BST210 HW4 Project Check-In 1

Group 7: Regression Heroes

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### 1. What is the general domain/subject area of this project?

The general domain/subject area of this project is cancer prevalence with respect to socioeconomic status in the United States.

### 2. What data will you use, and what is the source?

We are using data from “OLS Regression Challenge - dataset by nrippner | data.world”, which aggregates socioeconomic and clinical data from census.gov, clinicaltrials.gov, and cancer.gov. These cancer outcomes data were aggregated from cancer trials during 01/01/2010 through 06/01/2016 and socioeconomic and demographic data were aggregated from 2013 U.S. census data.

Source: <https://data.world/nrippner/ols-regression-challenge>

### 3. What primary questions will you seek to answer?

We are trying to explore the relationship between cancer-related death and the socioeconomic status of US counties. We will use predictors such as median income, age, and employment status to create a model that could predict the number of cancer diagnoses.

### 4. What secondary questions will you seek to answer?

Some secondary questions we are considering revolve around analysis of demographic data from the dataset. We are proposing to explore the relationship between race, income, and available medical resources in each county.

### 5. What outcome(s)/endpoint(s) will you use? (could be continuous, binary, polytomous, Poisson, survival,…and you may be considering more than one–and this may be updated/added to, as the semester progresses)

Our main outcome will be the cancer-related death rates in US counties from 2010 to 2016. This outcome is transformed from discrete count (count of cancer-related deaths) type, and we will use poisson distribution to model. For our secondary questions, we might

### 6. What is your draft Statistical Analysis Plan? (should be a very thorough, detailed, bullet point outline, demonstrating that you have broadly thought this through and included details - this may be updated of course as the semester proceeds, and with feedback) \*\* Note that we will be discussing all forms of outcome/endpoint data in this course, and at present have not yet covered each of these…so this plan may be updated/added to as the semester progresses, but you still should be able to plan out the structure and significant details of your plan. If your outcome data is other than continuous, you can still include for instance ‘Regression modeling involving ’Y’ outcome data of interest, involving these variables (list them)…’ and any other concerns or methods of interest (listing potential confounders, effect modifiers, potential use of splines or additive modeling, potential missing data considerations, data reduction methods, regularization methods, etc,…or none of these–you will want to consider what is most appropriate for your data and questions at hand). Recall the BST 210 Regression Models Overview Table from which most extensions arise.

**Exploratory Data Analysis:**

* Better understand the structure and complexity of our data set through str and summary functions in R
* Check for missing data points categorized by different covariates (by employment status of different age groups)
* Explore data patterns and modeling – Check for correlation between different socioeconomic status determining variables and cancer prevalence through a pairs plot.
  + One of the variables of interest of our project is the effect of region/state on cancer prevalence. Graph this relationship to observe for any patterns.

**Primary Inference Problem:**

* Poisson regression (within a generalized linear framework) will be our main modeling approach for statistical inference due to our primary outcome being rate/count data
* With 33 covariates of interest, we will need to employ model selection techniques, potentially finding a good in-between for various automated methods and integrating domain information
* Once we have identified some covariates of interest, we will perform various smoothing methods in order to determine the relative correlations in data
* Data cleaning and computing additional predictors (e.g., regions, climate … )
* As our data is on the county-level, we aim to add a dimension to our analysis by aggregating these data onto the state-level, which would allow us to study state-by-state differences in socioeconomic conditions and cancer outcomes
* Create comprehensive visualization of our demographic data to illustrate differences between US counties and states
* We will explore different imputation techniques to adjust for the 3046 incomplete observations in these data
* Identify and adjust for potential confounders (on the association of SES to cancer prevalence) such as, but not limited to, county and income.
* For any of these covariates which we are not interested in interpreting, we aim to adjust for them flexibly using splines/GAMs (and if not needed, we will keep using linear terms)

**Secondary Inference Problem:**

* We are potentially interested in looking at state-level and county-level differences in demographics and insurance status
* Here we will employ multinomial models in order to model the probability of some set of socioeconomic conditions and cancer outcomes being in a given county or state

