Data Analysis 3

Maksuda Aktar Toma, Jo Charbonneau, Ryan Lalicker November 25, 2024

Introduction

In this paper we will be looking at data related to calves. The data comes from an experiment designed to study the impact dietary treatments given to pregnant heifers had on the development of the calves. The study was conducted over a three year period and involved three different dietary treatments given to select groups of heifers in the final trimester. In total the data has 22 variables for 120 entires, though some data points are missing.

For more information on the experiment, the data, or any other files used in this paper see our Github page which can be found at https://github.com/RyanLalicker/Data-Analysis-2-STAT-325-825. The coding languages used in the paper are R and SAS. The corresponding code can be found in $Appendix\ A$ - $R\ Code$ and $Appendix\ B$ - $SAS\ Code$ respectively.

Variables

The experiment used three different dietary treatments. These were DDG, CON, and MET. For the first two trimesters the heifers were given one of seven developmental treatments, found in <code>Development.Treatment</code>, and then in the final trimester the each was given one of the three treatments mentioned above. This is recorded in the <code>Calan.Treatment</code> column of the dataset.

The heifers were placed into one of four pens by weight, which can be seen in the column Pen #. They were then artificially inseminated from an assigned sire, which we will assume was done randomly since the client says weight was not a factor. The sire is represented by the column of the same name and has six unique entries.

Upon the birth of the calves, several measurements were taken. These include the sex of the calf, weights taken at both birth and slaughter, and scores of both the calf's vigor and the ease of birth. The vigor score is on a scale of one to eleven where a score of one is very good and a score of ten or eleven indicates poor vitality for the calf. (Probo (2022)). The ease score goes from one to five where one indicates a quick and easy birth, two means a longer birth, three

means requires some assistance, and four or five indicates more assistance was needed. (Heins (2023)). Note, the variable names in the dataset line up with the descriptions above.

Other variables, such as the id of the calf, length of gestation for the heifer, and postmortem scoring such as hot carcass weight (HCW) are included as well. (Saner (2024)). Note two birthdays are included in the data, Birth.date and Birth.date.1. These variables will not be used in the models below so no further investigation was done on our part to determine the differences.

The client's main focus is the effect the third trimester treatment and the sex of a calf have on the calf's vigor score, ease of birth score, and final body weight. Therefore, these are the variables we will place more of an emphasis on, while exploring the effect some of the other variables may have.

Missing Values

The data contains some missing values. In total 53 rows in the dataset are missing at least one variable. Figure 1 shows which columns have the most missing data. As we can see the values for the variable DMI, which according to the USDA represents the dry matter intake for a cow, is missing for two-thirds of the entries. (USDA). Given the number of missing values is this large, it is probably best to not use this variable in our models. Some other variables, including the final body weight of the calf represented by Final.Calf.BW, are missing in 19 entries. Of the other four variables the client was most interested in, none have more than ten missing values.

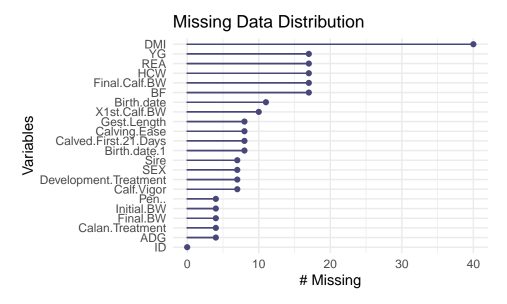


Figure 1: Chart counting the number of missing values for each variable within the data.

Cleaning the Dataset

Due to the missing values discussed above, we need to clean this dataset before continuing. There are several ways to handle missing values. One way is imputing them with some metric like the mean, median, or mode of the variable. Another is to just remove any rows with missing data.

We decided to cut all rows that contained missing values for variables we are interested in. These variables include the five variables the client is interested in, but also the pen number, sire, and the initial weight of the mother. The latter two are Sire and Initial.BW in the dataset. After removing missing values for these entries the dataset has 101 rows, which we feel is an ample amount for analyzing the data. Note all future figures and models come from this cleaned dataset and not the original.

We initially considered imputing the quantitative variables with the respective median values and using the mode for categorical variables as Memon, Wamala, and Kabano (2023) suggests. This has some issues though. For an example let's consider the third trimester treatment. The MET treatment was used in 40 cases, while the other two treatments were only used 38 times, meaning there are four missing values. If we mode impute this variable there will be 42 instances of the MET treatment. However, it seems very possible that the missing entries were split between the CON and DDG treatments to make an even 40 uses each. While imputing quantitative variables is less risky, we are not fully comfortable with that approach either since we are trying to analyze the data.

Summary Statistics

Let's take a closer look at what the three dependent variables the client is interested in. Figure 2 shows several summary statistics for each. Looking at the maximum values of the calving ease and calf vigor, we can see that the cleaned dataset does not contain any instances of poor scores for either. Both scores only goes from one to three. Note, the original dataset did have three instances of a vigor score of four or five, but each row was missing a final body weight so the entries were not included in the cleaned dataset. We can also see from the median and 75th percentiles that both seem very skewed towards the low end of the scale. While this is a good thing in terms of the health of the cows it could present some challenges for us later on.

				25th	75th		
Variable	Mean	Median	SD	Percentile	Percentile	Min	Max
Calving.Ease	1.118812	1	0.4071149	1	1	1	3
Calf.Vigor	1.257426	1	0.6107940	1	1	1	3
Final.Calf.BW	1291.584158	1292	128.9883148	1219	1365	932	1690

Figure 2: Summary statistics for dependent variables.

The final weight of calf is the third variable in Figure 2. The mean and median are relatively similar given the large standard deviation. While the previous two variables discussed give us some concerns about the skew, the final weight does not present the same issues. Further investigation into the approximate distribution of the final weight is needed though.

Let's look at a histogram and a Q-Q plot for the final weight of the calves in Figure 3. The bin width for the histogram comes from the Freedman-Diaconis rule. (William (2023)). The histogram appears to follow an approximately normal distribution. The Q-Q plot mostly follows this as most points follow the linear trend represented by the red line.

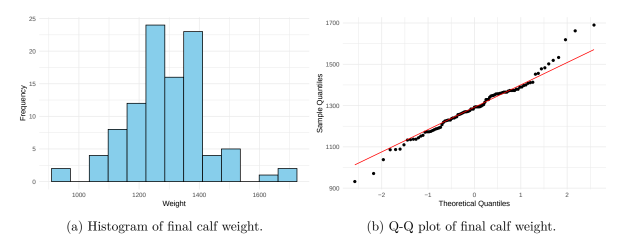


Figure 3: Plots used to check if the distribution of the final calf weight is normal.

Before moving on we want to look at plots of the scoring variables as well. While we suspect a heavy skew for each, the histograms in Figure 4 verify this. It is important to remember that these two variables are not continuous like the weight variable, so the types of models used will vary.

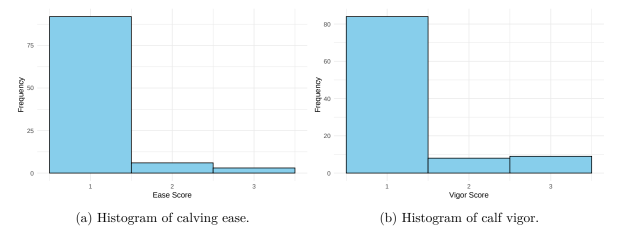


Figure 4: Histograms of scoring variables.

Exploring the Data

Before looking at potential models, let's explore how some of the variables interact with each other. While we will be able to include other explanatory variables, the client specifically mentions using the third trimester treatment and the sex of a calf as explanatory variables of interest. The table in Figure 5 shows the breakdown of treatment by sex. Note HFR stands for heifer and STR stands for steer. Although not every group has an equivalent number of subjects, this is nothing we are concerned about. Please note that the total occurrences per treatment are different than discussed above since rows containing missing values were removed.

	HFR	STR
CON DDG	19	15
DDG	14	20
MET	18	15

Figure 5: Table showing the breakdown of treatment by sex.

One of the key assumptions for regression models is that the explanatory variables should not be highly correlated to avoid multicollinearity. Here, we used a Pearson's Chi-squared test to examine the association between the categorical variables Calan.Treatment and SEX. The p-value (0.40811) suggests that there is no statistically significant association between these variables. While this result indicates that these variables are not strongly associated, it does not directly address multicollinearity, which is better assessed using metrics like Variance Inflation Factor (VIF) or condition indices in regression models. Therefore, we will do further

diagnostic checks for the model, such as calculating VIF values, to ensure multicollinearity is not an issue in the model.

Metric	Value
Statistic	1.79
Degrees of Freedom	2
P-Value	0.40811

Figure 6: Chi-squared test for treatment and sex.

Now let's consider how these variables affect the final Calf body weight. The boxplot shown in Figure 7 allows us to see this relationship graphically. We can see the steers are heavier on average than the heifers. The treatments seem to different variances as well, but their median values are not different by huge quantities. Both the CON and MET treatments had one steer large enough to be an outlier, while the DDG treatment had several outliers for both sexes.

