Data Manipulation, EDA, Statistical Learning Tools

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```
#Load, clean, manipulate, and tidy the data
# Import data
cancer_raw = readr::read_csv("./Data/Cancer_Registry.csv") %>%
  janitor::clean_names()
dim(cancer_raw)
## [1] 3047
head(cancer raw)
## # A tibble: 6 x 34
     avg_ann_count avg_deaths_per_~ target_death_ra~ incidence_rate med_income
##
             <dbl>
                               <dbl>
                                                 <dbl>
                                                                 <dbl>
                                                                            <dbl>
## 1
              1397
                                 469
                                                  165.
                                                                  490.
                                                                            61898
## 2
               173
                                  70
                                                  161.
                                                                  412.
                                                                            48127
                                  50
               102
                                                  175.
                                                                  350.
                                                                            49348
## 4
               427
                                 202
                                                  195.
                                                                  430.
                                                                            44243
## 5
                57
                                  26
                                                  144.
                                                                  350.
                                                                            49955
               428
## 6
                                 152
                                                  176
                                                                  505.
                                                                            52313
     ... with 29 more variables: pop_est2015 <dbl>, poverty_percent <dbl>,
## #
       study_per_cap <dbl>, binned_inc <chr>, median_age <dbl>,
       median_age_male <dbl>, median_age_female <dbl>, geography <chr>,
## #
## #
       avg_household_size <dbl>, percent_married <dbl>, pct_no_hs18_24 <dbl>,
## #
       pct_hs18_24 <dbl>, pct_some_col18_24 <dbl>, pct_bach_deg18_24 <dbl>,
## #
       pct_hs25_over <dbl>, pct_bach_deg25_over <dbl>,
## #
       pct_employed16_over <dbl>, pct_unemployed16_over <dbl>,
## #
       pct_private_coverage <dbl>, pct_private_coverage_alone <dbl>,
## #
       pct_emp_priv_coverage <dbl>, pct_public_coverage <dbl>,
## #
       pct_public_coverage_alone <dbl>, pct_white <dbl>, pct_black <dbl>,
## #
       pct_asian <dbl>, pct_other_race <dbl>, pct_married_households <dbl>,
       birth_rate <dbl>
# Check NA values for each column
n_NA = sapply(cancer_raw[1:34], function(x) sum(length(which(is.na(x)))))
n_NA
##
                                      avg_deaths_per_year
                avg_ann_count
##
##
            target_death_rate
                                            incidence_rate
##
##
                                               pop_est2015
                    med_income
##
##
              poverty_percent
                                             study_per_cap
##
##
                    binned_inc
                                                median_age
##
##
              median_age_male
                                        median_age_female
```

```
##
                                                         0
##
                     geography
                                       avg_household_size
##
##
                                           pct_no_hs18_24
              percent_married
##
                  pct_hs18_24
                                        pct_some_col18_24
##
                                                      2285
##
##
            pct_bach_deg18_24
                                             pct_hs25_over
##
##
          pct_bach_deg25_over
                                      pct_employed16_over
##
                                                       152
##
        pct_unemployed16_over
                                     pct_private_coverage
##
   pct_private_coverage_alone
##
                                    pct_emp_priv_coverage
##
                           609
##
          pct_public_coverage
                                pct_public_coverage_alone
##
                                                         0
                             0
##
                                                 pct_black
                     pct_white
##
                             0
                                                         0
##
                     pct_asian
                                           pct_other_race
##
                                                         0
##
       pct_married_households
                                                birth_rate
##
# Check the percentage of NA values for each column
percentage_NA = sapply(cancer_raw[1:34], function(x) sum(length(which(is.na(x)))) / 3047)
percentage_NA %>% data.frame()
##
                               0.00000000
## avg_ann_count
                               0.00000000
## avg_deaths_per_year
                               0.00000000
## target_death_rate
## incidence_rate
                               0.00000000
## med_income
                               0.00000000
## pop_est2015
                               0.00000000
  poverty_percent
                               0.0000000
## study_per_cap
                               0.0000000
                               0.00000000
## binned_inc
## median_age
                               0.0000000
