# final project

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### Motivation

Breast cancer prognosis is essential for guiding treatment and improving patient survival. This study uses logistic regression to identify key factors that influence whether patients survive or not (status: Alive or Dead). The goal is to build a clear and understandable model to predict survival and see how well it works across different racial groups. It also focuses on fairness by examining performance differences between groups and finding ways to improve generalizability of the model.

# Data Cleaning and Preprocessing

# **Exploratory Data Analysis**

Summary statistics

#### Distribution Visualization

Categorical Variables Our modified dataset contains 10 categorical variables:

- race: Patient's race (Black, White, Other).
- marital\_status: Patient's marital status (Divorced, Married, Separated, Single, Widowed).
- t\_stage: Tumor stage (T1, T2, T3, T4). ("T" refers to the size of the primary tumor)
- n\_stage: Lymph node stage (N1, N2, N3). (extent of cancer spread to nearby lymph nodes)
- x6th\_stage: Adjusted AJCC 6th stage (IIA, IIB, IIIA, IIIB, IIIC).
- differentiate: Tumor differentiation (Well, Moderately, Poorly, Undifferentiated). grade: Grade of the tumor (1–4).
- a\_stage: Tumor spread stage (Regional, Distant).
- estrogen\_status: Estrogen receptor status (Positive, Negative).
- progesterone status: Progesterone receptor status (Positive, Negative).

To examine the association of these variables with the binary outcome status (Alive/Dead), I used Cramér's V, which quantifies the strength of the association between two categorical variables based on the Chi-Square statistic. Cramér's V ranges from 0 (no association) to 1 (perfect association). This method helps identify the predictors most strongly associated with survival status, enabling us to prioritize variables for modeling.

The plot shows the strength of association (Cramér's V) between **survival status** and each categorical predictor:

- 1. **x6th\_stage** and **n\_stage** have the **highest Cramér's V values** (around 0.25), indicating they are the most informative predictors of survival status in the dataset. These variables reflect tumor stage and lymph node involvement, which are critical factors in breast cancer prognosis.
- 2. Estrogen\_status, progesterone\_status, and grade exhibit moderate associations (Cramér's V around 0.15–0.2). These variables provide meaningful information about hormone status and differentiation, making them important contributors to survival prediction.
- 3. **Differentiate** and **t\_stage** show moderate but slightly lower associations compared to the top variables, suggesting their relevance to survival status.
- 4. a\_stage, marital\_status, and race have lower Cramér's V values (less than 0.1), indicating weaker associations with survival status. Although these variables contribute limited information individually, they may still be useful in interaction terms or when combined with other predictors.

By focusing on variables with higher Cramér's V values, we can build more efficient and predictive models for survival status analysis.

Both plots confirm a disparity in survival outcomes across racial groups. Non-White patients, particularly Black patients, show a higher likelihood of death. Effect of Combining Groups: Combining Black and Other into Minority Non-White simplifies the comparison and reduce the racial disparities between Majority White and Minority Non-White groups.

Continuous Variables Histograms were created for each continuous variable to visually examine their distributions and identify potential skewness. The results revealed that, except for age, the other three variables exhibited significant right skewness, indicating the need for transformation if we want to perform linear regression. Though we are not doing linear regression, transformation has still been performed here, we see that after performing log-transformation, there is a improvement on the skewness of tumor\_size, regional\_node\_examined and reginol\_node\_positive.

Boxplots were then generated for each continuous variable stratified by survival status to visually assess their relationships with the binary outcome.

### Pairwise Relationships and Interactions

A correlation matrix was generated to examine the relationships between the variables. Small circles (near-zero correlations) are observed for age and all other variables; tumor\_size and regional\_node\_examined suggesting weak or no relationships. regional\_node\_positive is moderately associated with both tumor size (0.24) and regional nodes examined (0.41), which might influence modeling decisions. There are no strong correlations (close to +-1), suggesting no immediate multicollinearity issues among these variables.

# Modeling - Logistic Regression

Since the outcome is Binary (e.g., Alive/Dead), We decide to proceed with logistic regression instead of linear regression. Logistic regression outputs odds ratios, which are more interpretable for binary classification problems. For example, it tells you how much the odds of death increase with a unit increase in a predictor.

### Checking Assumptions and Transformations

**Binary or Dichotomous Response Variable** The response variable (status) has exactly two categories: Alive and Dead.

No Multicollinearity From the alias() output: grade2, grade3, and grade4 are collinear with other predictors. x6th\_stageIIIC is collinear with n\_stage or other levels of x6th\_stage. differentiate levels also overlap in prediction with grade.

Given that both x6th\_stage and n\_stage are strong predictors, with x6th\_stage showing a slightly higher association, n\_stage can be dropped as its information is already captured by x6th\_stage. Similarly, t\_stage, which is related to tumor size, can also be removed to avoid redundancy. The variable differentiate can be excluded as well, as it overlaps with grade, with both variables reflecting tumor differentiation. In the context of breast cancer, "regional node positive" indicates the presence of cancer cells in nearby lymph nodes, while "regional node examined" refers to the surgical removal and analysis of these nodes to determine cancer spread. From the previous correlation matrix, regional\_node\_positive shows a moderate association with both tumor size (0.24) and regional\_node\_examined (0.41), suggesting some redundancy. To simplify the model and minimize multicollinearity, we choose to drop regional\_node\_positive.

The results show that all VIF values are below 5, which indicates that multicollinearity is no longer a concern in your logistic regression model.

Linear Relationship to Log Odds From the plot, we see that there's a linear relationship between continuous predictors (log transformed) and the log odds of the outcome.

#### No Extreme Outliers

```
## Robustness weights w.r * w.x:
## 3546 weights are ~= 1. The remaining 478 ones are summarized as
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.2123 0.4537 0.5997 0.6159 0.7762 0.9983
##
## Number of observations: 4024
```

In large datasets, Cook's Distance can flag numerous observations as influential, often due to large sample size. These flagged points may represent true variability in the population rather than errors. The code results indicate that removing influential points harms the model, likely by disrupting essential data structure. Coefficients for variables like grade and intercept became unstable and unreliable, with extreme standard errors.

In contrast, the robust logistic regression (model\_robust) provided stable and reliable estimates while effectively mitigating the impact of influential observations. Key predictors, such as age\_log, rn\_examined\_log, grade, x6th\_stage, and race, retained coefficients similar to the standard logistic regression. The robustness weights further show that about 12% of the data (478 out of 4024 observations) had reduced influence, while the majority (~88%, or 3546 observations) remained largely unaffected with weights close to 1. This approach accounts for influential points by down-weighting their impact, ensuring the model is less sensitive to extreme values without the need to remove data.