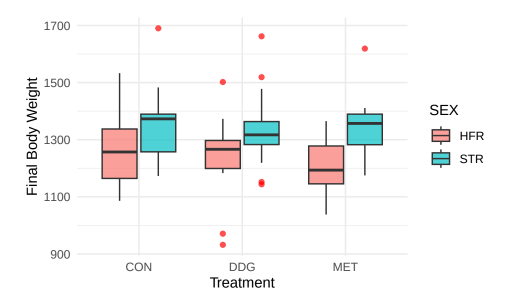
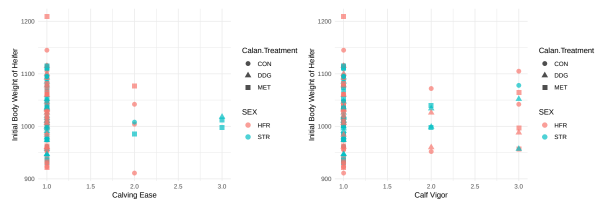


Figure 7: Boxplot of final calf body weight by treatment and sex.

Another variable we want to investigate graphically is the initial weight of the heifer that birthed the calf and see how it compares to both the ease and vigor score. In Figure 8 we can see this while also accounting for both the third trimester treatment with the shape of the data point and the sex of the calf with the color of the data point. This allows us to see, how the effect of the heifer's initial weight has, but also the trends of both treatment and sex.



- (a) Initial body weight vs. calf vigor.
- (b) Initial body weight vs. calving ease.

Figure 8: Scatterplot of heifer's initial body weight versus scoring variables, controlling for third trimester treatment and sex of the calf.

In both plots, we can see the initial weight of the mother heifer does not seem to have a huge effect on either score. It would appear that being heavier or lighter than average had little to no effect on getting a score other than one. The clustering of data at the lowest levels (1) for both calving ease and calf vigor highlights the imbalance in these outcome variables, potentially limiting the ability to draw robust conclusions. While this variable may be used in models later to test its significance more formally, we have doubts about its effect on either score variable.

Models for Calving Ease Score

Ordinal Logistic Regression Model

Since calving ease is a score from one to three in our dataset and one to five in real life, it can be considered an ordinal variable. This means instead of treating it as a quantitative variable as we did previously, it could be considered an ordered categorical variable. One method of modeling ordinal variables is with ordinal logistic regression. This uses one or more independent variables to predict the ordinal value of the dependent variable. A key assumption of ordinal logistic regression, outside of the dependent variable being ordinal, is no multicollinearity between independent variables. ("Ordinal Regression" (n.d.)).

Let's attempt to apply an ordinal logistic regression to the calving ease variable. In this simple case, we will use the third-trimester treatment and the calf's sex as the independent variables.

Ordinal logistic regression models are expressed as a logit function containing probabilities rather than a single variable. In this case, where Y is a random variable representing the

calving ease score and l = 1, 2, ..., 5 represents the possible values of Y in theory and l = 1, 2, 3 in our dataset, $\frac{P(Y \le l)}{P(Y > l)}$ is the cumulative probability of the easing score being less than or equal to level l versus greater than l. We can then write the model as

$$\ln \left(\frac{P(Y \leq l)}{P(Y > l)} \right) = \alpha_l - \beta_{trt} X_{trt} - \beta_{sex} X_{sex} - \beta_{trt:sex} (X_{trt} \times X_{sex})$$

The equation above the left side is the log odds or log of the cumulative probability, the logit function mentioned previously. On the right side of the equation α_l is the intercept of the model at the lth level, X_{trt} and X_{sex} represent the values for the treatment, and the sex respectively, and the β_i are slope values corresponding to the ith explanatory variable with $\beta_{trt:sex}$ being the interaction effect.

Some of the results of the model are displayed in Figure 9. Note the model treats CON and heifer as the starting values for the treatment and sex respectively. The first three rows of the coefficients table report the main effects of the other two treatments and steers. Calan.TreatmentDDG: SEXST respressints the interaction effect between the DDG treatment and steers, which is highly significant (p-value < 0.001). The interaction between the MET treatment and steers is not significant, though (p-value = 0.149). Overall, the model highlights significant effects for DDG and its interaction with calf sex, while other predictors do not show substantial influence.

	Value	Std. Error	t value	p-value
Calan.TreatmentDDG	-16.863	0.728	-23.147	0.000
Calan.TreatmentMET	-1.116	1.203	-0.928	0.353
SEXSTR	-0.926	1.208	-0.767	0.443
Calan.TreatmentDDG:SEXSTR	16.616	0.728	22.809	0.000
Calan.TreatmentMET:SEXSTR	2.473	1.714	1.443	0.149
1 2	1.735	0.625	2.776	0.006
2 3	2.939	0.781	3.763	0.000

Figure 9: Coefficients table for ordinal logistic regression.

The last two rows of Figure 9 provide estimates for the model's interests. For instance the row $1 \mid 2$ means for $l \le 1$ the model estimates $\alpha_1 = 1.735$ and for $l \le 2$ we can say $\alpha_2 = 2.939$. Both of these intercepts are significant as well. ("Ordinal Logistic Regression | r Data Analysis Examples" (2011)).

In summary of this model, we found a substantial difference between heifer calves from mothers on the CON treatment compared to steer calves with mothers on the DDG treatment as it relates to the easing score. The DDG steer interaction effect had an estimated positive

coefficient, but in the context of the model, this implies a decrease in the log odds. However, the MET treatment did not have a statistically significant effect.

Further output can be found in *Appendix D - Additional R Output*. From this output we can say the Residual Deviance has a value of 65.95121. This measures the goodness of fit for the model, where lower values indicate a better match between the observed and predicted data. The Akaike Information Criterion (AIC) of 79.95121 evaluates the trade-off between the model's fit and complexity, with smaller AIC values indicating a more efficient model.

Multinomial Logistic Regression (MLR)

In search of another model, we considered a multinomial logistic regression. The model predicts Calving Ease based on third trimester treatment, sex of the calf, and their interaction. The model converged successfully after several iterations, with a residual deviance of 58.418 and an AIC of 82.418. It is indicating this may not be a good choice for the final the model. The coefficients, as seen in output below, represent the log-odds of the calving ease levels relative to a reference category, but the large standard errors for interaction terms suggest potential numerical instability or sparse data for some levels.

```
# weights: 21 (12 variable)
initial value 110.959841
iter 10 value 29.432881
      20 value 29.222854
iter
iter 30 value 29.215412
iter
      40 value 29.209442
iter 50 value 29.208986
final value 29.208984
converged
Call:
multinom(formula = Calving.Ease ~ Calan.Treatment * SEX, data = data2)
Coefficients:
  (Intercept) Calan.TreatmentDDG Calan.TreatmentMET
                                                         SEXSTR
2
    -1.673984
                       -89.53191
                                          -1.159207 -0.9650507
  -21.595074
                         4.64936
                                            4.446292 1.2808117
  Calan.TreatmentDDG:SEXSTR Calan.TreatmentMET:SEXSTR
2
                  -6.615244
                                              1.313325
3
                  12.720477
                                             14.076217
Std. Errors:
  (Intercept) Calan.TreatmentDDG Calan.TreatmentMET
                                                         SEXSTR
```

2 0.6291544 6.210743e-07 1.206083 1.211298 3 604.7136553 9.531250e+02 955.622215 603.218409 Calan.TreatmentDDG:SEXSTR Calan.TreatmentMET:SEXSTR 2 7.567627e-08 1.899839

Residual Deviance: 58.41797

9.535708e+02

AIC: 82.41797

3

Another important thing is that multinomial logistic regression is designed to model nominal data, which is categorical data without an order. (Frost (2021)). We could still choose this Model for our ordinal variable if it would be better fitting than OLR Model. For these reasons, we chose to go in another direction and not recommend this model.

956.119462

Binomial Regression

As previously discussed, the data for the calving ease score is highly skewed with most calves getting a score of one. One way to reduce the effect of the skew could be to convert this to a binomial data set. In this case we would have two categories for the calving ease score: low, which corresponds to scores of one, and high, which is everything else. A type of model that fits binomial data like this is a binomial regression. While this may not make a huge difference given that we have no scores larger than three, we want to see how this model compares to the last.

For this model, we chose to use the same two explanatory variables, third trimester treatment and sex of the calf. In this case let π be the probability of a high score given the explanatory variables, X_{trt} and X_{sex} respectively. In other words $\pi = P(Y = high|X_{trt}, X_{sex})$ where Y is the same random variable as before, but is only ever low or high. Binomial regression uses π in a similar logit function as the ordinal logistic regression. Thus we can write the model as,

$$\ln\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 X_{trt} + \beta_2 X_{sex} + \beta_3 (X_{trt} \times X_{sex})$$

where β_0 is the intercept of the model and β_1 , β_2 and β_3 are the coefficients for X_{trt} , X_{sex} , and their interaction respectively. (Wiley (2013)). For this model MET is considered to be the default treatment, while steer is the default sex.

