## Figure 2

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Set up the workspace.

```
rm(list=ls(all.names=TRUE))
setwd("/Users/bpetros/Desktop/PHIS")
libs <- c("cowplot", "ggsignif", "tidyverse")
invisible(lapply(libs, function(x) suppressPackageStartupMessages(library(x, character.only = TRUE))))
options(stringsAsFactors=FALSE, scipen = 999)
theme_set(theme_classic())</pre>
```

Read in the cleaned input files.

```
# read in cleaned pt data
pt <- read.csv("cleaned/rsv_patient.csv")
pt$Date = ymd(pt$Discharge_Date)

# read in cleaned tested pt data
ptest <- read.csv("cleaned/tested_patient.csv")
ptest$Date = ymd(ptest$Discharge_Date)</pre>
```

This function takes a df with column "Admit\_Age\_In\_Days" (integer) as input. It creates a column called Age\_Group that labels rows as belonging to one of the following age groups: 0-3 months, 3-6 months, 6-12 months, 1-2 years, 2-4 years, and >= 5 years.

```
add_age_group <- function(df) {
    df$Age_Group <- cut(
        df$Admit_Age_In_Days,
        breaks = c(0, 90, 180, 365, 730, 1460, Inf),
        labels = c("0-3 months", "3-6 months", "6-12 months", "1-2 years", "2-4 years", "5-17 years"),
        right = FALSE)
    return(df)}</pre>
```

This function takes a df with column "column\_name" as input. The column must represent a binary variable using either 0s and 1s (integer) or "N" and "Y" (character). The function prints the percent of rows for which the column equals either 1 or "Y".

```
summarize_and_print_column <- function(df, column_name) {
   df_summary <- df %>%
      summarize(
      Percent = round(sum(.data[[column_name]] == "Y" | .data[[column_name]] == 1) / n() * 100, 2))
   cat(column_name, "- Percent:", df_summary$Percent, "%\n")}
```

This function takes a df generated from a PHIS database download as input (i.e., multiple specific column names are required). It calculates and prints demographic (e.g., age, race, ethnicity) and clinical (e.g., presence of comorbid conditions, outcomes) summary statistics.

```
analyze_dataframe <- function(df) {</pre>
  # counts and percent of patients in each category (ED, ED -> IP, IP)
  ed_discharged_home <- sum(df$Patient_Type_Title == "ED Visit")</pre>
  percent_ed_discharged_home <- (ed_discharged_home / nrow(df)) * 100</pre>
  ed_admitted <- sum(df$Patient_Type_Title != "ED Visit" & df$ED_entry == 1)
  percent_ed_admitted <- (ed_admitted / nrow(df)) * 100</pre>
  admitted_without_ed <- sum(df$Patient_Type_Title != "ED Visit" & df$ED_entry == 0)
  percent_admitted_without_ed <- (admitted_without_ed / nrow(df)) * 100</pre>
  # patient type df
  ptype_counts <- list(</pre>
   ED_Discharged_Home_Count = ed_discharged_home,
   ED_Admitted_Count = ed_admitted,
   Admitted Without ED Count = admitted without ed)
  # print results
  cat("ED Discharged Home:\n")
  cat("Count:", ed_discharged_home, "\n")
  cat("Percentage:", percent_ed_discharged_home, "%\n\n")
  cat("ED Admitted:\n")
  cat("Count:", ed_admitted, "\n")
  cat("Percentage:", percent_ed_admitted, "%\n\n")
  cat("Admitted Without ED:\n")
  cat("Count:", admitted_without_ed, "\n")
  cat("Percentage:", percent_admitted_without_ed, "%\n\n")
  # median (IQR) age, in months
  df_summary <- df %>%
    summarize(
      Median_Age_Months = round(median(Admit_Age_In_Days) / 30.44, 2),
      Q1 = round(quantile(Admit_Age_In_Days, 0.25) / 30.44, 2),
      Q3 = round(quantile(Admit_Age_In_Days, 0.75) / 30.44, 2))
  # print age statistics
  cat("Admit_Age_In_Mo - Median:", df_summary$Median_Age_Months, "months\n")
  cat("Admit_Age_In_Mo - IQR (Q1 - Q3):", df_summary$Q1, "months -", df_summary$Q3, "months\n")
  # report percent in each age category
  age_groups = table(df$Age_Group)
  age_percents <- round(table(df$Age_Group) / nrow(df) * 100, 2)
  cat("Admit_Age_In_Days - Percent:\n")
  print(age_percents)
  # report percent male
  df_gender_summary <- df %>%