**Predictive Problem:**

* Aside from the inference problem, we are also interested in the prediction problem and if there is time, we will explore the performance of our generalized linear models on prediction metrics (accuracy, AUC, etc.)
* We will also explore how penalized models (LASSO, Ridge, Elastic Net) and traditional ML algorithms (random forests/basic nnets for primary analysis, RFs/KNN/nnets/etc. for secondary analysis tasks) compare with the inference models we previously built
* Here we are also interested in exploring how the number of predictors can correlate with generalizability in these 3 forms of models

### 7. What are the biggest challenges you foresee in answering your proposed questions and completing this project? (logistical, statistical, etc, if there are any)

A challenge we could face is that our data is predominantly collected on the caucasian population rather than populations of ethnic minority groups. The lack of data of these underrepresented groups may lead to potential bias in our findings. The data on socioeconomic status is not stratified on ethnicity, making it difficult to determine outcomes by race. Additionally, a lot of the data is in percentages. Therefore, when we convert the data into counts, the data may not be independent between different variables.

### 8. Will you seek domain expertise? Why or why not? If so, from whom?

No, we will not seek domain expertise because the data is fairly general to all cancer cases. All our group members have some background in life sciences, molecular biology, and/or economics, which we believe is sufficient for the scope of the project. However, if need be, we can further scope the literature for additional information on cancer biology or economic terminology.

### 9. What software package(s) will you use to complete this project? (It is absolutely fine for different group members to use different packages; in fact, some tasks are easier in some packages over others and vice versa.)

We will use the R statistical programming language to analyze these data and complete this project. We plan on using tidyverse for transforming and cleaning data. Furthermore, this library of packages allows us to create effective EDA plots in order to explore our data efficiently. For modeling, we plan on using the lm and glm functions in base R for standard linear regression and Poisson regression (or other generalized models we may approach later), respectively.

### 10. Complete an initial round of exploratory analyses on your data that would be relevant to your plan and responses above, and include any plots, summaries, code and output. Please include exploratory analysis for outcome(s) of continuous form however/wherever possible even if your ultimate goals/questions involve a different form of outcome data such as binary, polytomous, etc. (You may consider this initial analysis as a potential sub-analysis later on.)

#### Exploratory Data Analysis

library(tidyverse)

## ── Attaching packages ─────────────────────────────────────── tidyverse 1.3.2 ──  
## ✔ ggplot2 3.3.6 ✔ purrr 0.3.4   
## ✔ tibble 3.1.8 ✔ dplyr 1.0.10  
## ✔ tidyr 1.2.0 ✔ stringr 1.4.1   
## ✔ readr 2.1.2 ✔ forcats 0.5.2   
## ── Conflicts ────────────────────────────────────────── tidyverse\_conflicts() ──  
## ✖ dplyr::filter() masks stats::filter()  
## ✖ dplyr::lag() masks stats::lag()

library(stringr)  
library(viridisLite) # nice colours

##### What’s the structure of our data?

dat <- read\_csv("cancer\_reg.csv")

## Rows: 3047 Columns: 34  
## ── Column specification ────────────────────────────────────────────────────────  
## Delimiter: ","  
## chr (2): binnedInc, Geography  
## dbl (32): avgAnnCount, avgDeathsPerYear, TARGET\_deathRate, incidenceRate, me...  
##   
## 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.

str(dat)