While both original and robust logistic regression are viable options for future analysis, we will proceed with the original logistic regression due to our limited familiarity with robust logistic regression techniques

**Independence of Errors** Since there are no group IDs, the independence assumption is satisfied.

#### Model Evaluation and Selection

We use original Logistic Regression as the primary models, considering its earlier performance.

full results are attached in the end Full Model: glm(formula = status  $\sim$  . , family = binomial, data = model\_data\_3) AIC: 3039.8

Forward Model glm(formula = status  $\sim$  race + marital\_status + x6th\_stage + grade + a\_stage + estrogen\_status + progesterone\_status + tumor\_size\_log + rn\_examined\_log + age\_log, family = binomial, data = model data 3) AIC: 3039.8

Backward Model:  $glm(formula = status \sim race + marital\_status + x6th\_stage + grade + estrogen\_status + progesterone\_status + rn\_examined\_log + age\_log, family = binomial, data = model_data_3) AIC:3037.5$ 

Stepwise Model:  $glm(formula = status \sim race + marital\_status + x6th\_stage + grade + estrogen\_status + progesterone\_status + rn\_examined\_log + age\_log, family = binomial, data = model\_data_3) AIC:3037.5$ 

```
## Best Model by Cp: 10 variables, raceBlack x6th_stageIIIA x6th_stageIIIB x6th_stageIIIC grade3 grade
```

## Best Model by BIC: 10 variables, raceBlack x6th\_stageIIIA x6th\_stageIIIB x6th\_stageIIIC grade3 grad

## Best Model by Adjusted R2: 10 variables, raceBlack x6th stageIIIA x6th stageIIIB x6th stageIIIC gra-

```
## [1] Predictor1 Predictor2 P_Value
## <0 rows> (or 0-length row.names)
```

Since resulting dataframe is empty, it suggests that none of the interaction terms tested in our dataset have p-values less than 0.05. This outcome aligns with our earlier exploratory analyses (Cramér's V, correlation matrix, and model diagnostics), which indicated that interaction effects are likely weak or non-existent in your data.

**Final Model** Since Best subset selection evaluates all possible combinations of predictors (including all levels of factor variables) to find the model with the best fit according to criteria like adjusted R2, Cp and BIC, each level of a factor variable (e.g., x6th\_stageIIIA) can be treated as an independent binary variable (dummy variable) in this process. Thus we will not use the best subset selection, as we need consider the factor as a whole for interpretability.

The final model was chosen through forward, backward, and stepwise selection methods using AIC as the selection criterion. The model balances interpretability and predictive performance, including only significant predictors that contribute to understanding survival outcomes:

```
glm(formula = status ~ race + marital_status + x6th_stage + grade + estrogen_status +
progesterone_status + rn_examined_log + age_log, family = binomial, data = model_data_3)
```

In logistic regression, coefficients represent the change in the log-odds of the outcome (status = Dead) for a one-unit increase in a predictor, holding other variables constant. Odds ratios provide a more interpretable measure.

The key coefficients included are 1. Race: - raceBlack: Positive coefficient implies higher odds of death for Black patients compared to the reference group (White patients). Odds ratio > 1 indicates increased risk for Black patients. - raceOther: Negative coefficient indicates slightly reduced odds of death for patients classified as "Other" compared to White patients.

- 2. Marital Status: The marital status categories have weak effects (some not statistically significant). For instance, being married or widowed appears to have no strong association with survival compared to the reference group.
- 3. Adjusted AJCC 6th stage (x6th\_stage): is the strongest predictor of death risk. Higher stages (IIIA, IIIB, IIIC) significantly increase the odds of death compared to stage IIA (reference group). Patients in stage IIIC have the highest odds of death, indicating advanced tumor progression is a critical predictor of survival.

- 4. Tumor Grade (grade): Grades 3 and 4 significantly increase the odds of death compared to Grade 1, indicating poorly differentiated tumors are associated with worse outcomes.
- 5. Hormone Receptor Status: estrogen\_statusPositive and progesterone\_statusPositive: Negative coefficients suggest reduced odds of death for patients with positive estrogen or progesterone receptor status. This reflects the improved prognosis often associated with hormone-sensitive tumors.
- 6. Log-transformed Predictors: -rn\_examined\_log: Negative coefficient indicates increased survival odds with more regional lymph nodes examined. This might reflect more aggressive or effective treatment strategies. -age\_log: Positive coefficient implies older patients are at higher risk of death.

So x6th\_stage, grade, estrogen\_statusPositive and progesterone\_statusPositive, age\_log and rn\_examined\_log are significant predictors affecting the risk.

**Model Performance** full results are attached in the end We did Cross-Validation (10 fold) ROC-AUC: 0.7400, the overall performance of the model is moderately good, with an acceptable ability to distinguish between "Alive" and "Dead" outcomes. Sensitivity: 0.985, the model is highly sensitive, meaning it correctly identifies most of the "Dead" cases. Specificity: 0.122, the model has very low specificity, indicating difficulty in correctly identifying "Alive" cases.

```
## ROC-AUC for White group: 0.7503654
## ROC-AUC for Black group: 0.7021491
## ROC-AUC for Other group: 0.658431
## ROC-AUC for Minority group: 0.7313469
```

Performance by Race Groups White Group (Majority): - ROC-AUC: 0.7504 - The model performs better for the majority race group, achieving the highest predictive ability among the subgroups.

Black Group (Minority): - ROC-AUC: 0.7021 - The performance for Black patients is lower than for White patients, indicating potential disparities in prediction accuracy.

Other Group (Minority): - ROC-AUC: 0.6584 - The lowest ROC-AUC is for the "Other" race group, suggesting the model struggles most with this subgroup.

Combined Minority Group (Black + Other): - ROC-AUC: 0.7313 - When grouped together, the model's performance for minorities improves but remains slightly lower than for the White group.

This disparity indicate a potential fairness issue, which need us to reduce the performance gap.

```
## Reweighted Model - ROC-AUC for White group: 0.7486181
## Reweighted Model - ROC-AUC for Minority group: 0.7358304
## Reweighted Model - ROC-AUC for Black group: 0.7051653
## Reweighted Model - ROC-AUC for Other group: 0.6657164
```

The reweighted model shows improved fairness compared to the original combined model: Reweighting has reduced the performance gap between the White and minority groups. The ROC-AUC for the minority group (0.7358) is now closer to that of the majority group (0.7486).

### Conclusion

#### **Table**

### Finding and Interretation

The final reweighted logistic regression model identifies significant predictors of mortality risk (Alive vs Dead) for breast cancer patients. The table above summarizes the parameter estimates, highlighting the direction, magnitude, and significance of each predictor. The study confirms that Adjusted AJCC 6th stage (x6th\_stage) and grade (Grade of the tumor) are the most critical factors in predicting mortality, which is consistent with clinical guidelines. Hormone receptor status also plays a vital role, as positive estrogen/progesterone receptors improve survival. Race also significantly impacts mortality risk. Black patients experience a higher risk of death compared to White patients. This disparity may stem from differences in access to healthcare, late diagnosis, or biological factors. The reweighted model successfully improves fairness across racial groups while maintaining overall predictive accuracy. Although the model performs slightly worse for minority groups, the gap has been reduced compared to the original model.

### Limitation

The model assumes no significant interaction effects, which aligns with the data but could miss complex relationships. The disparity in model performance for "Other" racial groups suggests the need for further data collection or alternative modeling techniques.

#### Contribution

Ada Guo wrote the abstract and introduction, compiled data cleaning and EDA results, and synthesized findings into cohesive sections for the report. Khue Nguyen analyzed results, wrote the conclusion and limitations, refined the report structure, and ensured clarity and completeness by adding references. Yixin Zheng wrote the methods and appendix section and constructed the Rmd file(cleaning, correlation matrix, Cramér's V, assumption checks, feature selection, model, cross-validation, fairness reweight).