    summarize(Percent_Male = round(sum(Gender_Title == "Male") / n() * 100, 2))
  cat("Gender_Title - Percent Male:", df_gender_summary$Percent_Male, "%\n")
  # report percent in each ethnicity category
```

```
Ethnicity_Summary <- df %>%
  summarize(
    Percent_Hispanic = round(sum(Ethnicity_Title == "Hispanic or Latino") / n() * 100, 2),
   Percent Not Hispanic = round(sum(Ethnicity Title == "Not Hispanic or Latino") / n() * 100, 2),
    Percent_Unknown = round(sum(Ethnicity_Title == "Unknown") / n() * 100, 2))
cat("Ethnicity_Title - Percent Hispanic or Latino:", Ethnicity_Summary$Percent_Hispanic, "%\n")
cat("Ethnicity_Title - Percent Not Hispanic or Latino:", Ethnicity_Summary$Percent_Not_Hispanic, "%\n
cat("Ethnicity Title - Percent Unknown:", Ethnicity Summary Percent Unknown, "%\n")
# report percent in each racial group
race_columns <- c("Race_White", "Race_Black", "Race_Asian", "Race_Pacific_Islander", "Race_American_I
# column for multiple races
df <- df %>%
 mutate(Multiple = ifelse(rowSums(select(., all_of(race_columns)) == "Y") > 1, "Y", "N"))
# percent in each individual category
Race_Summary <- df %>%
  summarize(across(all_of(race_columns), ~ round(sum(. == "Y" & Multiple == "N") / n() * 100, 2))) %>
 rename_with(~ paste0(.x, "_Percentage"), all_of(race_columns))
# percent in multiple categories
Multiple_Percentage <- df %>%
  summarize(Multiple_Percentage = round(sum(Multiple == "Y") / n() * 100, 2))
# percent unknown
Unknown_Percentage <- 100 - sum(Race_Summary) - Multiple_Percentage</pre>
# print race category percents
Race_Summary <- Race_Summary %>%
 bind_rows(data.frame(Multiple = Multiple_Percentage, Unknown = Unknown_Percentage)) %>%
  set_names(nm = c("Race_White_Percentage", "Race_Black_Percentage", "Race_Asian_Percentage",
                   "Race_Pacific_Islander_Percentage", "Race_American_Indian_Percentage",
                   "Race_Other_Percentage", "Multiple_Percentage", "Unknown_Percentage"))
print(Race_Summary)
# save counts for stat testing
Race_Summary <- df %>%
  summarize(across(all_of(c(race_columns, "Multiple")), ~ sum(. == "Y")))
# report test categories
test_columns <- c("culture", "PCR", "SC2_multi", "antigen", "DNA_probe", "unspecified")
# column for multiple tests
 mutate(Multiple = ifelse(rowSums(select(., all_of(test_columns)) > 0) > 1, 1, 0))
# percent in each individual category
Test_Summary <- df %>%
  summarize(across(all_of(test_columns), ~ round(sum(. > 0 & Multiple == 0) / n() * 100, 3))) %>%
 rename_with(~ pasteO(.x, "_Percentage"), all_of(test_columns))
# percent in multiple categories
Multiple_Percentage <- df %>%
  summarize(Multiple_Percentage = round(sum(Multiple == 1) / n() * 100, 3))
# percent without test
No_Test_Percentage <- 100 - sum(Test_Summary) - Multiple_Percentage
# print test category percents
Test_Summary <- Test_Summary %>%
  bind_rows(data.frame(Multiple = Multiple_Percentage, No_Test = No_Test_Percentage)) %>%
  set_names(nm = c("culture", "PCR", "SC2_multi", "antigen", "DNA_probe",
                   "unspecified", "Multiple", "No_Test"))
```

```
print(Test_Summary)
# save counts for stat testing
test_columns <- c("culture", "PCR", "SC2_multi", "antigen", "DNA_probe")
Test Summary <- df %>%
  summarize(across(all_of(c(test_columns, "Multiple")), ~ sum(. > 0)),
            No_Test = sum(rowSums(select(., all_of(test_columns))) == 0))
# print percent with CCC
summarize_and_print_column(df, "Complex_Chronic_Condition_Flag")
# print percent with neonatal CCC
summarize_and_print_column(df, "Premature_And_Neonatal_Flag")
# print percent with CCC per age group
complex_condition_counts <- table(df$Age_Group[df$Complex_Chronic_Condition_Flag == "Y"])</pre>
complex_condition_percentages <- round(complex_condition_counts / table(df$Age_Group) * 100, 2)</pre>
cat("Age Categories - Complex Chronic Condition Percentage:\n")
