Final Report

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```
#Load Data
census_api_key("abc8289fa2ba274ced76d97c7f8ee31666a2c931", overwrite = TRUE, install = TRUE)

## Your original .Renviron will be backed up and stored in your R HOME directory if needed.

## Your API key has been stored in your .Renviron and can be accessed by Sys.getenv("CENSUS_API_KEY").

## To use now, restart R or run `readRenviron("~/.Renviron")`

## [1] "abc8289fa2ba274ced76d97c7f8ee31666a2c931"

if (FALSE) {
    census_api_key("abc8289fa2ba274ced76d97c7f8ee31666a2c931", install = TRUE)

# First time, reload your environment so you can use the key without restarting R.
    readRenviron("~/.Renviron")

# You can check it with:
Sys.getenv("CENSUS_API_KEY")
}
```

#Research Question:

How do measles vaccination rates vary across the country and demographics in schools?

overall vaccination status vs. state, overall vaccination status vs. type of school, each type of exemption (personal, religious, and medical) vs. state exemption vs. type of school. To analyze vaccination and exemption rates by states, we will use spatial data to show the change in these rates across the country. Then, we can use two-sample t-tests to test for significance of vaccination and exemption rates between different types of schools. If there are significantly lower vaccination rates in private schools vs. other types of schools, this will support our main hypothesis.

#Variable Manipulation

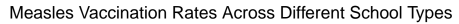
```
measles <- measles %>%
  filter(overall != (-1)) %>%
  mutate(numvaxx = round(enroll*overall*.01)) %>%
  mutate(unvaxx = enroll - numvaxx)%>%

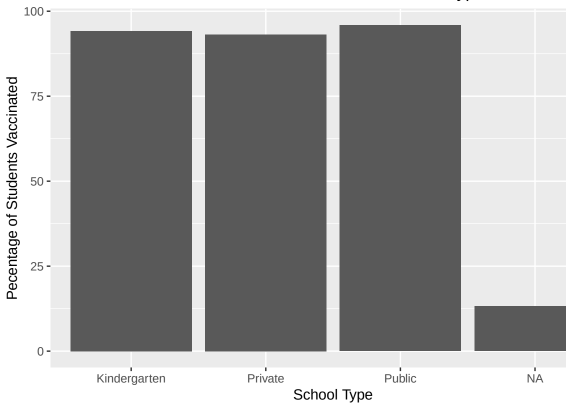
mutate(numxmed = round(enroll*.01*xmed)) %>%
  mutate(numxmed = ifelse(is.na(numxmed), 0, numxmed)) %>%

mutate(numxrel = round(enroll*.01*xrel)) %>%
  mutate(numxrel = ifelse(is.na(numxrel), 0, numxrel)) %>%