# Reference

# Table, Plots, and Code Results

# Tables

Table 1: Skim Summary for Categorical Variables

skim_variable	n_missing con	nplete_ra	t∉actor.ordere	d factor.n_uniqu	ufactor.top_counts
race	0	1	FALSE	3	Whi: 3413, Oth: 320, Bla: 291
marital_status	0	1	FALSE	5	Mar: 2643, Sin: 615, Div: 486, Wid: 235
t_stage	0	1	FALSE	4	T2: 1786, T1: 1603, T3: 533, T4: 102
n_stage	0	1	FALSE	3	N1: 2732, N2: 820, N3: 472
x6th_stage	0	1	FALSE	5	IIA: 1305, IIB: 1130, III: 1050, III: 472
differentiate	0	1	FALSE	4	Mod: 2351, Poo: 1111, Wel: 543, Und: 19
grade	0	1	FALSE	4	2: 2351, 3: 1111, 1: 543, 4: 19
a_stage	0	1	FALSE	2	Reg: 3932, Dis: 92
estrogen_status	0	1	FALSE	2	Pos: 3755, Neg: 269
progesterone_stati	us 0	1	FALSE	2	Pos: 3326, Neg: 698
status	0	1	FALSE	2	Ali: 3408, Dea: 616

Table 2: Skim Summary for Numeric Variables

skim_variable	n_missing con	nplete_rate	mean	$\operatorname{sd}$	p0	p25	p50	p75	p100	hist
age	0	1 5	3.972167	8.963134	30	47	54	61	69	
tumor_size	0	1 3	30.473658	21.119696	1	16	25	38	140	<b>L</b>
regional_node_examin	ned 0	1 1	4.357107	8.099675	1	9	14	19	61	
reginol_node_positive	0	1 4	4.158052	5.109331	1	1	2	5	46	<b></b>

Table 3: Summary Statistics Grouped by Survival Status

status	mean_age	$sd\_age$	$mean\_tumor\_sizesd$	_tumor_size	prop_white	prop_black_other	n_obs
Alive	53.75910	8.808420	29.26878	20.30317	0.8518192	0.1481808	3408
Dead	55.15097	9.698291	37.13961	24.11611	0.8279221	0.1720779	616

Table 4: Variance Inflation Factors for Predictors

Variable	GVIF	Df	GVIF_Ratio
age	1.106908	1	1.052097
race	1.058083	2	1.014215
marital_status	1.127489	4	1.015112
x6th_stage	1.967732	4	1.088293
grade	1.118473	3	1.018836
a_stage	1.210950	1	1.100432
tumor_size	1.365304	1	1.168462
estrogen status	1.484914	1	1.218570

Variable	GVIF	Df	GVIF_Ratio
progesterone_status regional node examined	1.434692 1.225729	1	1.197786 1.107127

Table 5: Best Subset Selection Summary

$Num\_Predictors$	$Adj_R2$	Cp	BIC
1	0.0541705	298.36391	-208.5087
2	0.0808751	177.39536	-316.4581
3	0.0879639	146.00298	-340.3144
4	0.0945059	117.12118	-361.9835
5	0.0989525	97.81162	-374.4942
6	0.1026866	81.76122	-383.9066
7	0.1060954	67.20233	-391.9243
8	0.1090479	54.73108	-397.9391
9	0.1112815	45.54176	-400.7425
10	0.1133639	37.04659	-402.8850

Table 6: Reweighted Logistic Regression Model Results

Predictor	Estimate	Std_Error	Odds_Ratio X9	5CILower.	X95CIUpper.	P.Value
(Intercept)	-5.126	1.161	0.006	0.001	0.058	1.02e-05
raceBlack	0.455	0.135	1.577	1.211	2.052	0.000713
raceOther	-0.426	0.147	0.653	0.490	0.871	0.003683
$marital\_statusMarried$	-0.248	0.134	0.781	0.600	1.015	0.064877
$marital\_statusSeparated$	0.696	0.350	2.006	1.011	3.981	0.046555
marital_statusSingle	0.034	0.162	1.035	0.753	1.422	0.833041
$marital\_statusWidowed$	0.123	0.204	1.131	0.759	1.687	0.544852
$x6th\_stageIIB$	0.533	0.136	1.704	1.305	2.225	9.02e-05
$x6th\_stageIIIA$	0.978	0.134	2.660	2.047	3.457	2.52e-13
$x6th\_stageIIIB$	1.565	0.287	4.781	2.726	8.386	4.82e-08
$x6th\_stageIIIC$	2.006	0.152	7.433	5.519	10.012	< 2e-16
grade2	0.506	0.172	1.659	1.184	2.324	0.003266
grade3	0.862	0.180	2.368	1.664	3.370	1.67e-06
grade4	1.825	0.521	6.203	2.233	17.232	0.000463
estrogen_statusPositive	-0.714	0.166	0.490	0.354	0.678	1.70 e-05
progesterone_statusPositive	e -0.503	0.121	0.605	0.477	0.767	3.37e-05
$rn\_examined\_log$	-0.277	0.076	0.758	0.653	0.880	0.000270
age_log	0.974	0.276	2.648	1.543	4.545	0.000410