print(complex_condition_percentages)
# print percent with positive test per age group
if ("pos" %in% colnames(df)) {
 summarize_and_print_column(df, "pos")
 pos_counts_by_age <- df %>%
   filter(!is.na(Age_Group)) %>%
    group by (Age Group) %>%
    summarize(
      Pos_1_Count = sum(pos == 1, na.rm = TRUE),
      Pos_0_Count = sum(pos == 0, na.rm = TRUE),
      Pos_1_Percent = round(mean(pos == 1, na.rm = TRUE) * 100, 5))
  cat("Age Groups - Percentage of rows where pos == 1:\n")
 print(data.frame(pos_counts_by_age))
  # report percent of positive tests by test type
  test_columns <- c("culture", "antigen", "PCR", "SC2_multi", "DNA_probe")
  if(all(test_columns %in% colnames(df))) {
    test_counts <- lapply(test_columns, function(test_col) {</pre>
      df %>%
        filter(.data[[test_col]] > 0) %>%
        summarise(
         Pos_1_Count = sum(.data[[test_col]] > 0 & pos == 1, na.rm = TRUE),
         Pos_0_Count = sum(.data[[test_col]] > 0 & pos == 0, na.rm = TRUE),
          Pos_1_Percent = round(sum(.data[[test_col]] > 0 & pos == 1) / sum(.data[[test_col]] > 0) *
        mutate(Test = test col)})
    test counts <- bind rows(test counts)</pre>
    cat("Test Counts - Positive cases where test > 0 and pos == 1 or pos == 0:\n")
   print(test_counts)}}
else {
 pos_counts_by_age = c()
 test_counts = c()}
# statistics of interest only for patients diagnosed with RSV
if ("flu" %in% colnames(df)) {
  # print percent with concurrent influenza
```

```
summarize_and_print_column(df, "flu")
  # print percent with concurrent SC2
  summarize_and_print_column(df, "sc2")
  # print percent admitted from ED
 ED_Dispo_Admitted_Percent <- df %>%
  filter(ED_entry == 1) %>%
  filter(!is.na(ED_Dispo)) %>%
  summarize(ED_Dispo_Admitted_Percent = round(sum(ED_Dispo == "ED_Admission") / n() * 100, 2))
  cat("Percent ED Patients Admitted:", ED_Dispo_Admitted_Percent$ED_Dispo_Admitted_Percent, "%\n")
  # only consider inpatients for the following statistics
  # print median (IQR) length of stay
 df = df %>% filter %>%
   filter(Patient_Type_Title != "ED Visit")
 LOS_Summary <- df %>%
    summarize(Median_Length_Of_Stay = round(median(Length_Of_Stay), 2),
    Q1_Length_Of_Stay = round(quantile(Length_Of_Stay, 0.25), 2),
    Q3_Length_Of_Stay = round(quantile(Length_Of_Stay, 0.75), 2))
  cat("Length_Of_Stay - Median:", LOS_Summary$Median_Length_Of_Stay, "\n")
  cat("Length_Of_Stay - Q1:", LOS_Summary$Q1_Length_Of_Stay, "\n")
  cat("Length_Of_Stay - Q3:", LOS_Summary$Q3_Length_Of_Stay, "\n")
  # print percent in each LOS category
 los_groups \leftarrow c(0, 1, 2, 4, 7, Inf)
  los_labels <- c("0-1 days", "1-2 days", "2-4 days", "5-7 days", ">7 days")
 df$LOS_Group <- cut(df$Length_Of_Stay, breaks = los_groups, labels = los_labels, right = FALSE)
 los_groups = table(df$LOS_Group)
 los_percents <- round(table(df$LOS_Group) / nrow(df) * 100, 2)</pre>
  cat("Length_Of_Stay - Percent:\n")
  print(los_percents)
  # print percent of IPs admitted to intensive care
  df$nicu_icu = ifelse(df$ICU_Flag == "Y" | df$NICU_Flag == "Y", "Y", "N")
  summarize_and_print_column(df, "nicu_icu")
  # print percent of IPs requiring mechanical ventilation
  summarize_and_print_column(df, "Mechanical_Vent_Flag")
  # print percent of IPs requiring ECMO
  summarize_and_print_column(df, "ECMO_Flag")
  # print percent of IPs expiring in hospital
  summarize_and_print_column(df, "Discharge_Mortality_Flag")}
  else {
    los_groups = c()}
return(list(ptype = ptype_counts, age = age_groups, los = los_groups, race = Race_Summary, test = Tes
```

This function takes two dfs with column "Age\_Group" (character) as input. It determines the distribution of the categorical variable Age\_Group in the reference\_df and randomly samples rows in target\_df, with

replacement, to generate a new target df with the same distribution of Age Group as reference df.