	Mo	del Fit	Statisti	ics			
Criterion Intercept Only Intercept and Covariate							
AIC			66.				
sc			82.290				
-2 Log L	(54.599			
Tes	ting Globa	al Null H	lypoth	esis:	BETA=0		
Test		Chi-S	quare	DF	Pr > ChiSq		
1.71 171	ad Datia	_	0062		0.2070		

Testing Global Null Hypothesis: BETA=0							
Test	Chi-Square	DF	Pr > ChiSq				
Likelihood Ratio	6.0963	5	0.2970				
Score	5.4692	5	0.3613				
Wald	3.0592	5	0.6909				

Joint Tests									
Effect DF Chi-Square Pr > ChiSq									
Calan.Treatment	2	2.1324	0.3443						
SEX	1	1.4189	0.2336						
Calan.Treatment*SEX	2	1.9791	0.3718						

Figure 10: Fit Statistics and other tests.

In the *Joint Tests* table above we can see the interaction terms are insignificant. This means we can look at main effects for each variable level. These can be seen in the *Analysis of Maximum Likelihood Estimates* below, but all terms outside of the intercept are insignificant as well.

Analysis of Maximum Likelihood Estimates									
Parameter			DF	Estimate	Standard Error	Wald Chi-Square	Pr > ChiSq		
Intercept			1	-1.3863	0.6455	4.6123	0.0317		
Calan.Treatment	CON		1	-1.2528	1.2199	1.0546	0.3044		
Calan.Treatment	DDG		1	-1.5581	1.2121	1.6524	0.1986		
SEX	HFR		1	-1.4469	1.2147	1.4189	0.2336		
Calan.Treatment*SEX	CON	HFR	1	2.4120	1.7154	1.9770	0.1597		
Calan.Treatment*SEX	DDG	HFR	1	-9.0820	225.2	0.0016	0.9678		

Figure 11: Analysis of Maximum Likelihood Estimates table.

Elsewhere in the output, which can be seen in *Appendix C - Additional SAS Output*, the odds ratios and subsequent confidence intervals also indicated no significant results.

While the lack of significant terms seems detrimental, the AIC of the model is 66.599, indicating a decent fit. However, we cannot compare this value to the AIC of the ordinal logistic model since they are different types of models. (Hyndman (2013)). This is backed up by the percent concordance (63.9%) and Somers' D (0.440), which suggest moderate predictive performance. Below, the odds ratio plot visually confirms the lack of significant associations for the predictors and their interactions.

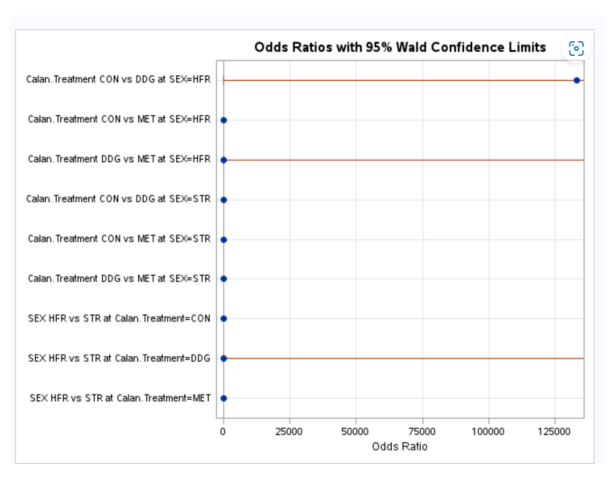


Figure 12: Plot for odds ratio.

The plots below in Figure 13 help evaluate the assumptions of a binomial regression model. The Cook's Distance plot shows that most observations have minimal influence on the model, with no Cook's distances exceeding common thresholds (e.g., 0.5), indicating no highly influential data points. The Deviance Residuals plot demonstrates that most residuals are centered near zero, reflecting a generally good model fit, but a few residuals exceed ± 2 , which may indicate potential outliers or areas of poor model performance.

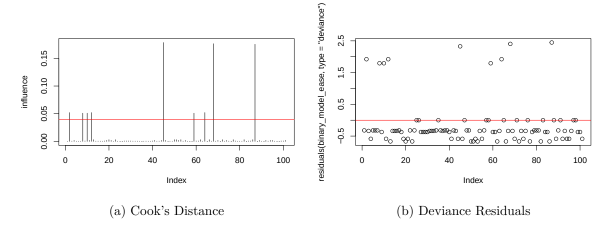


Figure 13: Graphs to test assumptions.

The Variance Inflation Factor (VIF) analysis highlights severe multicollinearity for Calan.Treatment (adjusted GVIF = 50.03) and its interaction with SEX (adjusted GVIF = 58.34), suggesting strong redundancy among these predictors. This appears to be a direct contradiction with Figure 6. Due to time constraints we were not able to invesigate this further though.

In contrast, the adjusted GVIF for SEX (1.68) indicates no significant multicollinearity for this variable alone. Overall, while the model does not appear to have issues with influential points, the presence of multicollinearity among predictors could complicate interpretation and may necessitate further investigation or variable transformation.

Variable	GVIF	Df	$GVIF^{(1/(2*Df))}$
Calan.Treatment	6.266542e + 06	2	50.033051
SEX	2.813087e + 00	1	1.677226
Calan.Treatment:SEX	1.158065e + 07	2	58.335540

Figure 14: VIF table

Conclusion for Calving Ease Score

We previously discussed three models for the calving ease score. As we already indicated, the multinomial logistic regression is not the direction we would recommend. Regarding to the other two, model preference depends on the cleint's desired goal. If these models are intended for purely predictive purposes, the binomial regression model may be effective. However, if

the client wants to study the impact the third-trimester treatment and the sex of the calf had on the ease score, then we strongly recommend the ordinal logistic regression model over the other models discussed. Additionally the multicolinearity assumption was not found in the ordinal logistic regression model, but further invesitgration should be done.

Models for Calf Vigor Score

Ordinal Logistic Regression Model

The similarities between the calving ease and calf vigor score make identifying potential models a much easier task, as what might fit calving ease score can also potentially fit the calf vigor score. We will first look at an ordinal logistic regression model. The model can once again be represented by the equation

$$\ln \left(\frac{P(Y \leq l)}{P(Y > l)} \right) = \alpha_l - \beta_{trt} X_{trt} - \beta_{sex} X_{sex} - \beta_{trt:sex} (X_{trt} \times X_{sex})$$

since the client is interested in the effect of the same explanatory variables on the score. This equation looks exactly the same as before, with the only changes being Y now representing the calf vigor score and l now ranging from one to eleven in real life. (Probo (2022)). Note, within out cleaned data set, l=1,2,3 as it did for the ease score.

While the dependent variable and equation are very similar to what we previously did the results of the model are not. In Figure 15 we can see the interaction terms are non-significant. The same can be said for the main effects meaning only the intercepts, α_l , are significant. This indicates neither of the explanatory variables seems to have much of an effect on the vigor score.

	Value	Std. Error	t value	p-value
Calan.TreatmentDDG	-0.021	0.852	-0.025	0.980
Calan.TreatmentMET	-0.179	0.845	-0.211	0.833
SEXSTR	-1.271	1.177	-1.080	0.280
Calan.TreatmentDDG:SEXSTR	1.229	1.452	0.847	0.397
Calan.TreatmentMET:SEXSTR	0.834	1.535	0.543	0.587
1 2	1.332	0.559	2.381	0.017
2 3	2.068	0.601	3.440	0.001

Figure 15: Coefficients table for ordinal logistic regression.

Additionally, the model returned an AIC of 127.059. This is not bad on its own and indicates the model fits decently. However, the issues regarding significance make us skeptical of the model's predictive capability. It may indicate that the two explanatory variables just do not significantly impact the vigor score. We will want to test other models before arriving at that conclusion, though. Note, the full model output can be found in $Appendix\ D$ - $Additional\ R$ Output.

Binomial Regression

Given the lack of significance with the ordinal logistic regression model, we will look into the binomial example again. As before, we are condensing the vigor score into a low category and a high category. All of the scores of one fit in the former and the rest are in the latter. Just as with the calving ease score, let π be the probability of a high score. Then, by using the same two explanatory variables again, we can write the model as

$$\ln\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 X_{trt} + \beta_2 X_{sex} + \beta_3 (X_{trt} \times X_{sex})$$

Once again, the Analysis of Maximum Likelihood Estimates table below indicates no significant interactions or main effects, outside of the intercept. Again though, the Model Fit Statistics table has a decent AIC store, so the model may still have predictive qualities despite showing no significant relationships among our explanatory variables.

	Mod	del Fit	Statisti	ics		
Criterion	Intercept	Interd	cept a	nd (Covariates	
AIC	9	3.548				101.469
SC	9	6.163				117.160
-2 Log L	9	1.548				89.469
						-
Tes	ting Globa	l Null H	lypoth	esis: I	BET	A=0
Test		Chi-Square		DF	Pr	> ChiSq
Likeliho	2.0784		5		0.8382	
Score		1.	8352	5		0.8714
Wald	1.	6922	5		0.8899	
		Joint 1	Tests			
Effect	DF	Wald Chi-Square		-	Pr > ChiSq	
Calan.Trea	2	1.1835		5	0.5534	
	1	0.0705		5	0.7907	
SEX				0.7551		000.