## spec\_tbl\_df [3,047 × 34] (S3: spec\_tbl\_df/tbl\_df/tbl/data.frame)  
## $ avgAnnCount : num [1:3047] 1397 173 102 427 57 ...  
## $ avgDeathsPerYear : num [1:3047] 469 70 50 202 26 152 97 71 36 1380 ...  
## $ TARGET\_deathRate : num [1:3047] 165 161 175 195 144 ...  
## $ incidenceRate : num [1:3047] 490 412 350 430 350 ...  
## $ medIncome : num [1:3047] 61898 48127 49348 44243 49955 ...  
## $ popEst2015 : num [1:3047] 260131 43269 21026 75882 10321 ...  
## $ povertyPercent : num [1:3047] 11.2 18.6 14.6 17.1 12.5 15.6 23.2 17.8 22.3 13.1 ...  
## $ studyPerCap : num [1:3047] 499.7 23.1 47.6 342.6 0 ...  
## $ binnedInc : chr [1:3047] "(61494.5, 125635]" "(48021.6, 51046.4]" "(48021.6, 51046.4]" "(42724.4, 45201]" ...  
## $ MedianAge : num [1:3047] 39.3 33 45 42.8 48.3 45.4 42.6 51.7 49.3 35.8 ...  
## $ MedianAgeMale : num [1:3047] 36.9 32.2 44 42.2 47.8 43.5 42.2 50.8 48.4 34.7 ...  
## $ MedianAgeFemale : num [1:3047] 41.7 33.7 45.8 43.4 48.9 48 43.5 52.5 49.8 37 ...  
## $ Geography : chr [1:3047] "Kitsap County, Washington" "Kittitas County, Washington" "Klickitat County, Washington" "Lewis County, Washington" ...  
## $ AvgHouseholdSize : num [1:3047] 2.54 2.34 2.62 2.52 2.34 2.58 2.42 2.24 2.38 2.65 ...  
## $ PercentMarried : num [1:3047] 52.5 44.5 54.2 52.7 57.8 50.4 54.1 52.7 55.9 50 ...  
## $ PctNoHS18\_24 : num [1:3047] 11.5 6.1 24 20.2 14.9 29.9 26.1 27.3 34.7 15.6 ...  
## $ PctHS18\_24 : num [1:3047] 39.5 22.4 36.6 41.2 43 35.1 41.4 33.9 39.4 36.3 ...  
## $ PctSomeCol18\_24 : num [1:3047] 42.1 64 NA 36.1 40 NA NA 36.5 NA NA ...  
## $ PctBachDeg18\_24 : num [1:3047] 6.9 7.5 9.5 2.5 2 4.5 5.8 2.2 1.4 7.1 ...  
## $ PctHS25\_Over : num [1:3047] 23.2 26 29 31.6 33.4 30.4 29.8 31.6 32.2 28.8 ...  
## $ PctBachDeg25\_Over : num [1:3047] 19.6 22.7 16 9.3 15 11.9 11.9 11.3 12 16.2 ...  
## $ PctEmployed16\_Over : num [1:3047] 51.9 55.9 45.9 48.3 48.2 44.1 51.8 40.9 39.5 56.6 ...  
## $ PctUnemployed16\_Over : num [1:3047] 8 7.8 7 12.1 4.8 12.9 8.9 8.9 10.3 9.2 ...  
## $ PctPrivateCoverage : num [1:3047] 75.1 70.2 63.7 58.4 61.6 60 49.5 55.8 55.5 69.9 ...  
## $ PctPrivateCoverageAlone: num [1:3047] NA 53.8 43.5 40.3 43.9 38.8 35 33.1 37.8 NA ...  
## $ PctEmpPrivCoverage : num [1:3047] 41.6 43.6 34.9 35 35.1 32.6 28.3 25.9 29.9 44.4 ...  
## $ PctPublicCoverage : num [1:3047] 32.9 31.1 42.1 45.3 44 43.2 46.4 50.9 48.1 31.4 ...  
## $ PctPublicCoverageAlone : num [1:3047] 14 15.3 21.1 25 22.7 20.2 28.7 24.1 26.6 16.5 ...  
## $ PctWhite : num [1:3047] 81.8 89.2 90.9 91.7 94.1 ...  
## $ PctBlack : num [1:3047] 2.595 0.969 0.74 0.783 0.27 ...  
## $ PctAsian : num [1:3047] 4.822 2.246 0.466 1.161 0.666 ...  
## $ PctOtherRace : num [1:3047] 1.843 3.741 2.747 1.363 0.492 ...  
## $ PctMarriedHouseholds : num [1:3047] 52.9 45.4 54.4 51 54 ...  
## $ BirthRate : num [1:3047] 6.12 4.33 3.73 4.6 6.8 ...  
## - attr(\*, "spec")=  
## .. cols(  
## .. avgAnnCount = col\_double(),  
## .. avgDeathsPerYear = col\_double(),  
## .. TARGET\_deathRate = col\_double(),  
## .. incidenceRate = col\_double(),  
## .. medIncome = col\_double(),  
## .. popEst2015 = col\_double(),  
## .. povertyPercent = col\_double(),  
## .. studyPerCap = col\_double(),  
## .. binnedInc = col\_character(),  
## .. MedianAge = col\_double(),  
## .. MedianAgeMale = col\_double(),  
## .. MedianAgeFemale = col\_double(),  
## .. Geography = col\_character(),  
## .. AvgHouseholdSize = col\_double(),  
## .. PercentMarried = col\_double(),  
## .. PctNoHS18\_24 = col\_double(),  
## .. PctHS18\_24 = col\_double(),  
## .. PctSomeCol18\_24 = col\_double(),  
## .. PctBachDeg18\_24 = col\_double(),  
## .. PctHS25\_Over = col\_double(),  
## .. PctBachDeg25\_Over = col\_double(),  
## .. PctEmployed16\_Over = col\_double(),  
## .. PctUnemployed16\_Over = col\_double(),  
## .. PctPrivateCoverage = col\_double(),  
## .. PctPrivateCoverageAlone = col\_double(),  
## .. PctEmpPrivCoverage = col\_double(),  
## .. PctPublicCoverage = col\_double(),  
## .. PctPublicCoverageAlone = col\_double(),  
## .. PctWhite = col\_double(),  
## .. PctBlack = col\_double(),  
## .. PctAsian = col\_double(),  
## .. PctOtherRace = col\_double(),  
## .. PctMarriedHouseholds = col\_double(),  
## .. BirthRate = col\_double()  
## .. )  
## - attr(\*, "problems")=<externalptr>