```
bootstrap_df_by_age <- function(target_df, reference_df, sample_size = nrow(target_df)) {
  # calc the distribution of Age_Group in reference_df
  reference_distribution <- table(reference_df$Age_Group)</pre>
  subset df <- data.frame()</pre>
  # loop through each age group
  for (age_group in unique(reference_df$Age_Group)) {
    # calc the target percent for this age group
    target_percentage <- reference_distribution[age_group] / sum(reference_distribution)</pre>
    # calc the sample size for this age group
    n_to_select <- round(target_percentage * sample_size)</pre>
    # select rows from target_df matching the age group
    available_rows <- target_df %>%
      filter(Age_Group == age_group)
    # make sure there are rows in this age group present
    if (nrow(available_rows) > 0) {
      # randomly sample rows with replacement
      sampled_rows <- available_rows %>% sample_n(size = n_to_select, replace = TRUE)
      subset_df <- bind_rows(subset_df, sampled_rows)}}</pre>
  return(subset_df)}
```

Add the columns "Age\_Group", "nicu\_icu", and "Race" to the dfs with diagnosed and tested patient information.

```
# create the "Age_Group" column from the Admit_Age_In_Days column
pt = add_age_group(pt)
ptest = add_age_group(ptest)
# create the "nicu_icu" column from the binary ICU flags
pt$nicu_icu = ifelse(pt$ICU_Flag == "Y" | pt$NICU_Flag == "Y", "Y", "N")
# create the "Race" column from the binary race columns
race_columns <- c("Race_White", "Race_Black", "Race_Asian", "Race_Pacific_Islander", "Race_American_Ind
pt$Race <- apply(pt[race_columns], 1, function(row) {</pre>
  y_columns <- names(row[row == "Y"])</pre>
  if (length(y_columns) == 1) {
    return(y_columns)
  } else if (length(y_columns) > 1) {
    return("Multiple")
  } else {
    return("Unknown")}})
ptest$Race <- apply(ptest[race_columns], 1, function(row) {</pre>
  y_columns <- names(row[row == "Y"])</pre>
  if (length(y_columns) == 1) {
    return(y_columns)
 } else if (length(y_columns) > 1) {
   return("Multiple")
  } else {
    return("Unknown")}})
```

Report the median (IQR) of monthly RSV cases across all hospitals during the intermediate period.

```
# print median (IQR) of cases between phases
pandemic <- pt %>%
  filter(Date >= as.Date('2020-04-01') & Date < as.Date('2021-04-01')) %%
  mutate(YearMonth = format(Date, "%Y-%m")) %>%
  group_by(YearMonth) %>%
  summarise(Count = n())
print(paste("Median:", median(pandemic$Count)))
## [1] "Median: 46.5"
print(paste("Q1:", quantile(pandemic$Count, probs = 0.25)))
## [1] "Q1: 32"
print(paste("Q3:", quantile(pandemic$Count, probs = 0.75)))
## [1] "Q3: 78.5"
rm(pandemic)
Produce summary statistics for patients tested for RSV in the pre-pandemic and post-emergence phases, and
identify statistically significant differences in demographic or clinical variables across the two phases (Table
1).