# Plots

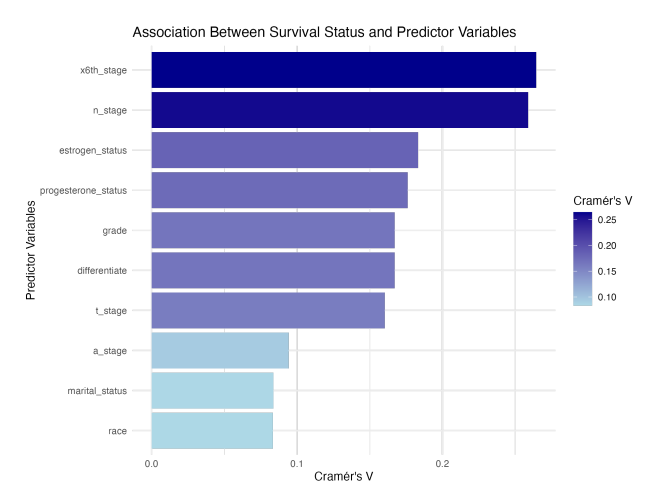


Figure 1: Cramér's V Associations

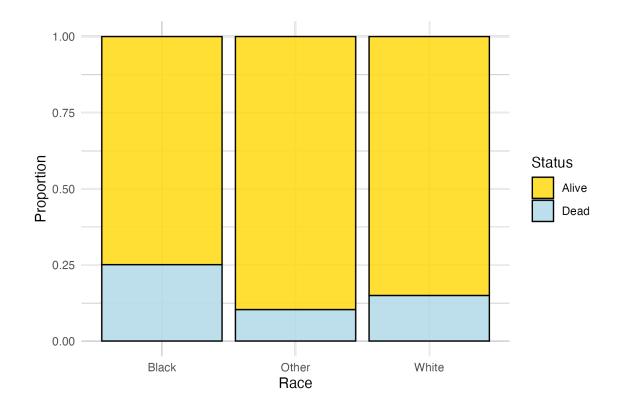


Figure 2: Survival Status by Race

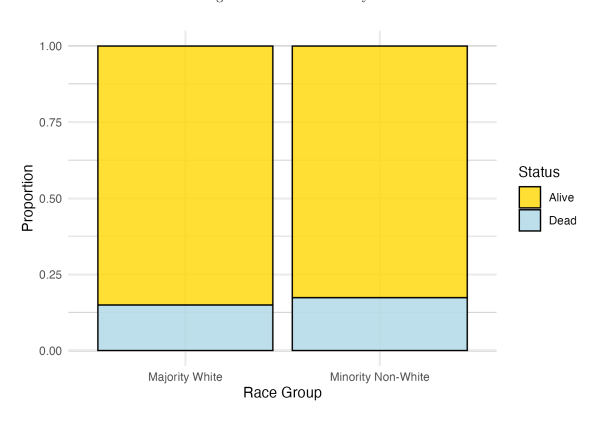


Figure 3: Survival Status by Combined Race Groups

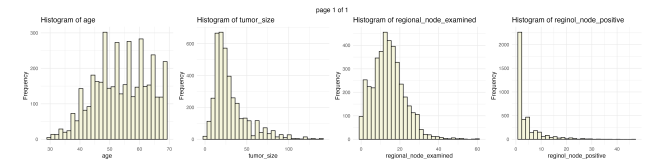


Figure 4: Histograms for Original Continuous Variables

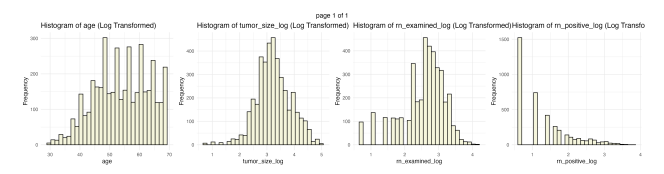


Figure 5: Histograms for Log-Transformed Continuous Variables

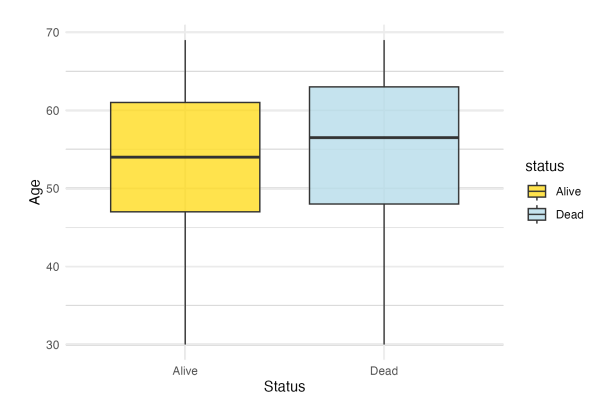


Figure 6: Age by Survival Status

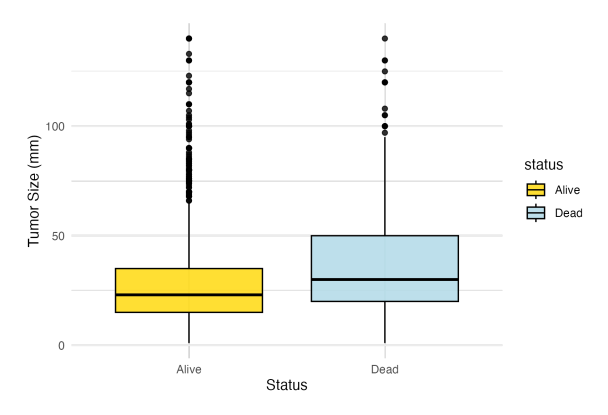


Figure 7: Tumor Size by Survival Status

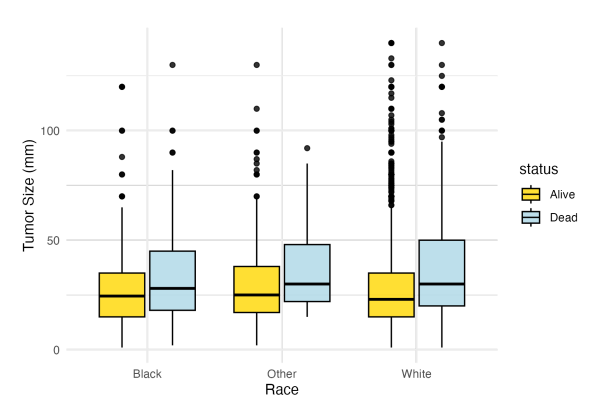


Figure 8: Tumor Size by Race and Survival Status

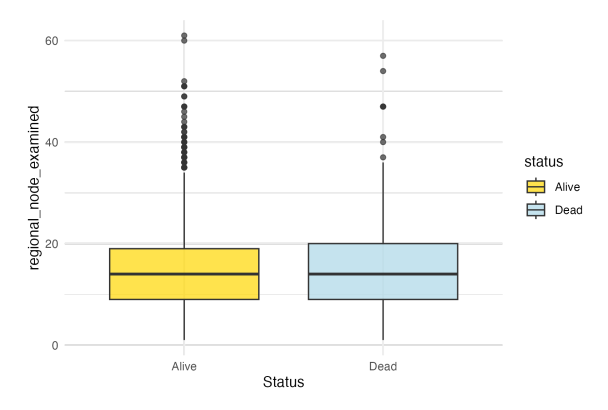


Figure 9: Regional Node Examined by Survival Status

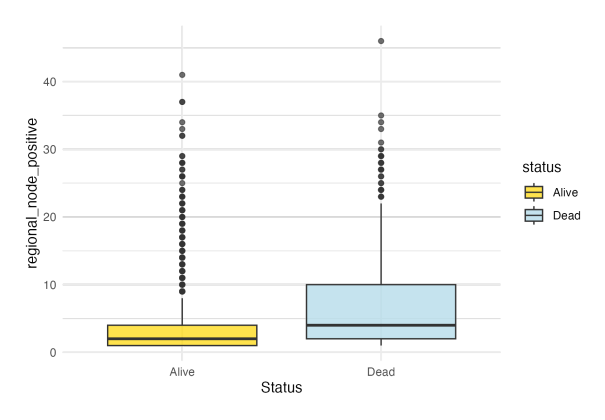


Figure 10: Regional Node Positive by Survival Status

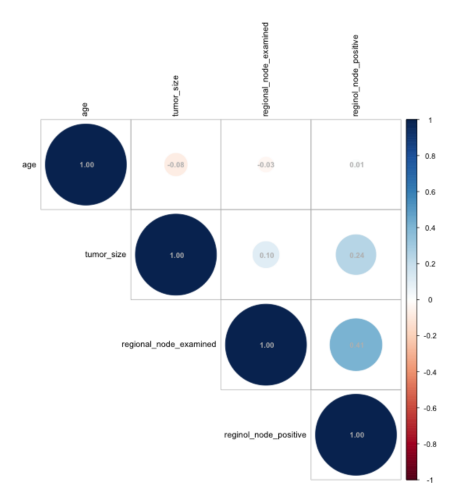


Figure 11: Correlation Matrix for Continuous Variables

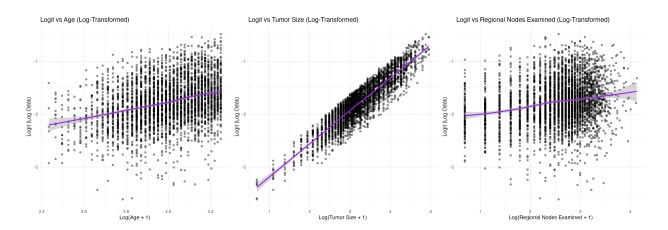


Figure 12: Log Odds Relationship with Predictors

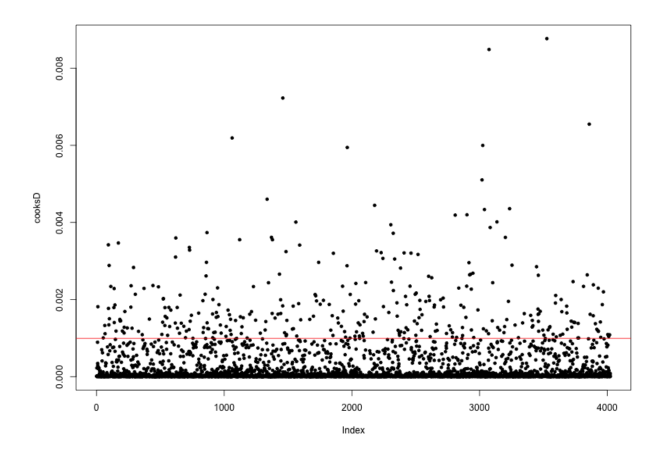


Figure 13: Cook's Distance for Outlier Detection

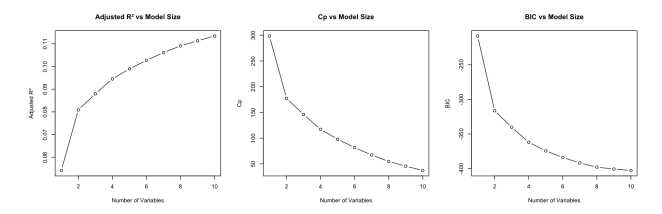


Figure 14: Best Subset Selection Performance Metrics

#### Code Results

For full code results, please refer to the .txt files available in the results folder of the GitHub repository: Yixin-Zheng/p8130\_finalproject.

