

# P8130\_final\_project

Leonor Rui

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## Appendix

- Data Import

```
survival_df = read_csv("data/Project_2_data.csv") |>
  janitor::clean_names()
```

```
## Rows: 4024 Columns: 16
## -- Column specification -----
## Delimiter: ","
## chr (11): Race, Marital Status, T Stage, N Stage, 6th Stage, differentiate, ...
## dbl (5): Age, Tumor Size, Regional Node Examined, Reginol Node Positive, Su...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
```

- Data Description

```
str(survival_df)
```

```
## spc_tbl_ [4,024 x 16] (S3: spec_tbl_df/tbl_df/tbl/data.frame)
## $ age : num [1:4024] 68 50 58 58 47 51 51 40 40 69 ...
## $ race : chr [1:4024] "White" "White" "White" "White" ...
## $ marital_status : chr [1:4024] "Married" "Married" "Divorced" "Married" ...
## $ t_stage : chr [1:4024] "T1" "T2" "T3" "T1" ...
## $ n_stage : chr [1:4024] "N1" "N2" "N3" "N1" ...
## $ x6th_stage : chr [1:4024] "IIA" "IIIA" "IIIC" "IIA" ...
## $ differentiate : chr [1:4024] "Poorly differentiated" "Moderately differentiated" "Moderat
## $ grade : chr [1:4024] "3" "2" "2" "3" ...
## $ a_stage : chr [1:4024] "Regional" "Regional" "Regional" "Regional" ...
## $ tumor_size : num [1:4024] 4 35 63 18 41 20 8 30 103 32 ...
## $ estrogen_status : chr [1:4024] "Positive" "Positive" "Positive" "Positive" ...
## $ progesterone_status : chr [1:4024] "Positive" "Positive" "Positive" "Positive" ...
## $ regional_node_examined: num [1:4024] 24 14 14 2 3 18 11 9 20 21 ...
## $ reginol_node_positive : num [1:4024] 1 5 7 1 1 2 1 1 18 12 ...
## $ survival_months : num [1:4024] 60 62 75 84 50 89 54 14 70 92 ...
## $ status : chr [1:4024] "Alive" "Alive" "Alive" "Alive" ...
## - attr(*, "spec")=
## .. cols(
```

```
## .. Age = col_double(),
## .. Race = col_character(),
## .. 'Marital Status' = col_character(),
## .. 'T Stage' = col_character(),
## .. 'N Stage' = col_character(),
## .. '6th Stage' = col_character(),
## .. differentiate = col_character(),
## .. Grade = col_character(),
## .. 'A Stage' = col_character(),
## .. 'Tumor Size' = col_double(),
## .. 'Estrogen Status' = col_character(),
## .. 'Progesterone Status' = col_character(),
## .. 'Regional Node Examined' = col_double(),
## .. 'Reginol Node Positive' = col_double(),
## .. 'Survival Months' = col_double(),
## .. Status = col_character()
## .. )
## - attr(*, "problems")=<externalptr>
```

Numeric variables include `age`, `tumor_size`, `regional_node_examined`, `reginol_node_positive`, and `survival_months`.

These are continuous variables that can be used for our later regression analysis.

Categorical variables include `race`, `marital_status`, `t_stage`, `n_stage`, `x6th_stage`, `differentiate`, `grade`, `a_stage`, `estrogen_status`, `progesterone_status`, and `status`.

These can be converted into factors for later analysis.

```
summary(survival_df)
```

```
##      age      race      marital_status      t_stage
## Min.   :30.00  Length:4024      Length:4024      Length:4024
## 1st Qu.:47.00  Class :character  Class :character  Class :character
## Median :54.00  Mode  :character  Mode  :character  Mode  :character
## Mean   :53.97
## 3rd Qu.:61.00
## Max.   :69.00
##      n_stage      x6th_stage      differentiate      grade
## Length:4024      Length:4024      Length:4024      Length:4024
## Class :character  Class :character  Class :character  Class :character
## Mode  :character  Mode  :character  Mode  :character  Mode  :character
##
##
##      a_stage      tumor_size      estrogen_status      progesterone_status
## Length:4024      Min.   : 1.00      Length:4024      Length:4024
## Class :character  1st Qu.: 16.00      Class :character  Class :character
## Mode  :character  Median : 25.00      Mode  :character  Mode  :character
##                      Mean    : 30.47
##                      3rd Qu.: 38.00
##                      Max.    :140.00
## regional_node_examined regional_node_positive survival_months
## Min.   : 1.00      Min.   : 1.000      Min.   : 1.0
## 1st Qu.: 9.00      1st Qu.: 1.000      1st Qu.: 56.0
```

```
## Median :14.00      Median : 2.000      Median : 73.0
## Mean   :14.36      Mean    : 4.158      Mean    : 71.3
## 3rd Qu.:19.00      3rd Qu.: 5.000      3rd Qu.: 90.0
## Max.   :61.00      Max.    :46.000      Max.    :107.0
##      status
## Length:4024
## Class :character
## Mode  :character
##
##
##
```