## median age male
                               0.00000000
                               0.0000000
## median_age_female
   geography
                               0.00000000
## avg_household_size
                               0.00000000
## percent_married
                               0.0000000
## pct_no_hs18_24
                               0.00000000
   pct_hs18_24
                               0.00000000
  pct_some_col18_24
                               0.74991795
## pct_bach_deg18_24
                               0.00000000
## pct_hs25_over
                               0.0000000
                               0.00000000
## pct_bach_deg25_over
## pct_employed16_over
                               0.04988513
## pct_unemployed16_over
                               0.00000000
## pct_private_coverage
                               0.0000000
## pct_private_coverage_alone 0.19986872
                               0.00000000
## pct_emp_priv_coverage
```

```
## pct white
                              0.00000000
## pct_black
                              0.00000000
## pct_asian
                              0.00000000
## pct other race
                              0.00000000
## pct_married_households
                              0.00000000
## birth rate
                              0.00000000
#Pulling quartiles for study_per_cap categorical manipulation
study.quart <- with(cancer_raw, study_per_cap[study_per_cap > 0]) %>%
  quantile(., probs = c(0.25, 0.5, 0.75))
#Variable Manipulation
cancer.df <- cancer_raw %>%
                               #Remove Rows with > 20% missing
  dplyr::select(-pct_some_col18_24) %>% #Remove for too many missing
  mutate(
   pct_non_white = pct_black + pct_asian + pct_other_race, #Creating white, non-white percentages vari
    state = str_split_fixed(geography, ", ", 2)[ ,2] %>% as.factor(), #pulling state variable and casti
   binned_inc_lb = str_split_fixed(binned_inc, ", ", 2)[ ,1] %>% parse_number(), #pulling numeric lowe
   binned_inc_ub = str_split_fixed(binned_inc, ", ", 2)[ ,2] %>% parse_number(), #pulling numeric uppe
   binned_inc_point = (binned_inc_lb + binned_inc_ub)/2, #computing point estimate from ub, lb (interva
    study_quantile = ifelse(study_per_cap == 0, "None",
                           ifelse(study_per_cap > 0 & study_per_cap <= study.quart[1], "Low",</pre>
                                  ifelse(study_per_cap > study.quart[1] & study_per_cap <= study.quart[</pre>
                                          ifelse(study_per_cap > study.quart[2] & study_per_cap <= study</pre>
                                                 "Very High")))),
   study_quantile = as.factor(study_quantile) %>% fct_relevel(., "None", "Low", "Moderate", "High", "V
   avg_deaths_yr_pop = avg_deaths_per_year/pop_est2015, #incorporate two vars into one (multicollinea
   avg_ann_count_pop = avg_ann_count/pop_est2015 #incorporate two vars into one (multicollinearity)
  ) %>%
  dplyr::select(-c(binned_inc, geography, study_per_cap))
#Imputing Values with less than 20% missing (two variables)
  • pct_employed16_over \sim 4\%
  • pct_private_coverage_alone ~ 20%
#library(qlmnet)
#library(tidyverse)
#Impute those missing less than 20%
#1. pct_employed16_over
#2. pct private coverage alone
#Set up appropriate test and train for pct_employed16_over (removing other missing % variable and respo
train.df <- cancer.df %>% dplyr::select(-c(pct_private_coverage_alone, target_death_rate)) %>% filter(!
test.df <- cancer.df %>% dplyr::select(-c(pct_private_coverage_alone, target_death_rate)) %>% filter(is
#Function for imputation (after correct test, train set up), charstring must literally be the character
impute.lasso <- function(train.df, test.df, charstring){</pre>
  if ((charstring %in% names(train.df))) {
#pull variable index
index <- which(names(train.df) == charstring)</pre>
```

pct_public_coverage

pct_public_coverage_alone 0.00000000

0.00000000

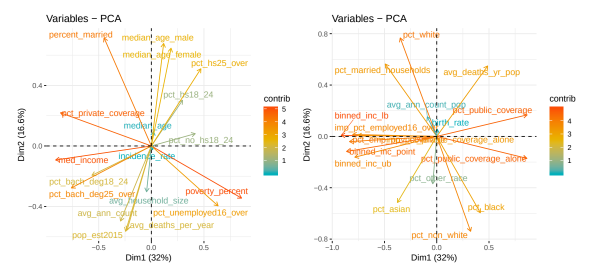
```
#Set up Matrices
#Create Design Matrix Train
X <- train.df[ ,-index] %>%
  names() %>%
  paste("~ ", paste(., collapse = "+")) %>%
  formula() %>%
  model.matrix(.,train.df)
#Create Design Matrix Test
X1 <- test.df[, -index] %>%
  names() %>%
  paste("~ ", paste(., collapse = "+")) %>%
  formula() %>%
  model.matrix(., test.df)
#Remove Intercept
X \leftarrow X[,-1]
X1 \leftarrow X1[,-1]
#Create Response vector (as matrix)
Y <- train.df[, index] %>% as.matrix()
#Optimize lambda
lambda.grid \leftarrow 10^{\circ}seq(-3,1,length = 100)
cv.lasso <- cv.glmnet(X, Y, alpha = 1, intercept = TRUE, lambda = lambda.grid, family = "gaussian")
#Grab optimal lambda
opt.lambda.lasso <- cv.lasso$lambda.min
#Run model
unemploy.lasso <- glmnet(X, Y, alpha = 1, intercept = TRUE, lambda = opt.lambda.lasso, family = "gaussi
#Return predictions
predict(unemploy.lasso, newx = X1)
    stop("Error: Incorrect variable name")
}
#Impute employed16_over_preds (first since it has less missing data ~4%)
employed16_over_preds <- impute.lasso(train.df = train.df, test.df, "pct_employed16_over")</pre>
#Set up appropriate test and train
train.df <- cancer.df %>% dplyr::select(-c(pct_employed16_over, target_death_rate)) %>% filter(!is.na(p
test.df <- cancer.df %>% dplyr::select(-c(pct_employed16_over, target_death_rate)) %>% filter(is.na(pct
#Impute pct_private_coverage_alone (second since it has more missing data ~20%)
pct_private_coverage_alone_preds <- impute.lasso(train.df = train.df, test.df, "pct_private_coverage_al
#Replace Imputed values
```

```
cancer.df <- cancer.df %>%
 mutate(imp_pct_employed16_over = ifelse(is.na(pct_employed16_over),
                                       employed16_over_preds, pct_employed16_over),
        imp_pct_private_coverage_alone = ifelse(is.na(pct_private_coverage_alone),
                                       pct_private_coverage_alone_preds, pct_private_coverage_alone)
       )
#Check
verif.df <- cancer.df %>%
 looks good so we will take out extraneous variables for final df.