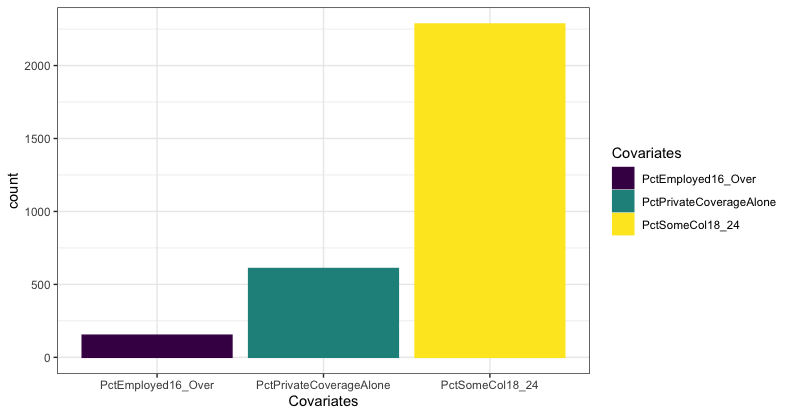
summary(dat)

## avgAnnCount avgDeathsPerYear TARGET\_deathRate incidenceRate   
## Min. : 6.0 Min. : 3 Min. : 59.7 Min. : 201.3   
## 1st Qu.: 76.0 1st Qu.: 28 1st Qu.:161.2 1st Qu.: 420.3   
## Median : 171.0 Median : 61 Median :178.1 Median : 453.5   
## Mean : 606.3 Mean : 186 Mean :178.7 Mean : 448.3   
## 3rd Qu.: 518.0 3rd Qu.: 149 3rd Qu.:195.2 3rd Qu.: 480.9   
## Max. :38150.0 Max. :14010 Max. :362.8 Max. :1206.9   
##   
## medIncome popEst2015 povertyPercent studyPerCap   
## Min. : 22640 Min. : 827 Min. : 3.20 Min. : 0.00   
## 1st Qu.: 38882 1st Qu.: 11684 1st Qu.:12.15 1st Qu.: 0.00   
## Median : 45207 Median : 26643 Median :15.90 Median : 0.00   
## Mean : 47063 Mean : 102637 Mean :16.88 Mean : 155.40   
## 3rd Qu.: 52492 3rd Qu.: 68671 3rd Qu.:20.40 3rd Qu.: 83.65   
## Max. :125635 Max. :10170292 Max. :47.40 Max. :9762.31   
##   
## binnedInc MedianAge MedianAgeMale MedianAgeFemale  
## Length:3047 Min. : 22.30 Min. :22.40 Min. :22.30   
## Class :character 1st Qu.: 37.70 1st Qu.:36.35 1st Qu.:39.10   
## Mode :character Median : 41.00 Median :39.60 Median :42.40   
## Mean : 45.27 Mean :39.57 Mean :42.15   
## 3rd Qu.: 44.00 3rd Qu.:42.50 3rd Qu.:45.30   
## Max. :624.00 Max. :64.70 Max. :65.70   
##   
## Geography AvgHouseholdSize PercentMarried PctNoHS18\_24   
## Length:3047 Min. :0.0221 Min. :23.10 Min. : 0.00   
## Class :character 1st Qu.:2.3700 1st Qu.:47.75 1st Qu.:12.80   
## Mode :character Median :2.5000 Median :52.40 Median :17.10   
## Mean :2.4797 Mean :51.77 Mean :18.22   
## 3rd Qu.:2.6300 3rd Qu.:56.40 3rd Qu.:22.70   
## Max. :3.9700 Max. :72.50 Max. :64.10   
##   
## PctHS18\_24 PctSomeCol18\_24 PctBachDeg18\_24 PctHS25\_Over   
## Min. : 0.0 Min. : 7.10 Min. : 0.000 Min. : 7.50   
## 1st Qu.:29.2 1st Qu.:34.00 1st Qu.: 3.100 1st