Figure 16: Fit Statistics and other tests

Analysis of Maximum Likelihood Estimates									
Parameter			DF	Estimate	Standard Error	Wald Chi-Square	Pr > ChiSq		
Intercept			1	1.8718	0.7596	6.0730	0.0137		
Calan.Treatment	CON		1	0.7666	1.2837	0.3567	0.5504		
Calan.Treatment	DDG		1	-0.4855	0.9431	0.2650	0.6067		
SEX	HFR		1	-0.2624	0.9884	0.0705	0.7907		
Calan.Treatment*SEX	CON	HFR	1	-1.0543	1.5377	0.4701	0.4929		
Calan.Treatment*SEX	DDG	HFR	1	0.1754	1.3091	0.0179	0.8934		

Figure 17: Analysis of Maximum Likelihood Estimates table.

More of the SAS output, including odds ratios and plot can be found in $Appendix\ C$ - $Additional\ SAS\ Output.$

As with the previous binomial regression model, we need to check the assumptions of the binomial regression. The adjusted GVIF values in Figure 18 for all predictors and their interaction are below 5, implying no significant multicollinearity in the model.

Variable	GVIF	Df	$GVIF^(1/(2*Df))$
Calan.Treatment	2.986258	2	1.314564
SEX	4.701880	1	2.168382
Calan.Treatment:SEX	11.078455	2	1.824399

Figure 18: VIF table

In Figure 19 we can see the Cook's Distance plot indicates that all observations have minimal influence on the model, with no notable outliers or highly influential points. The Deviance Residuals plot shows residuals tightly clustered around zero, suggesting an excellent model fit, though the lack of variability might indicate overfitting or an overly simplistic model specification. Overall, the diagnostics support the validity of the model, but the small deviance residuals may require further investigation to ensure appropriate model fit.

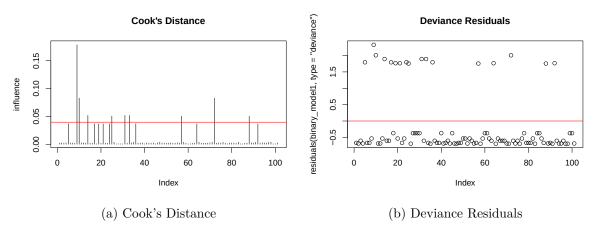


Figure 19: Graphs to test assumptions.

Conclusion for Calf Vigor Score

The two models discussed above both had a similar conclusion, which is the vigor score was not significantly affected by either the third trimester treatment, the calf's sex, or their interaction. Therefore recommending a model for this dataset will again depend on Client's goal. However, both models could still have some predictive value, which could be evaluated at a later date.

Note that we did fit a multinomial logistic regression model for the vigor score as well, but we ran into the same issues as before so no results are shown for the model in this paper.

Models for Final Calf Body Weight

Mixed Model

Now, we will look at models for investigating the effect the client's chosen explanatory variables have on the final body weight of the calf. Since the body weight is a quantitative variable, we can turn to linear models. We initially attempted to fit a simple model using only the third-trimester treatment and sex of the calf but ultimately chose to include the calf's sire as an additional explanatory variable. This slightly complicates the model as we believe the additional variable is better represented as a random effect, meaning we are working with a mixed model.

The proposed model and be represented as

$$Y_{ijk} = \mu + \alpha_i + \beta_j + (\alpha\beta)_{ij} + u_k + e_{ijk}$$

where Y_{ijk} represents the final weight for the *ith* treatment, the *j*th sex, and the *k*th sire. μ represents the overall mean, while α_i is the fixed effect for the *ith* treatment, β_j is the fixed effect for the *j*th sex, and $(\alpha\beta)_{ij}$ is their interaction. The random sire effect is represented by u_k and we assume u_k $N(0, \sigma_k^2)$. We also assume the residual term e_{ijkl} is distributed as $N(0, \sigma^2)$.

For a linear mixed model to work, the assumption regarding the distribution of the residuals must hold. These can be checked graphically in the conditional residual plots below. The histogram and Q-Q plot show if the residuals are approximately normal, while the plot in the top left evaluates any multicollinearity concerns. Thankfully the graphs give us no concerns as no trends are present in the top-left plot and the residuals are randomly distributed around zero, the histogram appears bell shaped around zero, and the Q-Q plot sees most points fall along the line, which is ideal. Therefore we can procede with our mixed model.

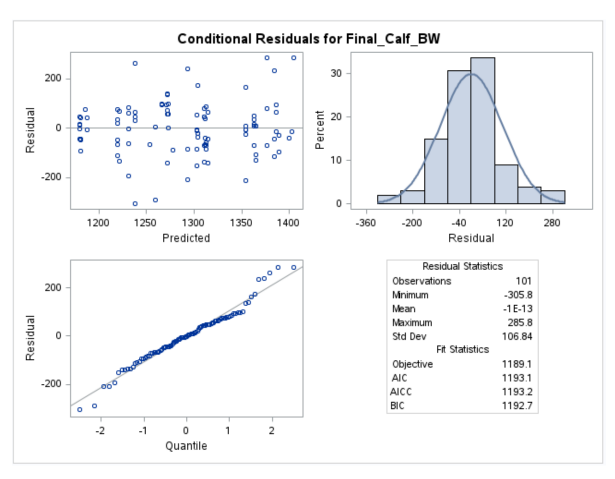


Figure 20: Plots to check residual assumption.

The *Fit Statistics* table below shows the model has an AIC of 1193.1. While this is not great, it is not far off from other models we have viewed in other data analyses. Also, the sire effect on the covariance structure is shown in the *Covariance Parameter Estimates* table.

		Co	nvergence	criteria	met.				
		Covaria	Covariance Parameter Estimates						
		Cov Pa	rm Subj	ect	Estim	ate			
		Interce	ot Sire		2372	2.00			
		Residu	al		12	484			
			Fit Stati	istics					
		-2 Res	Log Likelih	nood	118	9.1			
		AIC (Sn	naller is Be	etter)	119	3.1			
		AICC (S	maller is l	Better)	119	3.2			
		BIC (Sn	naller is Be	etter)	119	2.7			
1									
			3 Tests of						
	Effect		Num DF			F Value	Pr > F		
		Treatment	2		90	0.41	0.6660		
	SEX		1		90	25.16 0.23	<.0001		
	Calan.	Treatment*SEX	2	2 90			0.7919)	
						0.23	0.7010		
		1.	aget Squar	ae Mas	ne	0.23	0.7010		
		L	east Squar	es Mea	ans				
Effect		Lo Calan.Treatme		es Mea		Standard Error		t Value	Pr > t
Effect Calan.Treatmen	t*SEX				nate	Standard	DF	t Value 38.37	
		Calan.Treatme	nt SEX	Estim	nate 0.83	Standard Error	DF 90		<.0001
Calan.Treatmen	t*SEX	Calan.Treatme	nt SEX HFR	1270 1363	nate 0.83	Standard Error 33.1186	DF 90 90	38.37	<.0001
Calan.Treatmen	t*SEX t*SEX	Calan.Treatme	nt SEX HFR STR	1270 1363 1236	nate 0.83 3.88	Standard Error 33.1186 35.4134	DF 90 90 90	38.37 38.51	Pr > t <.0001 <.0001 <.0001 <.0001
Calan.Treatmen Calan.Treatmen Calan.Treatmen	t*SEX t*SEX t*SEX	Calan.Treatme CON CON DDG	nt SEX HFR STR HFR	1270 1363 1236 1353	0.83 3.88 6.86	Standard Error 33.1186 35.4134 36.2887	DF 90 90 90 90	38.37 38.51 34.08	<.0001 <.0001 <.0001

Figure 21: Covariance parameter, fit statistics, fixed effects, and least squares mean tables

The Type 3 Fixed Effects table tells us what fixed effects, if any, are statistically significant. As we have done when viewing other models in this paper, we must first look at the interaction effect, which is insignificant. Therefore, we turn to the main effects of the two explanatory variables. While the sex of the calf is highly significant (p-value < 0.0001), the treatment

effect is insignificant. Given that the interaction is insignificant, we chose not to evaluate the Differences of the Least Squares Means table, but we did include it in Appendix C - Additional SAS Output for the transparency.

ANCOVA Model

Another model that could fit the data is an ANCOVA model. These models take a continuous dependent variable and take at least one categorical variable along with a covariate. The dependent and categorical variables will once again be the third-trimester treatment and sex of the calf. The covariate must be a continuous, independent variable that is not a primary interest of the study. (Frost (2023)). We previously looked at the mother heifer's initial body weight in Figure 8, so we will make that the covariate.

The proposed ANCOVA model can be represented as:

$$Y_{ijk} = \mu + \alpha_i + \beta_j + (\alpha\beta)_{ij} + \gamma X_k + e_{ijk}$$

where Y_{ijk} represents the final calf body weight for the ith treatment, the jth sex, and the kth initial.BW. μ represents the overall mean, while α_i is the fixed effect for the ith treatment, β_j is the fixed effect for the jth sex, and $(\alpha\beta)_{ij}$ is their interaction. X_k is covariate effect of initial body weight where γ is the slope. We assume the residual error term e_{ijkl} is distributed as $N(0,\sigma^2)$.