# Code Appendix

```
knitr::opts chunk$set(echo = TRUE)
library(tidyverse)
library(janitor)
library(skimr)
library(dplyr)
library(ggplot2)
library(caret)
library(corrplot)
library(lsr)
library(vcd)
library(car)
library(gridExtra)
library(robustbase)
library(leaps)
library(pROC)
library(knitr)
# Import data and clean column names
data <- read.csv("./data/Project_2_data.csv") %>%
  clean_names()
# Select relevant covariates (variables 1-14) and outcome variable
model_data <- data %>%
  dplyr::select(-survival_months)
# Convert categorical variables to factors and relabel `grade`
model_data <- model_data %>%
  mutate(
    race = factor(race),
    marital_status = factor(marital_status),
    t_stage = factor(t_stage),
    n_stage = factor(n_stage),
    x6th_stage = factor(x6th_stage),
    differentiate = factor(differentiate),
    a_stage = factor(a_stage),
    estrogen_status = factor(estrogen_status),
    progesterone_status = factor(progesterone_status),
    status = factor(status, levels = c("Alive", "Dead")),
    grade = case_when(
      grade == "1" ~ "1",
      grade == "2" ~ "2",
      grade == "3" ~ "3",
      grade == " anaplastic; Grade IV" ~ "4",
      TRUE ~ NA_character_
    ) %>% factor(levels = c("1", "2", "3", "4"))
```

```
# Summarize structure of the cleaned dataset
summary(model_data)
# Summary statistics for continuous and categorical variables
skimmed data <- skim(model data)</pre>
skim_categorical <- skimmed_data %>%
  filter(skim_type == "factor") %>%
  select(-starts with("numeric"), -skim type) %>%
  na.omit()
write.csv(skim_categorical, "tables/skim_categorical_summary.csv", row.names = FALSE)
skim_numeric <- skimmed_data %>%
  filter(skim_type == "numeric") %>%
  select(-starts_with("factor"), -skim_type) %>%
  na.omit()
colnames(skim_numeric) <- gsub("^numeric\\.", "", colnames(skim_numeric))</pre>
write.csv(skim_numeric, "tables/skim_numeric_summary.csv", row.names = FALSE)
# Key statistics grouped by survival status
summary_by_status <- model_data %>%
  group by(status) %>%
  summarise(
    mean_age = mean(age, na.rm = TRUE),
    sd_age = sd(age, na.rm = TRUE),
    mean_tumor_size = mean(tumor_size, na.rm = TRUE),
    sd_tumor_size = sd(tumor_size, na.rm = TRUE),
    prop_white = mean(race == "White", na.rm = TRUE),
    prop_black_other = mean(race != "White", na.rm = TRUE),
    n_{obs} = n()
  )
write.csv(summary_by_status, "tables/summary_by_status.csv", row.names = FALSE)
# Define categorical variables to analyze
variables <- c("race", "marital_status", "t_stage", "n_stage", "x6th_stage",</pre>
               "differentiate", "grade", "a_stage", "estrogen_status", "progesterone_status")
# Initialize a vector to store Cramér's V results
results <- numeric(length(variables))
# Calculate Cramér's V for each variable
for (i in seq_along(variables)) {
  var <- variables[i]</pre>
  # Select outcome and predictor variable, omitting missing values
  df_temp <- model_data %>%
    dplyr::select(status, all_of(var)) %>%
    na.omit()
  # Convert both columns to factors
  x <- droplevels(as.factor(df_temp$status))</pre>
  y <- droplevels(as.factor(df_temp[[var]]))</pre>
  # Create contingency table and calculate Cramér's V
```

```
table_var <- table(x, y)</pre>
 results[i] <- cramersV(table_var)</pre>
# Create a dataframe with results
association_df <- data.frame(Variable = variables, CramersV = results)
# Plot Cramér's V values
cramerV_association <- ggplot(association_df, aes(x = reorder(Variable, CramersV), y = CramersV, fill =</pre>
  geom_bar(stat = "identity") +
  coord flip() +
  scale_fill_gradient(low = "lightblue", high = "darkblue") +
   title = "Association Between Survival Status and Predictor Variables",
   x = "Predictor Variables",
   y = "Cramér's V",
   fill = "Cramér's V"
  theme_minimal()
ggsave("plots/cramerV_association.png", plot = cramerV_association, width = 8, height = 6)
# Proportional Bar Plot for Survival Status by Race
race_barplot <- ggplot(model_data, aes(x = race, fill = status)) +</pre>
  geom_bar(position = "fill", alpha = 0.8, color = "black") +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme minimal() +
 labs(
   x = "Race",
   y = "Proportion",
   fill = "Status"
ggsave("plots/race_proportional_barplot.png", plot = race_barplot, width = 6, height = 4)
# Combine "Black" and "Other" into a single group "Minority Non-White"
model_data_race_combined <- model_data %>%
 mutate(
   race_combined = case_when(
     race == "White" ~ "Majority White",
     race %in% c("Black", "Other") ~ "Minority Non-White"
   ),
   race_combined = factor(race_combined, levels = c("Majority White", "Minority Non-White"))
# Proportional Bar Plot for Combined Race Groups
race_combined_barplot <- ggplot(model_data_race_combined, aes(x = race_combined, fill = status)) +</pre>
  geom_bar(position = "fill", alpha = 0.8, color = "black") +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme_minimal() +
 labs(
   x = "Race Group",
   y = "Proportion",
```

```
fill = "Status"
 )
ggsave("plots/race_combined_proportional_barplot.png", plot = race_combined_barplot, width = 6, height
continuous_vars <- model_data %>%
  dplyr::select(age, tumor_size, regional_node_examined, reginol_node_positive) %>%
 na.omit()
df1 <- as.data.frame(continuous vars)</pre>
hist_list <- lapply(names(continuous_vars), function(col) {</pre>
  ggplot(continuous_vars, aes_string(x = col)) +
    geom_histogram(fill = "beige", color = "black", bins = 30) +
    labs(
      title = paste("Histogram of", col),
     x = col,
      y = "Frequency"
    ) +
    theme_minimal()
})
# Arrange plots in a grid
hist_grid <- marrangeGrob(hist_list, nrow = 1, ncol = 4)</pre>
ggsave("plots/original_histograms_grid.png", hist_grid, width = 16, height = 4)
df_log <- df1 %>%
 dplyr::select(-tumor_size, -regional_node_examined, -regional_node_positive) %>%
    tumor_size_log = log(df1$tumor_size + 1),
    rn_examined_log = log(df1$regional_node_examined + 1),