Qu.:30.40   
## Median :34.7 Median :40.40 Median : 5.400 Median :35.30   
## Mean :35.0 Mean :40.98 Mean : 6.158 Mean :34.80   
## 3rd Qu.:40.7 3rd Qu.:46.40 3rd Qu.: 8.200 3rd Qu.:39.65   
## Max. :72.5 Max. :79.00 Max. :51.800 Max. :54.80   
## NA's :2285   
## PctBachDeg25\_Over PctEmployed16\_Over PctUnemployed16\_Over PctPrivateCoverage  
## Min. : 2.50 Min. :17.60 Min. : 0.400 Min. :22.30   
## 1st Qu.: 9.40 1st Qu.:48.60 1st Qu.: 5.500 1st Qu.:57.20   
## Median :12.30 Median :54.50 Median : 7.600 Median :65.10   
## Mean :13.28 Mean :54.15 Mean : 7.852 Mean :64.35   
## 3rd Qu.:16.10 3rd Qu.:60.30 3rd Qu.: 9.700 3rd Qu.:72.10   
## Max. :42.20 Max. :80.10 Max. :29.400 Max. :92.30   
## NA's :152   
## PctPrivateCoverageAlone PctEmpPrivCoverage PctPublicCoverage  
## Min. :15.70 Min. :13.5 Min. :11.20   
## 1st Qu.:41.00 1st Qu.:34.5 1st Qu.:30.90   
## Median :48.70 Median :41.1 Median :36.30   
## Mean :48.45 Mean :41.2 Mean :36.25   
## 3rd Qu.:55.60 3rd Qu.:47.7 3rd Qu.:41.55   
## Max. :78.90 Max. :70.7 Max. :65.10   
## NA's :609   
## PctPublicCoverageAlone PctWhite PctBlack PctAsian   
## Min. : 2.60 Min. : 10.20 Min. : 0.0000 Min. : 0.0000   
## 1st Qu.:14.85 1st Qu.: 77.30 1st Qu.: 0.6207 1st Qu.: 0.2542   
## Median :18.80 Median : 90.06 Median : 2.2476 Median : 0.5498   
## Mean :19.24 Mean : 83.65 Mean : 9.1080 Mean : 1.2540   
## 3rd Qu.:23.10 3rd Qu.: 95.45 3rd Qu.:10.5097 3rd Qu.: 1.2210   
## Max. :46.60 Max. :100.00 Max. :85.9478 Max. :42.6194   
##   
## PctOtherRace PctMarriedHouseholds BirthRate   
## Min. : 0.0000 Min. :22.99 Min. : 0.000   
## 1st Qu.: 0.2952 1st Qu.:47.76 1st Qu.: 4.521   
## Median : 0.8262 Median :51.67 Median : 5.381   
## Mean : 1.9835 Mean :51.24 Mean : 5.640   
## 3rd Qu.: 2.1780 3rd Qu.:55.40 3rd Qu.: 6.494   
## Max. :41.9303 Max. :78.08 Max. :21.326   
##