# pre-pandemic
pre_ptest = subset(ptest, Date < as.Date("2020-04-01"))</pre>
print("Pre-pandemic tested patients:")
## [1] "Pre-pandemic tested patients:"
pre_ptest_cat = analyze_dataframe(pre_ptest)
## ED Discharged Home:
## Count: 119092
## Percentage: 56.59621 %
##
## ED Admitted:
## Count: 73090
## Percentage: 34.73463 %
## Admitted Without ED:
## Count: 18242
## Percentage: 8.669163 %
## Admit_Age_In_Mo - Median: 8.57 months
## Admit_Age_In_Mo - IQR (Q1 - Q3): 3.12 months - 20.01 months
## Admit_Age_In_Days - Percent:
```

##

```
## 0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                       2-4 years
                                                                  5-17 years
##
         23.91
                     15.63
                                  20.29
                                              19.38
                                                            9.99
                                                                       10.81
## Gender Title - Percent Male: 55.84 %
## Ethnicity_Title - Percent Hispanic or Latino: 31.59 %
## Ethnicity_Title - Percent Not Hispanic or Latino: 62.8 %
## Ethnicity_Title - Percent Unknown: 5.61 %
     Race_White_Percentage Race_Black_Percentage Race_Asian_Percentage
                     43.94
## 1
                                               24
## 2
                         NA
                                               NA
                                                                      NA
##
     Race_Pacific_Islander_Percentage Race_American_Indian_Percentage
## 1
                                  0.35
## 2
                                                                     NA
                                    ΝA
##
     Race_Other_Percentage Multiple_Percentage Unknown_Percentage
## 1
                      9.91
                                             NA
                                                                 NA
## 2
                        NA
                                           1.15
##
     culture
               PCR SC2_multi antigen DNA_probe unspecified Multiple
## 1
       3.054 5.256
                            0 83.387
                                          4.774
                                                       0.011
## 2
          NA
                NA
                          NA
                                   NA
                                             NA
                                                          NA
                                                                3.518
##
                      No_Test
## 1
                            NA
## 2 0.000000000000008881784
## Complex_Chronic_Condition_Flag - Percent: 18.45 %
## Premature_And_Neonatal_Flag - Percent: 2.71 %
## Age Categories - Complex Chronic Condition Percentage:
##
   0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                       2-4 years 5-17 years
##
                     10.55
                                              14.64
                                                           27.68
                                                                       51.75
         13.77
                                  11.39
## pos - Percent: 21.5 %
## Age Groups - Percentage of rows where pos == 1:
##
       Age_Group Pos_1_Count Pos_0_Count Pos_1_Percent
## 1
     0-3 months
                       13230
                                    37074
                                               26.30010
## 2 3-6 months
                         9035
                                    23859
                                               27.46702
## 3 6-12 months
                         9658
                                    33040
                                               22.61933
       1-2 years
                         8297
                                    32475
                                               20.34975
## 5
       2-4 years
                         3789
                                    17229
                                               18.02741
                        1242
                                    21496
                                                5.46222
## 6 5-17 years
## Test Counts - Positive cases where test > 0 and pos == 1 or pos == 0:
     Pos_1_Count Pos_0_Count Pos_1_Percent
                                                 Test
## 1
             942
                         8833
                                     9.6368
                                              culture
## 2
           41883
                      139957
                                    23.0329
                                              antigen
## 3
            2305
                       10384
                                    18.1653
                                                  PCR
## 4
               0
                            0
                                        NaN SC2 multi
## 5
            1054
                       12674
                                     7.6777 DNA_probe
# post-emergence
post_ptest <- subset(ptest, Date >= as.Date("2021-04-01"))
print("Post-emergence tested patients:")
## [1] "Post-emergence tested patients:"
post_ptest_cat = analyze_dataframe(post_ptest)
```

## ED Discharged Home:

```
## Count: 479941
## Percentage: 66.33967 %
##
## ED Admitted:
## Count: 195318
## Percentage: 26.99776 %
## Admitted Without ED:
## Count: 48201
## Percentage: 6.662566 %
## Admit_Age_In_Mo - Median: 35.05 months
## Admit_Age_In_Mo - IQR (Q1 - Q3): 11.99 months - 88.21 months
## Admit_Age_In_Days - Percent:
##
##
   0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                       2-4 years 5-17 years
##
          8.50
                      5.73
                                  10.77
                                              15.63
                                                           17.57
                                                                       41.80
## Gender Title - Percent Male: 54.19 %
## Ethnicity_Title - Percent Hispanic or Latino: 30.67 %
## Ethnicity_Title - Percent Not Hispanic or Latino: 63.81 %
## Ethnicity_Title - Percent Unknown: 5.51 %
     Race_White_Percentage Race_Black_Percentage Race_Asian_Percentage
                                            23.07
## 1
                     48.89
                                                                    4.33
## 2
                        NA
                                               NA
##
     Race_Pacific_Islander_Percentage Race_American_Indian_Percentage
## 1
                                  0.45
                                                                   0.35
## 2
                                    NA
                                                                     NA
     Race_Other_Percentage Multiple_Percentage Unknown_Percentage
##
## 1
                     13.73
                                             NA
## 2
                        NA
                                           3.22
                                                               5.96
                PCR SC2_multi antigen DNA_probe unspecified Multiple
##
     culture
## 1
        0.26 10.777
                       80.782
                                 7.518
                                           0.142
## 2
          NA
                           NA
                                    NA
                                              NA
                                                           NA
                                                                 0.521
##
                      No_Test
## 1
## 2 0.000000000000007771561
## Complex_Chronic_Condition_Flag - Percent: 15.84 %
## Premature_And_Neonatal_Flag - Percent: 1.64 %
## Age Categories - Complex Chronic Condition Percentage:
##
   0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                      2-4 years 5-17 years
##
         18.73
                     11.84
                                  10.32
                                              10.35
                                                           12.63
                                                                       20.63
## pos - Percent: 7.48 %
## Age Groups - Percentage of rows where pos == 1:
       Age_Group Pos_1_Count Pos_0_Count Pos_1_Percent
## 1 0-3 months
                                    52518
                        8967
                                               14.58404
## 2 3-6 months
                        7209
                                    34261
                                               17.38365
## 3 6-12 months
                        9049
                                    68843
                                               11.61737
       1-2 years
                       11333
                                   101761
                                               10.02087
## 5
       2-4 years
                       10800
                                   116280
                                                8.49858
## 6 5-17 years
                        6740
                                   295699
                                                2.22855
## Test Counts - Positive cases where test > 0 and pos == 1 or pos == 0:
     Pos_1_Count Pos_0_Count Pos_1_Percent
                                                 Test
## 1
             183
                        1797
                                     9.2424
                                              culture
```

```
48200
## 2
           6583
                                 12.0165
                                           antigen
## 3
           8923
                     72421
                                 10.9695
                                               PCR.