The wide range of values in variables such as `tumor_size`, `regional_node_examined`, and `survival_months` indicates the need to explore relationships and their potential nonlinearities with survival, giving us a possible analytical regression model.

```
colSums(is.na(survival_df))
```

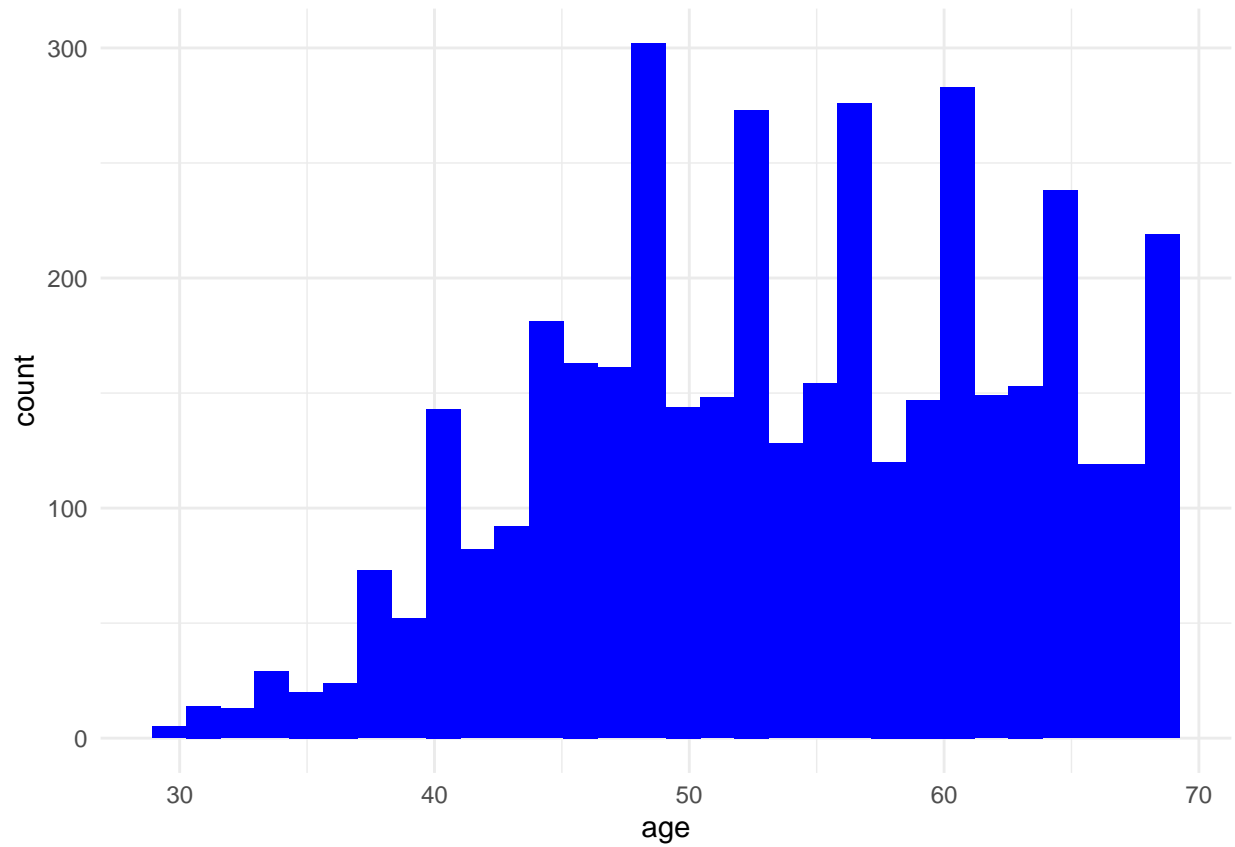
```
##           age           race      marital_status
##           0           0           0
##      t_stage      n_stage      x6th_stage
##           0           0           0
## differentiate      grade      a_stage
##           0           0           0
##      tumor_size      estrogen_status      progesterone_status
##           0           0           0
## regional_node_examined      reginol_node_positive      survival_months
##           0           0           0
##           status
##           0
```

We can conclude that no missing values are present in this dataset across all variables.

- Data Visualization

```
survival_df |>
  ggplot(aes(age)) +
  geom_histogram(fill = "blue") +
  theme_minimal()
```

```
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
```



The histogram shows the age distribution of patients. Most patients are aged between 40 and 70 years. The data is well spread across middle and older age groups, making it possible for age-related analysis. Therefore, age will likely be a significant predictor for later analysis.