From the summary above, we can observe that we have a lot of variables to consider to answer our main questions. Additionally, it tells us about the complexity about the data itself in that cancer prevalence is a factor of many socioeconomic factors.

##### How much missing data is there?

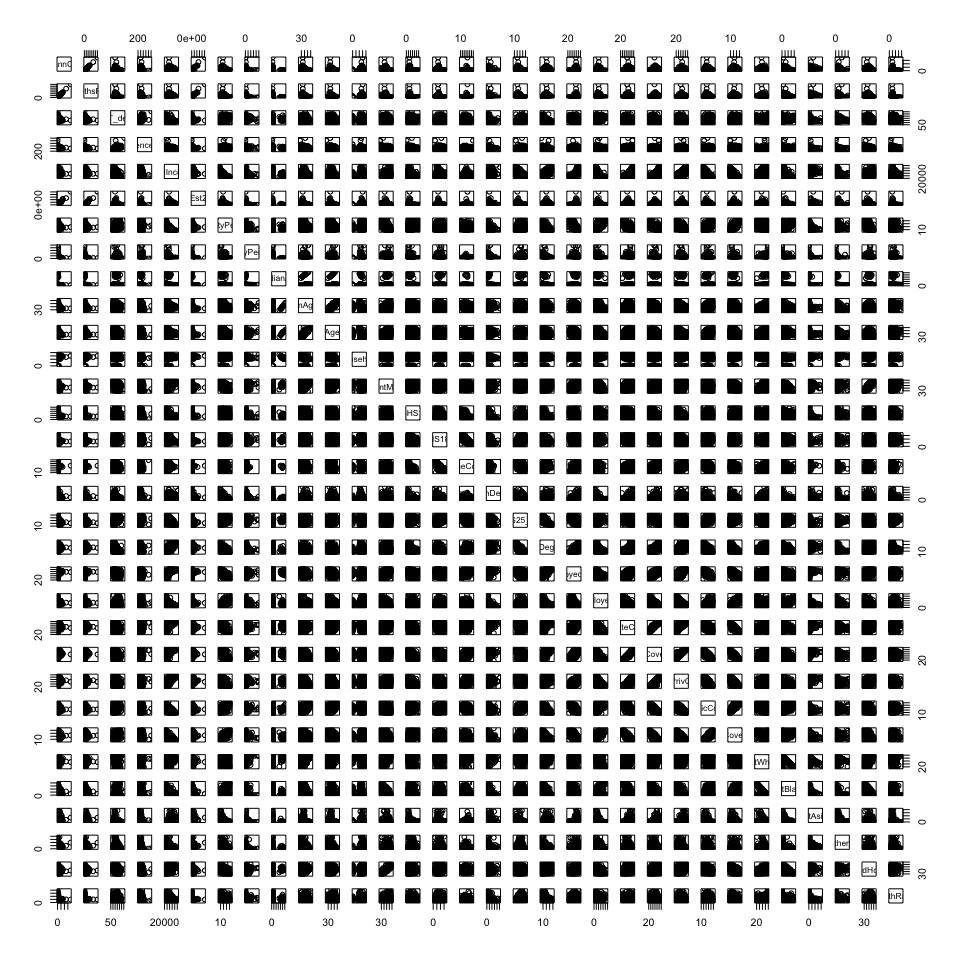
no\_geodat <- dat %>% select(-c("Geography", "binnedInc")) # no missing counties  
no\_geodat[!complete.cases(no\_geodat),] %>% # keep rows with NAs  
 pivot\_longer(colnames(no\_geodat), names\_to = "Covariates", values\_to = "Values") %>% # pivot into long table  
 filter(is.na(Values)) %>% # filter out all the non-na's  
 ggplot(aes(x = Covariates)) +   
 geom\_bar(position = "dodge", aes(col = Covariates, fill = Covariates)) +   
 theme\_bw() +   
 scale\_fill\_viridis\_d() +   
 scale\_color\_viridis\_d()