mutate(numxper = round(enroll*.01*xper)) %>%
  mutate(numxper = ifelse(is.na(numxper), 0, numxper)) %>%
```

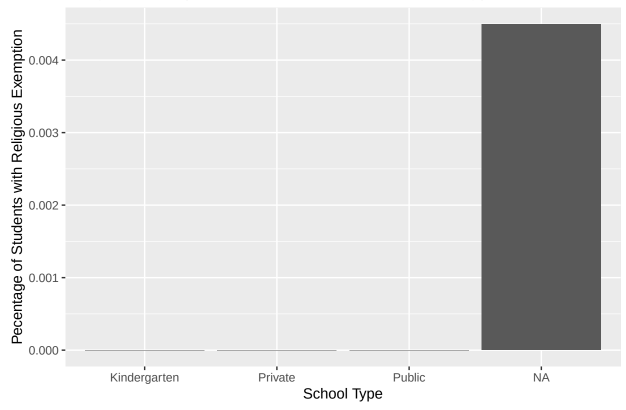
```
mutate(statefac = as.factor(state))
measlereg <- glm(cbind(numvaxx, unvaxx) ~ statefac + type, data=measles, family = binomial)
measlereg
## Call: glm(formula = cbind(numvaxx, unvaxx) ~ statefac + type, family = binomial,
      data = measles)
##
## Coefficients:
##
       (Intercept) statefacColorado
                                        statefacOhio
                                                          typePrivate
            3.7220
##
                           -1.6471
                                             -0.8457
                                                              -0.8424
        typePublic
##
           -0.6406
##
##
## Degrees of Freedom: 18645 Total (i.e. Null); 18641 Residual
    (20243 observations deleted due to missingness)
## Null Deviance:
                      114500
## Residual Deviance: 101300
                              AIC: 153400
summary(measlereg)
##
## Call:
## glm(formula = cbind(numvaxx, unvaxx) ~ statefac + type, family = binomial,
      data = measles)
##
## Deviance Residuals:
##
      Min
               1Q Median
                                 3Q
                                        Max
                                     17.086
## -59.896 -0.292 0.737
                              1.357
##
## Coefficients:
                   Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                   3.722031 0.049580 75.07 <2e-16 ***
## statefacColorado -1.647126  0.048295  -34.11
                                                <2e-16 ***
## statefacOhio
               ## typePrivate
                  <2e-16 ***
## typePublic
                  -0.640616 0.049617 -12.91
                                               <2e-16 ***
## --
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 114508 on 18645 degrees of freedom
## Residual deviance: 101255 on 18641 degrees of freedom
    (20243 observations deleted due to missingness)
## AIC: 153354
##
## Number of Fisher Scoring iterations: 5
```



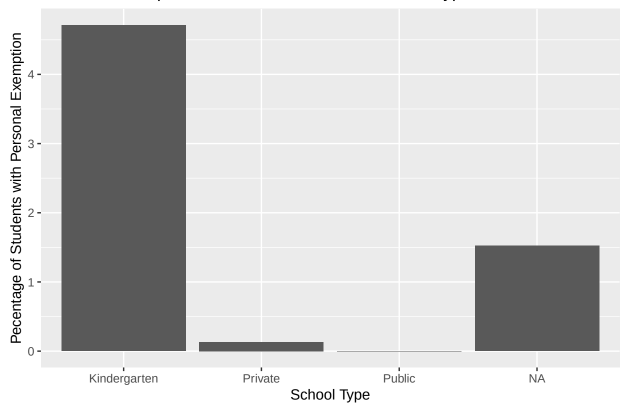


#Preliminary Bar Graphs

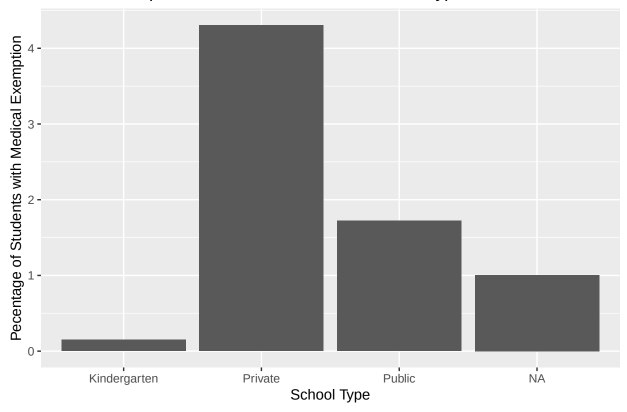
Religious Exemption Rates Across Different School Types



Personal Exemption Rates Across Different School Types



Medical Exemption Rates Across Different School Types



vaccination by state graph

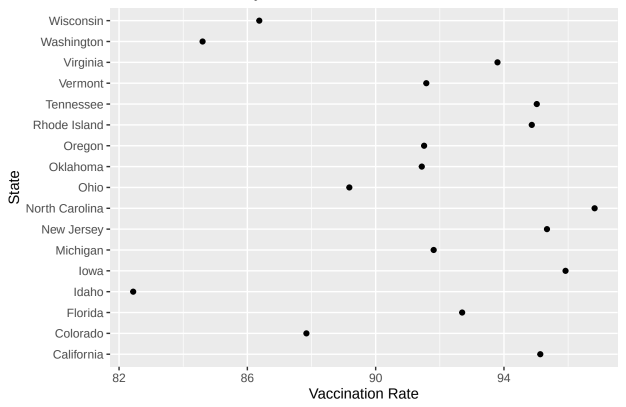
```
measles %>%
  count(state) %>%
  group_by(state)
```

```
## # A tibble: 17 x 2
## # Groups: state [17]
##
      state
                         n
##
      <chr>
                     <int>
   1 California
                     14224
##
   2 Colorado
                      1505
##
    3 Florida
                      2672
##
   4 Idaho
                       467
##
   5 Iowa
                      1163
   6 Michigan
                      2351
##
##
   7 New Jersey
                      2044
   8 North Carolina
                      2084
##
##
  9 Ohio
                      2917
## 10 Oklahoma
                      1052
## 11 Oregon
                       806
## 12 Rhode Island
                       215
## 13 Tennessee
                      1152
## 14 Vermont
                       338
## 15 Virginia
                      1413
```

```
## 16 Washington 1978
## 17 Wisconsin 2508

measles %>%
  filter(overall != (-1)) %>%
  group_by(state) %>%
  summarise(statemean = mean(overall)) %>%
  ggplot(aes(x = statemean, y = state)) +
  geom_point() +
  labs(x = "Vaccination Rate", y = "State", title = "Vaccination Rate by State")
```

Vaccination Rate by State



```
plotdata <- measles %>%
filter(overall != (-1)) %>%
group_by(state) %>%
summarise(statemean = mean(overall))
plot_usmap(data=plotdata, values = "statemean") +
  labs(title = "Vaccination Rate by State", fill = "Vaccination Rate") +
  theme(panel.background = element_rect(color = "black", fill = "lightblue"))
```

Vaccination Rate by State