          38740
                     549335
## 4
                                  6.5876 SC2 multi
## 5
             65
                        984
                                  6.1964 DNA_probe
# compare pre-pandemic and post-emergence phases
agepv = wilcox.test(pre_ptest$Admit_Age_In_Days/30.44, post_ptest$Admit_Age_In_Days/30.44, paired = FAL
print("Age p-value:")
## [1] "Age p-value:"
agepv
## [1] O
agecatpv = chisq.test(rbind(pre_ptest_cat$age, post_ptest_cat$age))$p.value
print("Age group p-value:")
## [1] "Age group p-value:"
agecatpv
## [1] 0
sexpv = fisher.test(rbind(table(pre_ptest$Gender_Title)[1:2], table(post_ptest$Gender_Title)[1:2]))$p.v.
print("Sex p-value:")
## [1] "Sex p-value:"
sexpv
ethnpv = fisher.test(rbind(table(pre_ptest\$Ethnicity_Title)[1:2], table(post_ptest\$Ethnicity_Title)[1:2]
print("Ethnicity p-value:")
## [1] "Ethnicity p-value:"
ethnpv
## [1] 0.000000000000005942188
racepv = chisq.test(rbind(pre_ptest_cat$race, post_ptest_cat$race))$p.value
print("Race p-value:")
## [1] "Race p-value:"
```

```
racepv
## [1] 0
cccpv = fisher.test(rbind(table(pre_ptest$Complex_Chronic_Condition_Flag[pre_ptest$Complex_Chronic_Cond
                                                 table(post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Complex_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chronic_Chron
print("CCC p-value:")
## [1] "CCC p-value:"
cccpv
# Fisher's for each age group's CCC distribution
cc_pv <- list()
age_groups <- unique(as.character(ptest$Age_Group))</pre>
for (age_group in age_groups) {
   table_pre <- table(pre_ptest$Complex_Chronic_Condition_Flag[pre_ptest$Age_Group == age_group & pre_pt
    table_post <- table(post_ptest$Complex_Chronic_Condition_Flag[post_ptest$Age_Group == age_group & pos
   pvalue <- fisher.test(rbind(table_pre, table_post))$p.value</pre>
    cc_pv[[age_group]] <- pvalue}</pre>
# print results
for (i in 1:length(age_groups)) {
    cat("Age Group:", age_groups[i], "- p-value:", cc_pv[[i]], "\n")}
## Age Group: 2-4 years - p-value: 0
## Age Group: 5-17 years - p-value: 0
## Age Group: 3-6 months - p-value: 0.00000002722992
## Age Group: 6-12 months - p-value: 0.0000001276932
prempv = fisher.test(rbind(table(pre_ptest$Premature_And_Neonatal_Flag[pre_ptest$Premature_And_Neonatal
print("Premature & neonatal CCC p-value:")
## [1] "Premature & neonatal CCC p-value:"
prempv
ptypepv = chisq.test(rbind(unlist(pre_ptest_cat$ptype), unlist(post_ptest_cat$ptype)))$p.value
print("Patient type p-value:")
## [1] "Patient type p-value:"
```

```
ptypepv
## [1] 0
testpv = chisq.test(rbind(unlist(pre_ptest_cat$test), unlist(post_ptest_cat$test)))$p.value
print("Test type p-value:")
## [1] "Test type p-value:"
testpv
## [1] O
# Fisher's for each age group's test positivity
pos pv <- list()</pre>
age_groups <- unique(as.character(ptest$Age_Group))</pre>
for (age_group in age_groups) {
 table_pre <- table(pre_ptest$pos[pre_ptest$Age_Group == age_group])</pre>
 table_post <- table(post_ptest$pos[post_ptest$Age_Group == age_group])</pre>
 pvalue <- fisher.test(rbind(table_pre, table_post))$p.value</pre>
 pos_pv[[age_group]] <- pvalue}</pre>
for (i in 1:length(age_groups)) {
 cat("Age stratified test positivity:", age_groups[i], " p-value:", pos_pv[[i]], "\n")}
## Age stratified test positivity: 1-2 years p-value: 0
## Age stratified test positivity: 2-4 years p-value: 0
## Age stratified test positivity: 0-3 months p-value: 0
## Age stratified test positivity: 6-12 months p-value: 0
pos_pv = fisher.test(rbind(table(pre_ptest$pos), table(post_ptest$pos)))$p.value
print("Positive test p-value:")
## [1] "Positive test p-value:"
pos_pv
## [1] 0
pos_cult_pv = fisher.test(rbind(table(pre_ptest$pos[pre_ptest$culture > 0]), table(post_ptest$pos[post_
print("Positive culture p-value:")
## [1] "Positive culture p-value:"
pos_cult_pv
```

## [1] 0.6152608

```
pos_ant_pv = fisher.test(rbind(table(pre_ptest$pos[pre_ptest$antigen > 0]), table(post_ptest$pos[post_p
print("Positive antigen p-value:")
## [1] "Positive antigen p-value:"
pos_ant_pv
## [1] 0
pos_pcr_pv = fisher.test(rbind(table(pre_ptest$pos[pre_ptest$PCR > 0]), table(post_ptest$pos[post_ptest