Interestingly, there is some missing data in the reporting of county employment for residents age 16 and over. We observe more missing data in percentages of county residents with only private healthcare coverage, and missing data in the majority of counties for reports of the percent of county residents between 18 and 24 years old with with some college as their highest attained education. One hypothesis for why there may be a large amount of missing data in educational reporting for 18 to 24 year olds with some college may be because some counties may not have high emphasis on higher education and many individuals in this age range that are pursuing college will probably be in more college-oriented counties.

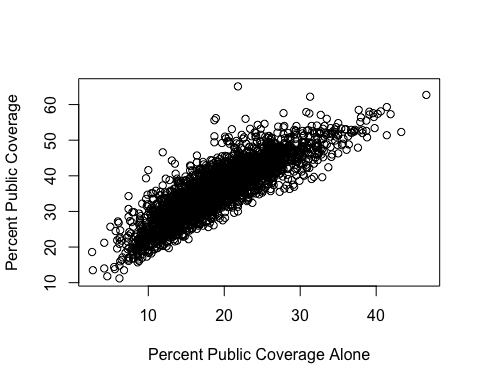
##### Exploring data patterns and modeling

pairs(dat %>% select(where(is.numeric)))



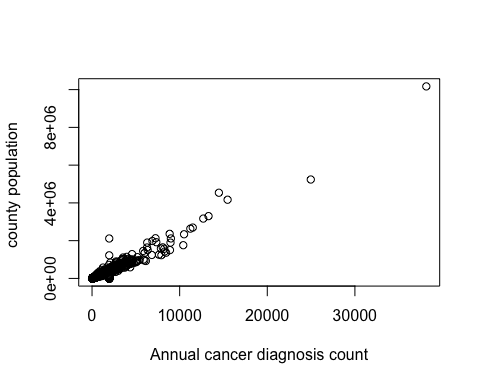
However, showing all of the variables is somewhat ineffective in showing the the relationship between the variables. A few variables that are strongly correlated are singled out for better visibility.

plot(dat$PctPublicCoverageAlone, dat$PctPublicCoverage, xlab = "Percent Public Coverage Alone", ylab = "Percent Public Coverage")



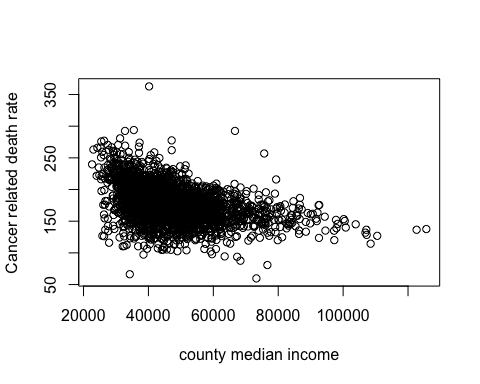
It is self explanatory that the higher the percentage of the county population is under public healthcare, the higher the percentage of the county single persons is under public healthcare.

plot(dat$avgAnnCount, dat$popEst2015, xlab = "Annual cancer diagnosis count", ylab = "county population ")



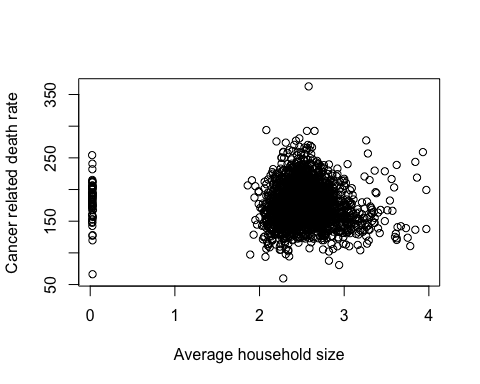
This is another expected correlation. The higher the population is, the more cancer occurance will be in the population.