    rn_positive_log = log(df1$reginol_node_positive + 1)
log_hist_list <- lapply(names(df_log), function(col) {</pre>
  ggplot(df_log, aes_string(x = col)) +
    geom_histogram(fill = "beige", color = "black", bins = 30) +
    labs(
      title = paste("Histogram of", col, "(Log Transformed)"),
     x = col,
     y = "Frequency"
    ) +
    theme_minimal()
})
# Arrange plots in a grid
log_hist_grid <- marrangeGrob(log_hist_list, nrow = 1, ncol = 4)</pre>
ggsave("plots/log_transformed_histograms_grid.png", log_hist_grid, width = 16, height = 4)
# Age by survival status
age_boxplot <- ggplot(model_data, aes(x = status, y = age, fill = status)) +
  geom_boxplot(alpha = 0.7) +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme_minimal() +
  labs(
    x = "Status",
```

```
y = "Age"
 )
ggsave("plots/age_by_status_boxplot.png", plot = age_boxplot, width = 6, height = 4)
# Tumor size by survival status
tumor_boxplot <- ggplot(model_data, aes(x = status, y = tumor_size, fill = status)) +</pre>
  geom_boxplot(alpha = 0.8, color = "black") +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme minimal() +
 labs(
   x = "Status",
   y = "Tumor Size (mm)"
ggsave("plots/tumor_size_by_status_boxplot.png", plot = tumor_boxplot, width = 6, height = 4)
# Tumor size by race and survival status
tumor_race_boxplot <- ggplot(model_data, aes(x = race, y = tumor_size, fill = status)) +</pre>
  geom_boxplot(alpha = 0.8, color = "black") +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme minimal() +
 labs(
   x = "Race",
   y = "Tumor Size (mm)"
ggsave("plots/tumor_size_by_race_status_boxplot.png", plot = tumor_race_boxplot, width = 6, height = 4)
# Regional Node Examined by survival status
rn_examined_boxplot <- ggplot(model_data, aes(x = status, y = regional_node_examined, fill = status)) +
  geom_boxplot(alpha = 0.7) +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme_minimal() +
 labs(
   x = "Status",
   y = "regional_node_examined"
  )
ggsave("plots/rn_examined_by_status_boxplot.png", plot = rn_examined_boxplot, width = 6, height = 4)
# Reginol Node Positive by survival status
rn_positive_boxplot <- ggplot(model_data, aes(x = status, y = reginol_node_positive, fill = status)) +</pre>
  geom_boxplot(alpha = 0.7) +
  scale_fill_manual(values = c("Alive" = "gold", "Dead" = "lightblue")) +
 theme_minimal() +
 labs(
   x = "Status",
    y = "regional_node_positive"
ggsave("plots/rn_positive_by_status_boxplot.png", plot = rn_positive_boxplot, width = 6, height = 4)
correlation_matrix <- cor(continuous_vars, use = "pairwise.complete.obs")</pre>
# Pairwise relationships (correlation matrix for continuous variables)
```

```
correlation_plot <- function() {</pre>
  corrplot(
    correlation_matrix,
    method = "circle",
    type = "upper",
   tl.col = "black"
   addCoef.col = "grey",
   number.cex = 0.8,
   tl.cex = 0.9
 )
}
png("plots/correlation_matrix_plot.png", width = 800, height = 600)
correlation_plot()
dev.off()
unique(model_data$status)
# Fit an initial logistic regression model
model_data_1 <- model_data %>%
 mutate(
    race = relevel(race, ref = "White"), # Set "White" as reference
    grade = relevel(grade, ref = "1"),  # Set "Grade 1" as reference
    x6th_stage = relevel(x6th_stage, ref = "IIA") # Set "IIA" as reference
alias_results <- capture.output(alias(glm(status ~ ., data = model_data_1, family = binomial)))</pre>
# Save the results to a text file in the `results` folder
writeLines(alias_results, "results/alias_results_model1.txt")
model_data_2 <- model_data_1 %>%
 dplyr::select(-differentiate, -n_stage, -t_stage, -reginol_node_positive)
# Perform multicollinearity check with alias()
alias_results_2 <- capture.output(alias(glm(status ~ ., data = model_data_2, family = binomial)))
writeLines(alias_results_2, "results/alias_results_model2.txt")
# Fit a logistic regression model
model_vif <- glm(status ~ ., data = model_data_2, family = binomial)</pre>
# Calculate VIF
vif_values <- vif(model_vif)</pre>
vif_df <- as.data.frame(vif_values)</pre>
vif_df <- tibble::rownames_to_column(vif_df, var = "Variable")</pre>
colnames(vif df) <- c("Variable", "GVIF", "Df", "GVIF Ratio")</pre>
write.csv(vif_df, "tables/vif.csv", row.names = FALSE)
continuous_vars_log_odds <- model_data_2 %>%
 dplyr::select(age, tumor_size, regional_node_examined, status) %>% # Include 'status'
 na.omit()
# Log-transform tumor size and regional nodes examined
df_log_odds <- continuous_vars_log_odds %>%
 mutate(
    age_log = log(age + 1),
    tumor_size_log = log(tumor_size + 1),
```

```
rn_examined_log = log(regional_node_examined + 1)
  )
linearity_test <- glm(status ~ age_log + tumor_size_log + rn_examined_log,</pre>
                       data = df_log_odds,
                       family = binomial)
df_log_odds$logit <- predict(linearity_test, type = "link")</pre>
plot1 <- ggplot(df_log_odds, aes(x = age_log, y = logit)) +</pre>
  geom_point(alpha = 0.4) +
  geom_smooth(method = "loess", color = "purple") +
  labs(title = "Logit vs Age (Log-Transformed)",
       x = "Log(Age + 1)", y = "Logit (Log Odds)") +
  theme_minimal()
plot2 <- ggplot(df_log_odds, aes(x = tumor_size_log, y = logit)) +</pre>
  geom_point(alpha = 0.4) +
  geom_smooth(method = "loess", color = "purple") +
  labs(title = "Logit vs Tumor Size (Log-Transformed)", x = "Log(Tumor Size + 1)", y = "Logit (Log Odds
  theme_minimal()
plot3 <- ggplot(df_log_odds, aes(x = rn_examined_log, y = logit)) +</pre>
  geom point(alpha = 0.4) +
  geom_smooth(method = "loess", color = "purple") +
  labs(title = "Logit vs Regional Nodes Examined (Log-Transformed)",
       x = "Log(Regional Nodes Examined + 1)", y = "Logit (Log Odds)") +
  theme_minimal()
log_odds_grid <- grid.arrange(plot1, plot2, plot3, ncol = 3)</pre>
ggsave("plots/logit_grid_plot.png", plot = log_odds_grid, width = 18, height = 6)
# Update
model_data_3 <- model_data_2 %>%
    tumor_size_log = log(tumor_size + 1),
    rn_examined_log = log(regional_node_examined + 1),