print("Positive PCR p-value:")
## [1] "Positive PCR p-value:"
pos_pcr_pv
pos_dna_pv = fisher.test(rbind(table(pre_ptest$pos[pre_ptest$DNA_probe > 0]), table(post_ptest$pos[post
print("Positive DNA probe p-value:")
## [1] "Positive DNA probe p-value:"
pos_dna_pv
## [1] 0.08967613
rm(list = ls(pattern = "pv|tab"))
rm(i)
Produce summary statistics for patients diagnosed with RSV in the pre-pandemic and post-emergence phases,
and identify statistically significant differences in demographic or clinical variables across the two phases
(Table 2).
pt = pt %>% filter(Date < as.Date("2020-04-01") | Date >= as.Date("2021-04-01"))
pt$phase = ifelse(pt$Date < as.Date("2020-04-01"), "Pre-Pandemic", "Post-Emergence")</pre>
# pre-pandemic
pre_pt = subset(pt, Date < as.Date("2020-04-01"))</pre>
print("Pre-pandemic diagnosed patients:")
## [1] "Pre-pandemic diagnosed patients:"
pre_pt_cat = analyze_dataframe(pre_pt)
```

```
## ED Discharged Home:
## Count: 62933
## Percentage: 31.13476 %
## ED Admitted:
## Count: 105867
## Percentage: 52.37544 %
## Admitted Without ED:
## Count: 33331
## Percentage: 16.4898 %
## Admit_Age_In_Mo - Median: 6.87 months
## Admit_Age_In_Mo - IQR (Q1 - Q3): 2.6 months - 17.48 months
## Admit_Age_In_Days - Percent:
##
##
   0-3 months 3-6 months 6-12 months
                                         1-2 years
                                                      2-4 years 5-17 years
##
         27.99
                     17.85
                                 18.85
                                              17.72
                                                          11.16
                                                                       6.43
## Gender_Title - Percent Male: 55.31 %
## Ethnicity_Title - Percent Hispanic or Latino: 25.45 %
## Ethnicity_Title - Percent Not Hispanic or Latino: 69.83 %
## Ethnicity_Title - Percent Unknown: 4.72 %
     Race_White_Percentage Race_Black_Percentage Race_Asian_Percentage
## 1
                     58.61
                                            18.43
                                                                    2.8
## 2
                        NΑ
                                               NA
                                                                     NΑ
     Race_Pacific_Islander_Percentage Race_American_Indian_Percentage
## 1
                                 0.53
## 2
                                   NA
                                                                    NA
##
     Race_Other_Percentage Multiple_Percentage Unknown_Percentage
## 1
                     10.31
                                            NA
                                                                NA
## 2
                        NA
                                           1.49
     culture PCR SC2_multi antigen DNA_probe unspecified Multiple No_Test
                             20.293
                                        0.274
## 1
       0.282 1.09
                          0
                                                     0.002
                                                                 NA
## 2
          NA
                                                              0.441
               NA
                         NA
                                 NA
                                           NA
                                                        NΑ
                                                                    77.618
## Complex_Chronic_Condition_Flag - Percent: 16.84 %
## Premature_And_Neonatal_Flag - Percent: 3.1 %
## Age Categories - Complex Chronic Condition Percentage:
##
  0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                      2-4 years 5-17 years
                                                                      52.76
##
         12.14
                      8.91
                                 11.89
                                              16.67
                                                          29.22