plot(dat$medIncome,dat$TARGET\_deathRate, ylab = "Cancer related death rate", xlab = "county median income")



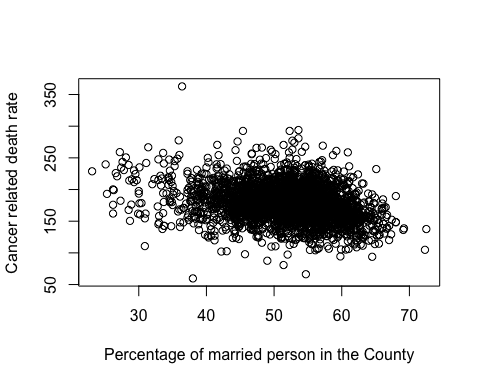
This is an example of significant predictors for our model. Annual income is likely very correlated with the # of cancer related death rate. However, we need to further explore what kind of model will be best fit for the data.

plot(dat$AvgHouseholdSize,dat$TARGET\_deathRate, ylab = "Cancer related death rate", xlab = "Average household size")



As the plot suggests, there isn’t a clear relationship between cancer-related death rate and the average household size. However, furtherwork could help illustrate some latent meanings.

plot(dat$PercentMarried,dat$TARGET\_deathRate, ylab = "Cancer related death rate", xlab = "Percentage of married person in the County")



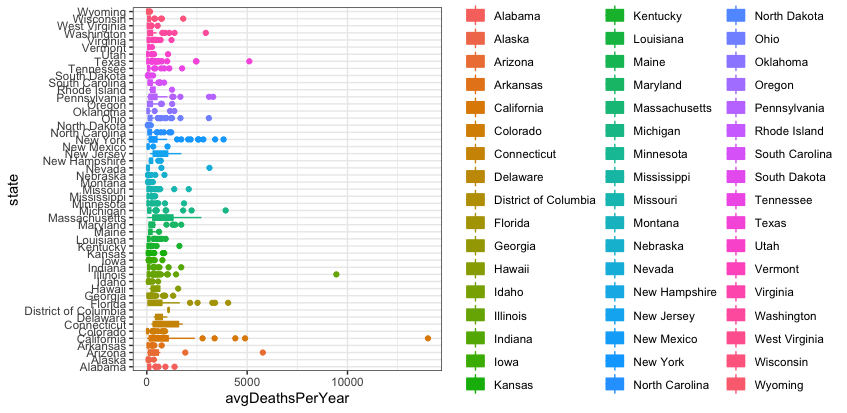
It seems that there is a negative association between percent of people married and cancer-related death rate. This seems to be another valuable predictor that we could use in our model.

These graphs are a brief look into the possible associations within our data set. We need to be careful that some of these relationships might be confounded by other co-variates, We also need to conduct confounding analysis (both classical, and statistical), effect modifier analysis to figure out the nature of the associations.

##### Look at the state-wide view

dat <- dat %>%  
 mutate(state = str\_extract(Geography, "[^,]+$")) %>% # regex to select everything after ','  
 mutate(state = str\_trim(state))

dat %>%  
 ggplot(aes(x = state, y = avgDeathsPerYear, colour = state, fill = state)) +  
 geom\_boxplot() +   
 theme\_bw() +   
 coord\_flip()



It appears that there are some potential outliers when studying the relationship between states and average deaths per year due to cancer. Particularly, there may be potential outliers in the data for California, Illinois, and Arizona. We would need to further examine the reasons for such a large deviation from the rest of the data. Additionally, it appears that the more populated states, such as California and New York, have a higher average death rate compared to less populated states like Wyoming.

### 11.

*We attest that no member of this group is using these data or same/similar questions in any other course or course project, at HSPH.*

*Hongkai Wang, Stella Nam, Ryan Wang*