This ANCOVA model adjusts the response variable Yijk (final calf body weight) for the covariate initial body weight X_k while modeling the main effects of treatment (α_i) , sex (β_j) , and their interaction $((\alpha\beta)_{ij})$. The residual errors are assumed to be normally distributed with constant variance.

Now, we can view the output of this model after it was fit in SAS. The ANCOVA model for the final body weight (Final_Calf_BW) is significant overall, with an F-value of 5.72 and a p-value < 0.0001, indicating that the predictors collectively explain a significant portion of the variability in final calf body weight. The R-Square value of 0.267 shows that 26.7% of the variability in Final_Calf_BW is accounted for by the model.

The GLM Procedure Dependent Variable: Final Calf BW Source DF Sum of Squares Mean Square F Value Pr > FModel 6 444984.875 74164.146 5.72 <.0001 Error 94 1218813.659 12966.103 Corrected Total 1663798.535 100 R-Square Coeff Var Root MSE Final_Calf_BW Mean 0.267451 8.816212 113.8688 1291.584 Source DF Type I SS Mean Square F Value Pr > FCalan.Treatment 2 13661.9170 6830.9585 0.53 0.5922 SEX 294400.0151 294400.0151 22.71 <.0001 1 2 8623.2008 0.33 0.7179 Calan.Treatment*SEX 4311.6004 Initial BW 128299.7422 128299.7422 9.90 0.0022 1 Source DF Type III SS Mean Square F Value Pr > F Calan.Treatment 2 8383.2348 4191.6174 0.32 0.7246 SEX 284570.2895 284570.2895 21.95 <.0001 1 Calan.Treatment*SEX 2 16304.4068 8152.2034 0.63 0.5355 128299.7422 Initial BW 128299.7422 9.90 0.0022 1

Figure 22: Model information along with Type I and Type III test results.

The final table in the figure below shows the Type 3 sums of squares. Here we can see the treatment by sex interaction is insignificant. In terms of main effects, sex (p-value < 0.0001) and the initial body weight of the mother (p-value = 0.0022) are significant, but the treatment is insignificant. This emphasizes that the sex of the calf and mother's initial body weight are key variables driving differences in the final weight of the calf, while treatment appears negligible.

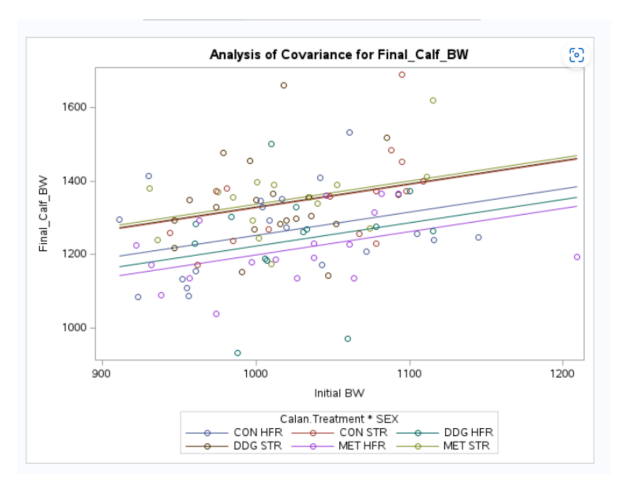


Figure 23: ANCOVA Plot-1

This ANCOVA plot examines the relationship between Initial Body Weight (Initial BW) and Final Calf Body Weight (Final_Calf_BW) for combinations of the treatment (Calan.Treatment) and SEX. Each line represents a fitted regression line, showing how Initial BW influences Final_Calf_BW for each group. The generally parallel nature of the lines indicates no significant interaction between Calan.Treatment and SEX, as the slopes are similar across groups. The positive slopes demonstrate that Initial BW is a strong predictor of Final_Calf_BW, with higher initial weights leading to higher final weights. There is some variability within each group, as shown by the scatter of points around the lines, but overall, Initial BW appears to adjust group differences effectively.

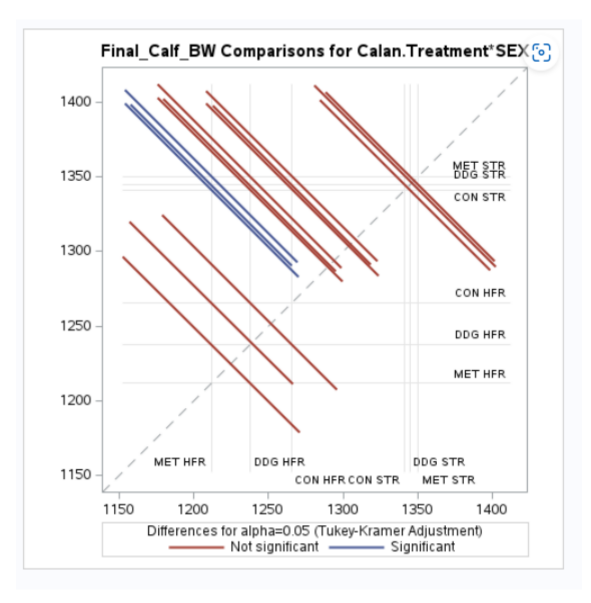


Figure 24: Tukey-Kramer adjustments plot

The plot displays pairwise comparisons of Final_Calf_BW across treatment and sex combinations, using Tukey-Kramer adjustments for multiple comparisons. Red lines represent comparisons where the differences between groups are not statistically significant, while blue lines indicate significant differences at the 0.05 level. The majority of comparisons are not significant, as evidenced by the predominance of red lines. However, some significant differences are noted, such as between MET STR and CON HFR. Overall, the visualization demonstrates that while certain treatment-sex combinations show significant differences, the interaction effect between treatment and sex does not exhibit a strong pattern of significance, supporting

the earlier conclusion of non-significant interaction effects.

We also need to check the assumptions of the ANCOVA model. This scatter plot below of residuals against predicted values checks for homoscedasticity (constant variance of residuals). The residuals are scattered relatively evenly around the zero line, showing no clear pattern or trend, which supports the assumption of homoscedasticity.

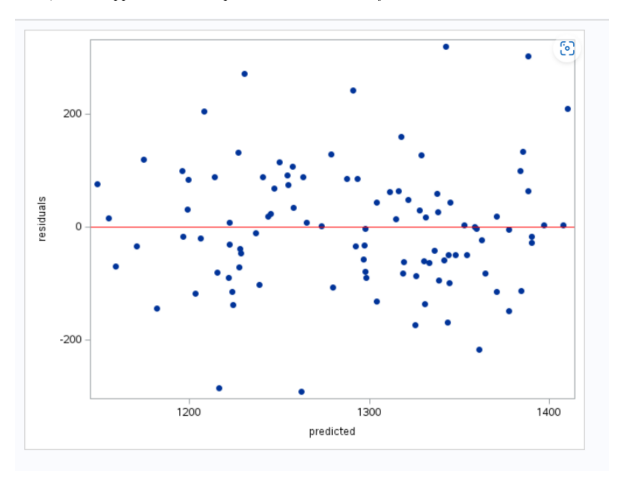


Figure 25: Residuals vs Predicted Plot

The histogram shows the distribution of residuals, which appears approximately symmetric and bell-shaped. The overlaid normal curve aligns well with the histogram, further supporting the assumption of normality.

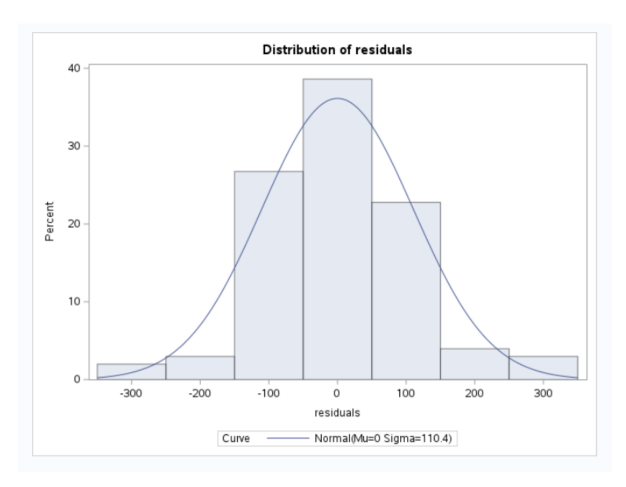


Figure 26: Histogram

For the Q-Q plot, most points lie close to the diagonal line, indicating normality. However, slight deviations at the tails suggest minor departures from normality, but these are not severe enough to invalidate the model assumptions.