    age_log = log(age + 1)
  ) %>%
  dplyr::select(-tumor_size, -regional_node_examined, -age)
# Fit logistic regression model with log-transformed predictors
full_model <- glm(status ~ .,</pre>
                 data = model_data_3, family = binomial)
# Calculate Cook's Distance
cooksD <- cooks.distance(model_vif)</pre>
# Plot Cook's Distance
png("plots/cooks_distance_plot.png", width = 800, height = 600)
plot(cooksD, pch = 20)
abline(h = 4 / nrow(model_data_3), col = "red")
dev.off()
```

```
# Identify influential observations
influential_obs <- which(cooksD > 4 / nrow(model_data_3))
# Build Model
model_no_outliers <- glm(status ~ .,</pre>
                         data = model_data_3[-influential_obs, ], family = binomial)
model_robust <- glmrob(status ~ .,</pre>
                       data = model_data_3, family = binomial, method = "Mqle")
# Extract summaries
full_coefficients <- summary(full_model)$coefficients</pre>
no_outliers_coefficients <- summary(model_no_outliers)$coefficients</pre>
robust_coefficients <- summary(model_robust)$coefficients</pre>
# Identify unstable coefficients
unstable_coeffs <- which(
  (abs(full_coefficients[, "Estimate"] - no_outliers_coefficients[, "Estimate"]) > 2) |
  (abs(full_coefficients[, "Std. Error"] - no_outliers_coefficients[, "Std. Error"]) > 2)
unstable coeffs df <- data.frame(
  Variable = rownames(full_coefficients)[unstable_coeffs],
  Full_Model_Coefficient = full_coefficients[unstable_coeffs, "Estimate"],
  Full_Model_SE = full_coefficients[unstable_coeffs, "Std. Error"],
  No_Outliers_Coefficient = no_outliers_coefficients[unstable_coeffs, "Estimate"],
  No_Outliers_SE = no_outliers_coefficients[unstable_coeffs, "Std. Error"],
  Robust_Coefficient = robust_coefficients[rownames(full_coefficients)[unstable_coeffs], "Estimate"],
  Robust_SE = robust_coefficients[rownames(full_coefficients)[unstable_coeffs], "Std. Error"]
write.csv(unstable_coeffs_df, "tables/unstable_coefficients.csv", row.names = FALSE)
# Extract robustness weights
robust_summary <- capture.output(summary(model_robust))</pre>
start_line <- grep("Robustness weights w.r \\* w.x:", robust_summary)</pre>
end_line <- start_line + 5</pre>
desired_section <- robust_summary[start_line:end_line]</pre>
cat(paste(desired_section, collapse = "\n"))
# Forward selection
forward_model <- step(glm(status ~ ., data = model_data_3, family = binomial), scope = list(lower = ~1,
# Backward elimination
backward_model <- step(glm(status ~ ., data = model_data_3, family = binomial), direction = "backward",
# Stepwise selection
stepwise_model <- step(glm(status ~ ., data = model_data_3, family = binomial), scope = list(lower = ~1
writeLines(capture.output(summary(full_model)), "results/full_model_summary.txt")
writeLines(capture.output(summary(forward_model)), "results/forward_model_summary.txt")
writeLines(capture.output(summary(backward_model)), "results/backward_model_summary.txt")
writeLines(capture.output(summary(stepwise_model)), "results/stepwise_model_summary.txt")
best_subset <- regsubsets(`status` ~ ., data = model_data_3, nvmax = ncol(model_data_3) - 1)</pre>
best_summary <- summary(best_subset)</pre>
```

```
subset_table <- data.frame(</pre>
  Num_Predictors = 1:length(best_summary$adjr2),
  Adj_R2 = best_summary$adjr2,
 Cp = best_summary$cp,
 BIC = best_summary$bic
write.csv(subset_table, "tables/best_subset_summary.csv", row.names = FALSE)
best_adjr2_model <- which.max(best_summary$adjr2)</pre>
adjr2_predictor <- names(coef(best_subset, best_adjr2_model))[-1]</pre>
best_cp_model <- which.min(best_summary$cp)</pre>
cp_predictor <- names(coef(best_subset, best_cp_model))[-1]</pre>
best_bic_model <- which.min(best_summary$bic)</pre>
bic_predictor <- names(coef(best_subset, best_bic_model))[-1]</pre>
cat("Best Model by Adjusted R2: ", best_adjr2_model, "variables,", adjr2_predictor, "\n")
cat("Best Model by Cp: ", best_cp_model, "variables,", cp_predictor, "\n")
cat("Best Model by BIC: ", best_bic_model, "variables,", bic_predictor, "\n")
png("plots/best_subset_plots.png", width = 1800, height = 600, res = 150)
par(mfrow = c(1, 3))
plot(best_summary$adjr2,
     type = "b",
     main = "Adjusted R2 vs Model Size",
     xlab = "Number of Variables",
     ylab = "Adjusted R<sup>2</sup>")
plot(best_summary$cp,
     type = "b",
     main = "Cp vs Model Size",
     xlab = "Number of Variables",
     ylab = "Cp")
plot(best_summary$bic,
     type = "b",
     main = "BIC vs Model Size",
     xlab = "Number of Variables",
     ylab = "BIC")
par(mfrow = c(1, 1))
dev.off()
# Define predictors
predictors <- c("race", "marital_status", "x6th_stage", "grade",</pre>
                 "estrogen_status", "progesterone_status", "rn_examined_log", "age_log")
# Create a dataframe to store results
interaction results <- data.frame(</pre>
 Predictor1 = character(),
 Predictor2 = character(),
 P Value = numeric(),
  stringsAsFactors = FALSE
# Loop through each pair of predictors
for (i in seq_along(predictors)) {
 for (j in seq_along(predictors)) {
   if (i < j) {</pre>
```

```
predictor1 <- predictors[i]</pre>
      predictor2 <- predictors[j]</pre>
      # Fit interaction model
      formula <- as.formula(paste("status ~", paste(predictors, collapse = " + "),</pre>
                                    "+", predictor1, "*", predictor2))
      interaction_model <- glm(formula, family = binomial, data = model_data_3)</pre>
      # Extract interaction term
      interaction_term <- paste(predictor1, predictor2, sep = ":")</pre>
      coef_names <- names(coef(interaction_model))</pre>
      if (interaction_term %in% coef_names) {
        # Extract p-value for the interaction term
        p_value <- coef(summary(interaction_model))[interaction_term, "Pr(>|z|)"]
        # Append to results if p-value < 0.05
        if (p_value < 0.05) {</pre>
          interaction_results <- rbind(interaction_results,</pre>
                                         data.frame(Predictor1 = predictor1,