#Looks good, so we will replace for our final data set
cancer.df <- cancer.df %>%
 dplyr::select(-c(pct_employed16_over, pct_private_coverage_alone))
#Check it out
str(cancer.df)
## Classes 'tbl_df', 'tbl' and 'data.frame':
                                             3047 obs. of 38 variables:
                                 : num 1397 173 102 427 57 ...
## $ avg_ann_count
                                 : num 469 70 50 202 26 152 97 71 36 1380 ...
## $ avg_deaths_per_year
## $ target_death_rate
                                : num 165 161 175 195 144 ...
                                 : num 490 412 350 430 350 ...
## $ incidence_rate
## $ med income
                                 : num 61898 48127 49348 44243 49955 ...
## $ pop_est2015
                                : num 260131 43269 21026 75882 10321 ...
                                : num 11.2 18.6 14.6 17.1 12.5 15.6 23.2 17.8 22.3 13.1 ...
## $ poverty_percent
## $ median_age
                                 : num 39.3 33 45 42.8 48.3 45.4 42.6 51.7 49.3 35.8 ...
                                : num 36.9 32.2 44 42.2 47.8 43.5 42.2 50.8 48.4 34.7 ...
## $ median_age_male
                                : num 41.7 33.7 45.8 43.4 48.9 48 43.5 52.5 49.8 37 ...
## $ median_age_female
                                : num 2.54 2.34 2.62 2.52 2.34 2.58 2.42 2.24 2.38 2.65 ...
## $ avg_household_size
## $ percent_married
                                 : num 52.5 44.5 54.2 52.7 57.8 50.4 54.1 52.7 55.9 50 ...
## $ pct_no_hs18_24
                                : num 11.5 6.1 24 20.2 14.9 29.9 26.1 27.3 34.7 15.6 ...
                                : num 39.5 22.4 36.6 41.2 43 35.1 41.4 33.9 39.4 36.3 ...
## $ pct_hs18_24
## $ pct_bach_deg18_24
                                : num 6.9 7.5 9.5 2.5 2 4.5 5.8 2.2 1.4 7.1 ...
## $ pct_hs25_over
                                 : num 23.2 26 29 31.6 33.4 30.4 29.8 31.6 32.2 28.8 ...
## $ pct_bach_deg25_over
                                : num 19.6 22.7 16 9.3 15 11.9 11.9 11.3 12 16.2 ...
## $ pct_unemployed16_over
                                : num 8 7.8 7 12.1 4.8 12.9 8.9 8.9 10.3 9.2 ...
                                 : num 75.1 70.2 63.7 58.4 61.6 60 49.5 55.8 55.5 69.9 ...
## $ pct_private_coverage
## $ pct_emp_priv_coverage
                                 : num 41.6 43.6 34.9 35 35.1 32.6 28.3 25.9 29.9 44.4 ...
## $ pct_public_coverage
                                 : num 32.9 31.1 42.1 45.3 44 43.2 46.4 50.9 48.1 31.4 ...
## $ pct_public_coverage_alone
                                : num 14 15.3 21.1 25 22.7 20.2 28.7 24.1 26.6 16.5 ...
                                 : num 81.8 89.2 90.9 91.7 94.1 ...
## $ pct white
                                 : num 2.595 0.969 0.74 0.783 0.27 ...
## $ pct_black
## $ pct_asian
                                 : num 4.822 2.246 0.466 1.161 0.666 ...
## $ pct_other_race
                                 : num 1.843 3.741 2.747 1.363 0.492 ...
## $ pct_married_households
                                 : num 52.9 45.4 54.4 51 54 ...