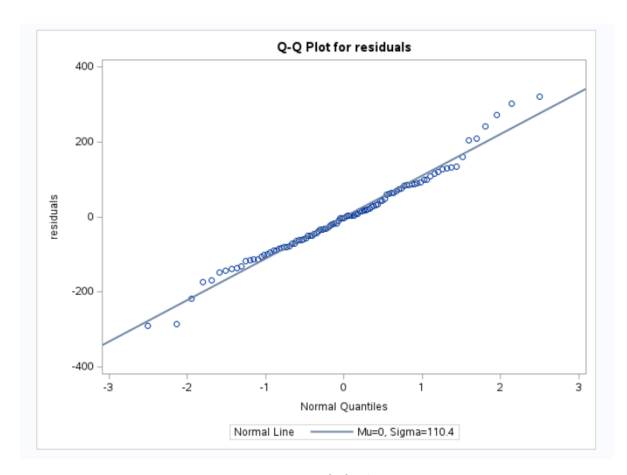


Figure 27: Q-Q Plot

The diagnostic plots and tests collectively support that the assumptions of homoscedasticity, normality, and zero-mean residuals are reasonably met. While minor deviations are observed, they are unlikely to significantly impact the validity of the model.

Conclusion for Final Calf Body Weight

From the results of both models we can say the sex of the calf had a significant impact on the final body weight of the calf. There was not a significant result for the third trimester treatment effects though. This indicates the treatments did not have a meaningful effect on the final body weight. Between the two models, we found the the ANCOVA model offered clear and interpretable results, making it our recommended approach for future studies.

It is important to note that in the ANCOVA model, the initial body weight of the mother was deemed significant. In the future including more models that account for this variable

and others within the data set may give cattle producers a better idea of how to maximize the weight of their calves before slaughter.

Future Work

This dataset contained many missing values. We ended up performing our analysis on only 84.17% of the original data. Repeating our analysis methods with imputed data would allow for more of the data to be used and may lead to slightly different results. While we previously explained why we did not impute this dataset, seeing if the results of the models differed would be interesting.

Additionally, future analysis of this data should attempt to control for more variables in the models. In our final model, the ANCOVA model for the final body weight of the calves, we found information about the mother heifer, her body weight, had a significant impact on the model. We were unable to apply this variable and others to additional models due to time constraints, but would recommend this to anyone working on this data in the future.

References

- Frost, Jim. 2021. "Multinomial Logistic Regression: Overview & Example." Statistics By Jim. 2021. https://statisticsbyjim.com/regression/multinomial-logistic-regression/.
- ——. 2023. "ANCOVA: Uses, Assumptions & Example." Statistics By Jim. 2023. https://statisticsbyjim.com/anova/ancova/#comments.
- Heins, Glenda, Brad & Pereira. 2023. "Monitoring Calving Traits to Improve Cow and Calf Health." University of Minnesota Extension. 2023. https://extension.umn.edu/dairy-milking-cows/calving-traits#:~:text=Calving%20ease%20score%201%3A%20quick, score%204%3A%20used%20obstetrical%20chains.
- Hyndman, Ron J. 2013. "Facts and Fallacies of the AIC." 2013. https://robjhyndman.com/hyndsight/aic/.
- Memon, Shaheen MZ., Robert Wamala, and Ignace H. Kabano. 2023. "A Comparison of Imputation Methods for Categorical Data." *Informatics in Medicine Unlocked* 42: 101382. https://doi.org/https://doi.org/10.1016/j.imu.2023.101382.
- "Ordinal Logistic Regression | r Data Analysis Examples." 2011. UCLA. 2011. https://stats.oarc.ucla.edu/r/dae/ordinal-logistic-regression/.
- "Ordinal Regression." n.d. University of St. Andrews. n.d. https://www.st-andrews.ac.uk/media/ceed/students/mathssupport/ordinal%20logistic%20regression.pdf.
- Probo, Maria Cristina, Monica & Veronesi. 2022. "Clinical Scoring Systems in the Newborn Calf: An Overview." *Animals (Basel)*. https://doi.org/doi.10.3390/ani12213013.
- Saner, Brianna, Randy & Buseman. 2024. "How Many Pounds of Meat Can We Expect from a Beef Animal?" 2024. https://beef.unl.edu/beefwatch/2020/how-many-pounds-meat-can-we-expect-beef-animal.
- USDA. "5017-1: Calculating Dry Matter Intake from Pasture." https://www.ams.usda.gov/rules-regulations/organic/handbook/5017-1#:~:text=DMI%20is%20the%20level%20of, life%20and%20level%20of%20production.
- William, Nkugwa Mark. 2023. "How to Determine Bin Width for a Histogram (r and Python)." 2023. https://nkugwamarkwilliam.medium.com/how-to-determine-bin-width-for-a-histogram-r-and-pyth-653598ab0d1c.

Appendix A - R Code

```
library(knitr)
library(dplyr)
library(ggplot2)
library(naniar)
library(reshape2)
library(GGally)
library(janitor)
library(emmeans)
library(MASS)
library(multcomp)
library(lme4)
library(nnet)
data <- read.csv("data.csv")</pre>
# Placeholder of original data
origdata <- data
data <- data %>%
  mutate(across(everything(), ~ ifelse(. %in% c(".", ""), NA, .)))
length(unique(na.omit(data$Sire)))
length(unique(na.omit(data$Development.Treatment)))
sum(rowSums(is.na(data)) > 0)
rows_with_na <- sum(rowSums(is.na(data)) > 0)
unique_entries <- unique(data$Calan.Treatment)</pre>
num_unique <- length(unique_entries)</pre>
frequency_table <- table(data$Calan.Treatment)</pre>
gg_miss_var(data) +
  ggtitle("Missing Data Distribution") +
  theme_minimal()
var_used <- c("Calan.Treatment", "SEX", "Calving.Ease",</pre>
               "Calf.Vigor", "Final.Calf.BW", "Pen..", "Initial.BW", "Sire")
cleaned_data <- data[complete.cases(data[, var_used]), ]</pre>
dep_vars <- c("Calving.Ease", "Calf.Vigor", "Final.Calf.BW")</pre>
custom_names <- c("Calving Ease", "Calf Vigor", "Final Calf Weight")</pre>
```

```
cleaned_data$Calving.Ease <- as.numeric(cleaned_data$Calving.Ease)</pre>
cleaned_data$Calf.Vigor <- as.numeric(cleaned_data$Calf.Vigor)</pre>
cleaned_data$Final.Calf.BW <- as.numeric(cleaned_data$Final.Calf.BW)</pre>
calc stats <- function(var) {</pre>
  mean_val <- mean(var, na.rm = TRUE)</pre>
  median_val <- median(var, na.rm = TRUE)</pre>
  sd_val <- sd(var, na.rm = TRUE)</pre>
  quantiles <- quantile(var, probs = c(0.25, 0.75), na.rm = TRUE)
  max <- max(var, na.rm = TRUE)</pre>
  min <- min(var, na.rm = TRUE)</pre>
  c(mean = mean_val, median = median_val, sd = sd_val,
    Q1 = quantiles[1], Q3 = quantiles[2], Min = min, Max = max)
summary_table <- t(sapply(dep_vars, function(var)</pre>
  calc_stats(cleaned_data[[var]])))
summary_table <- as.data.frame(summary_table)</pre>
kable(summary_table, col.names = c("Variable", "Mean",
                                     "Median", "SD", "25th Percentile",
                                     "75th Percentile", "Min", "Max"),
      caption = "Summary Statistics for Dependent Variables")
bin_width <- round(2*IQR(cleaned_data$Final.Calf.BW)/</pre>
                      (length(cleaned_data$Final.Calf.BW))^(1/3), 2)
ggplot(cleaned_data, aes(x = Final.Calf.BW)) +
  geom histogram(binwidth = bin_width, color = "black", fill = "skyblue") +
  labs(
    x = "Weight",
    y = "Frequency"
  ) +
  theme_minimal()
ggplot(cleaned_data, aes(sample = Final.Calf.BW)) +
  stat_qq() +
  stat_qq_line(color = "red") +
  labs(
    x = "Theoretical Quantiles",
    y = "Sample Quantiles"
```

```
) +
  theme_minimal()
ggplot(cleaned_data, aes(x = Calving.Ease)) +
  geom_histogram(binwidth = 1, color = "black", fill = "skyblue") +
  labs(
   x = "Ease Score",
    y = "Frequency"
  )+
  theme_minimal()
ggplot(cleaned_data, aes(x = Calf.Vigor)) +
  geom_histogram(binwidth = 1, color = "black", fill = "skyblue") +
  labs(
    x = "Vigor Score",
    y = "Frequency"
  ) +
  theme_minimal()
data2 <- cleaned_data
crosstab_treatment_sex <- table(data2$Calan.Treatment, data2$SEX)</pre>
kable(
  crosstab_treatment_sex)
chi_sq_test <- chisq.test(crosstab_treatment_sex)</pre>
chi_sq_table <- data.frame(</pre>
  Metric = c("Statistic", "Degrees of Freedom", "P-Value"),
  Value = c(round(chi_sq_test$statistic, 2),
            chi_sq_test$parameter,
            format.pval(chi_sq_test$p.value)),
 stringsAsFactors = FALSE
)
kable(chi_sq_table, align = c("l", "c"), row.names = FALSE)
ggplot(data2, aes(x = Calan.Treatment, y = Final.Calf.BW, fill = SEX)) +
  geom_boxplot(outlier.color = "red", alpha = 0.7) +
  labs(title = "Boxplot of Final Calf Body Weight by Treatment and Sex",
       x = "Treatment", y = "Final Body Weight") +
  theme_minimal()
```

```
data2$Initial.BW <- as.numeric(data2$Initial.BW)</pre>
ggplot(data2, aes(x = Calving.Ease, y = Initial.BW,
                   color = SEX, shape = Calan.Treatment)) +
  geom point(size = 3, alpha = 0.7) +
  labs(x = "Calving Ease", y = "Initial Body Weight of Heifer") +
  theme minimal()
ggplot(data2, aes(x = Calf. Vigor, y = Initial. BW,
                   color = SEX, shape = Calan.Treatment)) +
  geom_point(size = 3, alpha = 0.7) +
  labs(x = "Calf Vigor", y = "Initial Body Weight of Heifer") +
  theme_minimal()
data2$Calving.Ease <- factor(data2$Calving.Ease, ordered = TRUE)
ease_model_3 <- polr(Calving.Ease ~ Calan.Treatment * SEX,</pre>
                      data = data2, Hess = TRUE)
coefs <- coef(summary(ease model 3))</pre>
p_values <- pnorm(abs(coefs[, "t value"]), lower.tail = FALSE) * 2</pre>
coefs <- cbind(coefs, "p-value" = p_values)</pre>
kable(coefs,
      digits = 3,
      format = "markdown")
data2$Calf.Vigor <- factor(data2$Calf.Vigor, ordered = TRUE)</pre>
vigor_model_1 <- polr(Calf.Vigor ~ Calan.Treatment * SEX,</pre>
                       data = data2, Hess = TRUE)
summary(vigor_model_1)
coefs <- coef(summary(vigor_model_1))</pre>
p values <- pnorm(abs(coefs[, "t value"]), lower.tail = FALSE) * 2</pre>
coefs <- cbind(coefs, "p-value" = p_values)</pre>
#print(coefs)
kable(coefs,
      digits = 3,
      format = "markdown")
```