                                                    Predictor2 = predictor2,
                                                    P_Value = p_value,
                                                     stringsAsFactors = FALSE))
        }
      }
   }
 }
}
# Sort results by p-value
interaction_results <- interaction_results[order(interaction_results$P_Value), ]</pre>
print(interaction_results)
set.seed(123)
control <- trainControl(method = "cv", number = 10, classProbs = TRUE, summaryFunction = twoClassSummar</pre>
cv_model <- train(</pre>
  status ~ race + marital_status + x6th_stage + grade + estrogen_status +
    progesterone_status + rn_examined_log + age_log,
  data = model_data_3,
 method = "glm",
 family = "binomial",
 trControl = control,
 metric = "ROC"
performance_summary <- capture.output(cv_model)</pre>
writeLines(performance_summary, "results/cv_model_performance.txt")
# Split data by race group
white_data <- model_data_3 %>% filter(race == "White")
minority_data <- model_data_3 %>% filter(race != "White")
black_data <- model_data_3 %>% filter(race == "Black")
other_data <- model_data_3 %>% filter(race == "Other")
```

```
# Predict probabilities for White group
pred_white <- predict(cv_model, newdata = white_data, type = "prob")[, "Dead"]</pre>
roc white <- roc(white data$status, pred white)</pre>
auc_white <- auc(roc_white)</pre>
# Predict probabilities for Minority group
pred_minority <- predict(cv_model, newdata = minority_data, type = "prob")[, "Dead"]</pre>
roc_minority <- roc(minority_data$status, pred_minority)</pre>
auc_minority <- auc(roc_minority)</pre>
# Predict probabilities for Black group
pred_black <- predict(cv_model, newdata = black_data, type = "prob")[, "Dead"]</pre>
roc_black <- roc(black_data$status, pred_black)</pre>
auc_black <- auc(roc_black)</pre>
# Predict probabilities for Other group
pred_other <- predict(cv_model, newdata = other_data, type = "prob")[, "Dead"]</pre>
roc_other <- roc(other_data$status, pred_other)</pre>
auc_other <- auc(roc_other)</pre>
# Print AUC results
cat("ROC-AUC for White group:", auc_white, "\n")
cat("ROC-AUC for Black group:", auc_black, "\n")
cat("ROC-AUC for Other group:", auc_other, "\n")
cat("ROC-AUC for Minority group:", auc_minority, "\n")
model_data_4 <- model_data_3 %>%
  mutate(weight = case_when(
    race == "White" ~ 1,
    race == "Black" ~ 1.5,
    race == "Other" ~ 2
  ))
minority_data <- model_data_4 %>% filter(race != "White")
# Refit the model with weights
reweighted_model <- train(</pre>
  status ~ race + marital_status + x6th_stage + grade + estrogen_status +
    progesterone status + rn examined log + age log,
  data = model_data_4,
  method = "glm",
 family = "binomial",
 trControl = control,
  weights = weight
)
# Evaluate performance on subgroups
pred_white_weighted <- predict(reweighted_model, newdata = white_data, type = "prob")[, "Dead"]</pre>
roc_white_weighted <- roc(white_data$status, pred_white_weighted)</pre>
auc_white_weighted <- auc(roc_white_weighted)</pre>
pred_minority_weighted <- predict(reweighted_model, newdata = minority_data, type = "prob")[, "Dead"]</pre>
roc_minority_weighted <- roc(minority_data$status, pred_minority_weighted)</pre>
auc_minority_weighted <- auc(roc_minority_weighted)</pre>
```

```
pred_black_weighted <- predict(reweighted_model, newdata = black_data, type = "prob")[, "Dead"]</pre>
roc_black_weighted <- roc(black_data$status, pred_black_weighted)</pre>
auc_black_weighted <- auc(roc_black_weighted)</pre>
pred_other_weighted <- predict(reweighted_model, newdata = other_data, type = "prob")[, "Dead"]</pre>
roc other weighted <- roc(other data$status, pred other weighted)</pre>
auc_other_weighted <- auc(roc_other_weighted)</pre>
# Print updated results
cat("Reweighted Model - ROC-AUC for White group:", auc_white_weighted, "\n")
cat("Reweighted Model - ROC-AUC for Minority group:", auc_minority_weighted, "\n")
cat("Reweighted Model - ROC-AUC for Black group:", auc_black_weighted, "\n")
cat("Reweighted Model - ROC-AUC for Other group:", auc_other_weighted, "\n")
# Extract coefficients and confidence intervals from the reweighted model
summary_reweighted <- summary(reweighted_model$finalModel)</pre>
coefficients <- coef(summary_reweighted)</pre>
# Calculate Odds Ratios and 95% Confidence Intervals
odds ratios <- exp(coefficients[, "Estimate"])</pre>
lower_ci <- exp(coefficients[, "Estimate"] - 1.96 * coefficients[, "Std. Error"])</pre>
upper_ci <- exp(coefficients[, "Estimate"] + 1.96 * coefficients[, "Std. Error"])
# Combine into a table
results_table <- data.frame(</pre>
 Predictor = rownames(coefficients),
  Estimate = round(coefficients[, "Estimate"], 3),
  Std_Error = round(coefficients[, "Std. Error"], 3),
 Odds_Ratio = round(odds_ratios, 3),
 '95% CI (Lower)' = round(lower_ci, 3),
  '95% CI (Upper)' = round(upper_ci, 3),
  `P-Value` = format.pval(coefficients[, "Pr(>|z|)"], digits = 3)
# Clean up row names
rownames(results table) <- NULL</pre>
write.csv(results_table, "tables/reweighted_model_results.csv", row.names = FALSE)
skim categorical <- read.csv("tables/skim categorical summary.csv")</pre>
knitr::kable(skim_categorical, caption = "Skim Summary for Categorical Variables")
skim_numeric <- read.csv("tables/skim_numeric_summary.csv")</pre>
knitr::kable(skim_numeric, caption = "Skim Summary for Numeric Variables")
summary_by_status <- read.csv("tables/summary_by_status.csv")</pre>
knitr::kable(summary_by_status, caption = "Summary Statistics Grouped by Survival Status")
vif_table <- read.csv("tables/vif.csv")</pre>
knitr::kable(vif_table, caption = "Variance Inflation Factors for Predictors")
best_subset_summary <- read.csv("tables/best_subset_summary.csv")</pre>
knitr::kable(best_subset_summary, caption = "Best Subset Selection Summary")
```

reweighted\_model\_results <- read.csv("tables/reweighted\_model\_results.csv")
knitr::kable(reweighted\_model\_results, caption = "Reweighted Logistic Regression Model Results")</pre>