup Wean Mass was the Linear Model and this model was tested using Cross Validation. The correlation of predicted values to actual values showed a value of 0.6596914 meaning the Linear Model could be expected to predict Pup Wean Mass with about a 66 percent accuracy.

# **Conclusions**

In both question one, that explored the effect of diet on Maternal Mass Change, and question two, that explored the predictive ability of maternal diet on pup wean mass at partition, we found relatively large predictive errors in our models (cross-validation revealed RMSE of +/- 16kg for question 1 and 33% for question two). Although the data-set provided many predictive variables, many of these were categorical data and only two, Dietary Energy Density and Diet Diversity, where numeric in nature. In addition to this the number of observation in the dataset were limited (question one n=76, and question two n=56) which resulted in many rows of data but few columns with ultimately resulted in added complexity in our models and the need to reduce the number of predictor variable to prevent overfitting the model.

In question one the residuals suggest good model fit but the relatively large prediction error suggest the set of covariates used in this model are not sufficient in predicting maternal mass change or more observation would be required to fit a more reliable model. It was found from the data that females who fed on redfish, sand lance, or cod experience greater mass change than individuals who feed on Pollock, but other factors outside of maternal diet, such as individual physiology or characteristics associated with individual foraging behavior are likely responsible for a large proportion of variation in maternal mass change. Diet diversity appeared to have negative relationship with maternal mass change, possibly indicating a narrow diet focusing on more profitable prey species is beneficial in increasing mass gain, although these results were not quite statistically significant.

In question two it was determined through both AIC and Cross Validation that the simpler Linear Model had a higher accuracy for predicting Pup Wean Mass with the two available numeric predictor variables. Access to additional observation could increase the predictive ability of the Linear Model as well as allow the fitting of more complex model such as a Mixed Model without overfitting. Although we feel the models provided fair accuracy in predicting Pup Wean Mass at partition, the lack of number of observations may have significantly impacted the reliability and validity of the statistical analysis.

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