Appendix B - SAS Code

```
/* Binomial Calving_Ease Model */
data data;
    set data;
    if 'Calving.Ease'n in (2, 3) then Binary_Ease = "High";
    else if 'Calving.Ease'n = 1 then Binary_Ease = "Low";
run;
proc freq data=data;
    tables Binary_Ease;
run;
/* Logistic regression model */
proc logistic data=data;
    class 'Calan.Treatment'n SEX / param=ref;
    model Binary_Ease(event='High') = 'Calan.Treatment'n|SEX;
    oddsratio 'Calan.Treatment'n;
    oddsratio SEX;
run;
```

```
/*Binomial Calf Vigor*/
data data;
    set data;
    /* Recode Calf. Vigor: 1 = Low, 2 and 3 = High */
    if 'Calf.Vigor'n = 1 then Binary_Vigor = "Low";
    else if 'Calf.Vigor'n in (2, 3) then Binary_Vigor = "High";
run;
proc freq data=data;
   tables Binary_Vigor;
run;
data data;
    set data;
    Binary_Vigor = strip(Binary_Vigor);
run;
data data;
    set data;
    Binary_Vigor = propcase(Binary_Vigor);
run;
```

```
proc logistic data=data descending;
    class 'Calan.Treatment'n SEX / param=ref;
    model Binary_Vigor = 'Calan.Treatment'n|SEX;
    oddsratio 'Calan.Treatment'n;
    oddsratio SEX;
run;
/*Mixed Model*/
data data;
    set data;
    if not missing('Final.Calf.BW'n) then Final_Calf_BW =
    input('Final.Calf.BW'n, best12.);
run;
proc mixed data=data method=reml plots=residualpanel;
    class 'Calan.Treatment'n SEX Sire;
    model Final_Calf_BW = 'Calan.Treatment'n|SEX ;
    random intercept / subject=Sire;
    lsmeans 'Calan.Treatment'n*SEX / adjust=tukey pdiff;
run;
/*ANCOVA*/
proc glm data=data ;
    class 'Calan.Treatment'n SEX;
    model Final_Calf_BW = 'Calan.Treatment'n|SEX 'Initial BW'n ;
    means 'Calan.Treatment'n / tukey cldiff;
    lsmeans 'Calan.Treatment'n*SEX / adjust=tukey pdiff cl;
    output out=diagnostics r=residuals p=predicted;
run;
/* Diagnostic plots */
proc sgplot data=diagnostics;
    scatter x=predicted y=residuals / markerattrs=(symbol=circlefilled);
    refline 0 / axis=y lineattrs=(color=red);
run;
proc univariate data=diagnostics normal;
    var residuals;
```

histogram residuals / normal;

run;

qqplot residuals / normal(mu=est sigma=est);

Appendix C - Additional SAS Output

Binomial Regression Model for Calving Ease

SEX HFR vs STR at Calan.Treatment=DDG

SEX HFR vs STR at Calan.Treatment=MET

	Association of Predicted Pro	babilitie	s and Obse	rved Re	esponses		
	Percent Concordant	63.9	Somers'	D	0.440		
	Percent Discordant	19.9	Gamma		0.524		
	Percent Tied	16.2	Tau-a		0.072		
	Pairs	828	С		0.720		
	Odds Ratio Estimates a	and Wald	d Confidence	e Interv	als		
Odds	Ratio		Estimate	95% C	onfidence	Limi	
Calan.	Treatment CON vs DDG at SEX	(=HFR	>999.999	<0.0	01 >9	99.99	
Calan.	Treatment CON vs MET at SEX	=HFR	3.188	0.3	00	33.89	
Calan.Treatment DDG vs MET at SEX=HFR			<0.001	<0.0	01 >9	>999.99	
Calan.Treatment CON vs DDG at SEX=STR			1.357	0.0	78	23.61	
Calan.	Treatment CON vs MET at SEX	=STR	0.286	0.0	26	3.12	
Calan	Treatment DDG vs MET at SEX	=STR	0.211	0.0	20	2.26	
SEX H	IFR vs STR at Calan.Treatment	=CON	2.625	0.2	44	28.19	

Figure 28: Association of Predicted Probabilities and Observed Responses and Odds Ratio Estimates and Wald Confidence Intervals tables.

< 0.001

0.235

< 0.001

0.022

>999.999

2.544

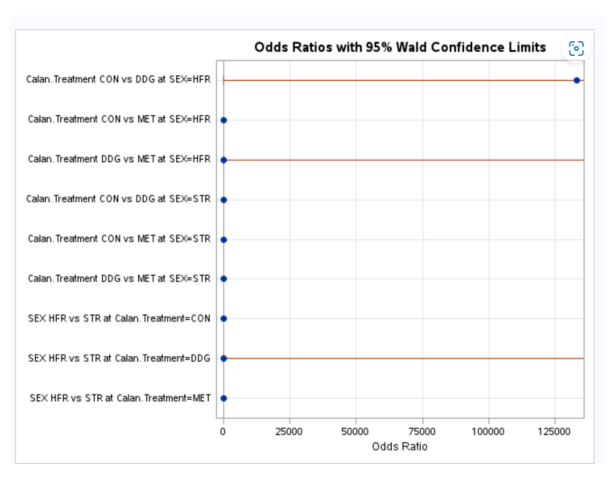


Figure 29: Odds Ratios with 95% Wald Confidence Limits table.

Binomial Regression Model for Calf Vigor

	Mod	del Fit 9	Statisti	ics			
Criterion	Intercept	Only	Interd	tercept and Covariates			
AIC	9	3.548			101.46	9	
sc	9	6.163			117.16	0	
-2 Log L	9	1.548			89.46	9	
Tes	ting Globa	Null H	ypoth	esis: I	BETA=0		
Test		Chi-Sq	uare	DF	Pr > ChiSq		
Likeliho	od Ratio	2.0784		5	0.8382		
Score		1.8352		5	0.8714		
Wald		1.	6922	5	0.8899		
		Joint 1	ests				
				Wal	-		
Effect		DF	Chi-Square		e Pr > ChiS	q	
Calan.Trea	tment	2		1.183	5 0.553	4	
SEX		1		0.070	5 0.790	7	
alan Trea	tment*SEX	2		0.755	1 0.685	6	

Figure 30: $Fit\ Statistics$ and other tests.

Association of Predicted Probabilities and Observed Responses							
Percent Concordant	51.0	Somers' D	0.189				
Percent Discordant	32.1	Gamma	0.228				
Percent Tied	16.9	Tau-a	0.053				
Pairs	1428	С	0.595				

Odds Ratio Estimates and Wald Confidence Intervals							
Odds Ratio	Estimate	95% Confidence Lim					
Calan.Treatment CON vs DDG at SEX=HFR	1.023	0.189	5.526				
Calan.Treatment CON vs MET at SEX=HFR	0.750	0.143	3.941				
Calan.Treatment DDG vs MET at SEX=HFR	0.733	0.124	4.346				
Calan.Treatment CON vs DDG at SEX=STR	3.498	0.349	35.071				
Calan.Treatment CON vs MET at SEX=STR	2.153	0.174	26.644				
Calan.Treatment DDG vs MET at SEX=STR	0.615	0.097	3.908				
SEX HFR vs STR at Calan.Treatment=CON	0.268	0.027	2.697				
SEX HFR vs STR at Calan.Treatment=DDG	0.917	0.170	4.930				
SEX HFR vs STR at Calan.Treatment=MET	0.769	0.111	5.338				

Figure 31: Association of Predicted Probabilities and Observed Responses and Odds Ratio Estimates and Wald Confidence Intervals tables.