## $ birth_rate
                                 : num 6.12 4.33 3.73 4.6 6.8 ...
## $ pct_non_white
                                 : num 9.26 6.96 3.95 3.31 1.43 ...
## $ state
                                 : Factor w/ 51 levels "Alabama", "Alaska",..: 48 48 48 48 48 48 4
## $ binned_inc_lb
                                 : num 61494 48022 48022 42724 48022 ...
## $ binned_inc_ub
                                 : num 125635 51046 51046 45201 51046 ...
## $ binned_inc_point
                                : num 93565 49534 49534 43963 49534 ...
```

\$ study_quantile

: Factor w/ 5 levels "None", "Low", "Moderate", ...: 5 2 2 4 1 4 1 1 1

```
## $ avg_deaths_yr_pop
                                    : num 0.0018 0.00162 0.00238 0.00266 0.00252 ...
## $ avg_ann_count_pop
                                   : num 0.00537 0.004 0.00485 0.00563 0.00552 ...
                                 : num 51.9 55.9 45.9 48.3 48.2 44.1 51.8 40.9 39.5 56.6 ...
## $ imp pct employed16 over
## $ imp_pct_private_coverage_alone: num 54.7 53.8 43.5 40.3 43.9 ...
dim(cancer.df)
## [1] 3047
#Check new percentage missing after removing one and imputing two
# Check the percentage of NA values for each column
\#percentage\_NA = apply(cancer.df, 2, function(x) sum(length(which(is.na(x)))) / nrow(cancer.df))
#percentage_NA %>% data.frame() %>% knitr::kable()
#No more missing data and we only had to throw out one variable
#PCA Analysis for Variable Selection
*Plot is a bit messy, take subsets of the data and repeat
#Scale and perform pca (take out non-continuous vars)
cancer.pca <- cancer.df %>%
  dplyr::select(-c(state, study_quantile, target_death_rate)) %>%
  scale() %>%
  as.data.frame() %>%
  prcomp()
#str(cancer.pca)
name.vec <- cancer.df %>% dplyr::select(-c(state, study quantile, target death rate)) %>% names()
pca.viz1 <- fviz_pca_var(cancer.pca,</pre>
             col.var = "contrib",
             gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
             repel = TRUE,
             select.var = list(name = name.vec[1:18]))
pca.viz2 <- fviz_pca_var(cancer.pca,</pre>
             col.var = "contrib",
             gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
             repel = TRUE,
             select.var = list(name = name.vec[19:35]))
pca.viz1 + pca.viz2
```



Here, we used PCA component analysis on the scaled continuous predictor set, projecting our p=35 dimensional predictor space onto the 2 dimensional PCA space with the two principal component vectors explaining the highest % of variability in the data(basically the best basis in \mathcal{R}^{\in} to visualize how related each of our continuous predictors are to each other) vectors pointing in the same directions are explaining very similar types of the variance in the data (i.e. related, potentially multicollinearity), and the length(color) describes the magnitude or strength of how much of the variance in the data the predictor is explaining. It takes a while staring at it to understand exactly what is going on, and I split into two plots for clarity. Here are my takeaways:

- 1. Median age female and male explain a good amount of variability but are very related, we should take a average of the two for an average median age
- 2. Percent white and percent married are explaining similiar variability at similiar strength, pct races are explaining different types of variability in the data, while pct_white and pct_non_white are explaining inverse types of variability in the data (180 degree angle) makes me think we should keep seperate race percentages or pct_non_white, but not both (obviously) one while be better than the other.
- 3. Avg_ann_count, avg deaths, pop_est2015 are all explaining the same type and proportion of variability, should only use one (my best guess is pop_est2015 based on magnitude.)
- 4. pct_private, pct_public_cov, and pct_public alone are all explaining different types of variability but at sufficient magnitude and should be kept. However pct_private_cov is highly correlated with income, and should not be included if any type of income variable is in the model.
- 5. median_age is not a strong explanotary variable (by magnitude), and weirdly does not equal median_male + median_female /2. So I say we lose median_age and keep a variable for avg_median = median_male + median_female /2.
- 6. Drop the mutate avg_ann_count_pop, low magnitude and not explaining anything significantly. See (3.) for recommendation on which var to select there.
- 7. For income, all median_income, binned_inc_point estimate, binned_inc_lb, and binned_ub very related. I think we should either use median_income OR binned_inc_point estimate, but retain the lb, and ub as they seem to be explained different types of variability at good magnitude.
- 8. pct_bach_deg25_over is better than pecent_bach_deg18_24, explaining similar variability. Maybe take an average of the two or only include pct_bach_deg25_over.
- 9. Incidence rate has relatively small magnitude, but in a direction almost no other variable takes, so that should be kept I think.
- 10. House hold size looks relatively unrelated

11.