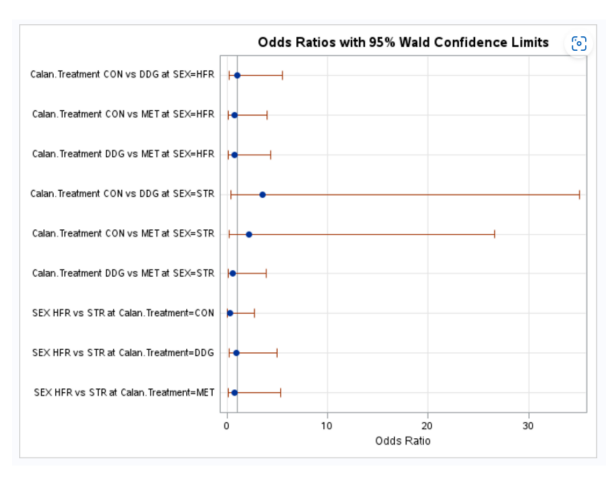


Figure 32: Odds Ratios with 95% Wald Confidence Limits table.

Linear Mixed Model for Calf Final Body Weight

			Differences of	of Least	Squares Me	ans					
Effect	Calan.Treatment	SEX	_Calan.Treatment	_SEX	Estimate	Standard Error	DF	t Value	Pr > t	Adjustment	Adj P
Calan.Treatment*SEX	CON	HFR	CON	STR	-93.0477	38.9261	90	-2.39	0.0189	Tukey-Kramer	0.1707
Calan.Treatment*SEX	CON	HFR	DDG	HFR	33.9699	40.4261	90	0.84	0.4030	Tukey-Kramer	0.9591
Calan.Treatment*SEX	CON	HFR	DDG	STR	-82.8198	36.4148	90	-2.27	0.0253	Tukey-Kramer	0.2155
Calan.Treatment*SEX	CON	HFR	MET	HFR	40.0888	37.2865	90	1.08	0.2852	Tukey-Kramer	0.8900
Calan.Treatment*SEX	CON	HFR	MET	STR	-90.7112	38.7579	90	-2.34	0.0215	Tukey-Kramer	0.1890
Calan.Treatment*SEX	CON	STR	DDG	HFR	127.02	41.9550	90	3.03	0.0032	Tukey-Kramer	0.0367
Calan.Treatment*SEX	CON	STR	DDG	STR	10.2279	38.2801	90	0.27	0.7899	Tukey-Kramer	0.9998
Calan.Treatment*SEX	CON	STR	MET	HFR	133.14	39.2920	90	3.39	0.0010	Tukey-Kramer	0.0130
Calan.Treatment*SEX	CON	STR	MET	STR	2.3365	41.1044	90	0.06	0.9548	Tukey-Kramer	1.0000
Calan.Treatment*SEX	DDG	HFR	DDG	STR	-116.79	39.3930	90	-2.96	0.0039	Tukey-Kramer	0.0435
Calan.Treatment*SEX	DDG	HFR	MET	HFR	6.1189	40.3190	90	0.15	0.8797	Tukey-Kramer	1.0000
Calan.Treatment*SEX	DDG	HFR	MET	STR	-124.68	42.5172	90	-2.93	0.0043	Tukey-Kramer	0.0474
Calan.Treatment*SEX	DDG	STR	MET	HFR	122.91	36.4353	90	3.37	0.0011	Tukey-Kramer	0.0136
Calan.Treatment*SEX	DDG	STR	MET	STR	-7.8914	38.5694	90	-0.20	0.8383	Tukey-Kramer	0.9999
Calan.Treatment*SEX	MET	HFR	MET	STR	-130.80	39.6146	90	-3.30	0.0014	Tukey-Kramer	0.0168

Figure 33: $Differences\ of\ Least\ Squares\ Means\ table.$

ANCOVA for Calf Final Body Weight

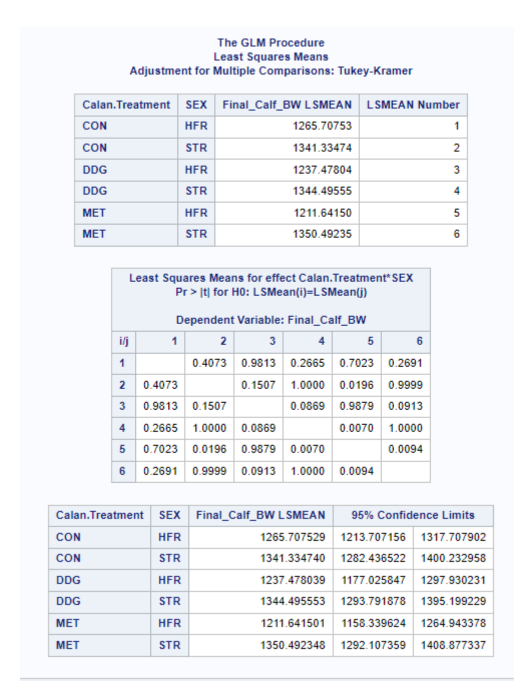


Figure 34: Tukey's HSD test and comparisons for ANCOVA model.

Appendix D - Additional R Output

Ordinal Logistic Regression for Easing

```
Call:
```

```
polr(formula = Calving.Ease ~ Calan.Treatment * SEX, data = data2,
    Hess = TRUE)
```

Coefficients:

	Value	Std. Error	t value
Calan.TreatmentDDG	-16.8625	0.7285	-23.1471
Calan.TreatmentMET	-1.1165	1.2030	-0.9281
SEXSTR	-0.9264	1.2079	-0.7669
${\tt Calan.TreatmentDDG:SEXSTR}$	16.6163	0.7285	22.8091
Calan.TreatmentMET:SEXSTR	2.4732	1.7137	1.4432

Intercepts:

```
Value Std. Error t value
1|2 1.7349 0.6250 2.7759
2|3 2.9393 0.7811 3.7632
```

Residual Deviance: 65.95121

AIC: 79.95121

Binomial regression for Easing

```
Call:
```

```
glm(formula = Calving.Ease.bin ~ Calan.Treatment * SEX, family = binomial,
    data = data2)
```

Coefficients:

	Estimate	Std. Error	${\tt z}$ value	Pr(> z)	
(Intercept)	-1.6740	0.6292	-2.661	0.0078	**
Calan.TreatmentDDG	-16.8921	1743.2486	-0.010	0.9923	
Calan.TreatmentMET	-1.1592	1.2061	-0.961	0.3365	
SEXSTR	-0.9651	1.2113	-0.797	0.4256	
Calan.TreatmentDDG:SEXSTR	16.5867	1743.2492	0.010	0.9924	
Calan.TreatmentMET:SEXSTR	2.4120	1.7154	1.406	0.1597	

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

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 60.695 on 100 degrees of freedom Residual deviance: 54.599 on 95 degrees of freedom

AIC: 66.599

Number of Fisher Scoring iterations: 17

Ordinal Logistic Regression for Vigor

Call:

```
polr(formula = Calf.Vigor ~ Calan.Treatment * SEX, data = data2,
    Hess = TRUE)
```

Coefficients:

	Value	Std. Error	t value
Calan.TreatmentDDG	-0.02146	0.8519	-0.02519
Calan.TreatmentMET	-0.17876	0.8454	-0.21145
SEXSTR	-1.27076	1.1768	-1.07981
${\tt Calan.TreatmentDDG:SEXSTR}$	1.22910	1.4516	0.84673
Calan.TreatmentMET:SEXSTR	0.83354	1.5350	0.54301

Intercepts:

Value Std. Error t value 1|2 1.3316 0.5592 2.3812 2|3 2.0682 0.6012 3.4399

Residual Deviance: 113.059

AIC: 127.059

Binomial regression for Vigor

Call:

```
glm(formula = Calf.Vigor ~ Calan.Treatment * SEX, family = binomial,
    data = data2)
```

Coefficients:

```
Estimate Std. Error z value Pr(>|z|)
(Intercept) -1.32176 0.56273 -2.349 0.0188 *
Calan.TreatmentDDG 0.02247 0.86076 0.026 0.9792
```

```
      Calan.TreatmentMET
      -0.28768
      0.84656
      -0.340
      0.7340

      SEXSTR
      -1.31730
      1.17817
      -1.118
      0.2635

      Calan.TreatmentDDG:SEXSTR
      1.23029
      1.45768
      0.844
      0.3987

      Calan.TreatmentMET:SEXSTR
      1.05494
      1.53786
      0.686
      0.4927
```

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

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 91.548 on 100 degrees of freedom Residual deviance: 89.469 on 95 degrees of freedom

AIC: 101.47

Number of Fisher Scoring iterations: 5