## flu - Percent: 2.19 %
## sc2 - Percent: 0 %
## Percent ED Patients Admitted: 50.55 %
## Length_Of_Stay - Median: 3
## Length_Of_Stay - Q1: 2
## Length_Of_Stay - Q3: 5
## Length_Of_Stay - Percent:
##
## 0-1 days 1-2 days 2-4 days 5-7 days >7 days
       3.10
              19.91
                        36.62
                                 22.90
                                          17.47
## nicu_icu - Percent: 32.25 %
## Mechanical_Vent_Flag - Percent: 11.9 %
## ECMO_Flag - Percent: 0.22 %
## Discharge_Mortality_Flag - Percent: 0.24 %
```

```
# post-emergence
post_pt <- subset(pt, Date >= as.Date("2021-04-01"))
print("Post-emergence diagnosed patients:")
## [1] "Post-emergence diagnosed patients:"
post_pt_cat = analyze_dataframe(post_pt)
## ED Discharged Home:
## Count: 62192
## Percentage: 44.58016 %
##
## ED Admitted:
## Count: 64719
## Percentage: 46.39155 %
##
## Admitted Without ED:
## Count: 12595
## Percentage: 9.028286 %
## Admit_Age_In_Mo - Median: 11.89 months
## Admit_Age_In_Mo - IQR (Q1 - Q3): 4.2 months - 28.32 months
## Admit_Age_In_Days - Percent:
##
##
   0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                      2-4 years 5-17 years
         18.18
                     14.56
                                 17.40
                                              20.00
                                                          18.46
                                                                      11.40
## Gender_Title - Percent Male: 54.75 %
## Ethnicity_Title - Percent Hispanic or Latino: 27.09 %
## Ethnicity_Title - Percent Not Hispanic or Latino: 67.74 %
## Ethnicity_Title - Percent Unknown: 5.17 %
##
     Race_White_Percentage Race_Black_Percentage Race_Asian_Percentage
## 1
                     57.03
                                            19.15
## 2
                        NA
                                               NA
                                                                     NA
##
     Race_Pacific_Islander_Percentage Race_American_Indian_Percentage
## 1
                                  0.8
                                                                   0.3
                                   NA
                                                                    NA
     Race_Other_Percentage Multiple_Percentage Unknown_Percentage
## 1
                     11.21
                                             NA
                                                                NA
## 2
                                           2.72
                                                              5.49
                        NA
               PCR SC2_multi antigen DNA_probe unspecified Multiple No_Test
     culture
       0.126 2.632
                      31.042
                               4.645
                                          0.042
## 2
          NA
                NA
                          NA
                                  NΔ
                                             NΔ
                                                         NA
                                                                0.29 61.223
## Complex_Chronic_Condition_Flag - Percent: 11.69 %
## Premature_And_Neonatal_Flag - Percent: 2.57 %
## Age Categories - Complex Chronic Condition Percentage:
##
   0-3 months 3-6 months 6-12 months
                                          1-2 years
                                                      2-4 years 5-17 years
##
         11.98
                      6.46
                                  7.74
                                               9.45
                                                          13.21
                                                                      25.46
## flu - Percent: 1.38 %
## sc2 - Percent: 2.73 %
## Percent ED Patients Admitted: 38.7 %
```

## Length\_Of\_Stay - Median: 2

```
## Length_Of_Stay - Q1: 1
## Length_Of_Stay - Q3: 4
## Length_Of_Stay - Percent:
##
## 0-1 days 1-2 days 2-4 days 5-7 days >7 days
##
      3.76
              23.35
                      39.83
                               20.99
                                        12.08
## nicu icu - Percent: 27.16 %
## Mechanical_Vent_Flag - Percent: 7.36 %
## ECMO_Flag - Percent: 0.15 %
## Discharge_Mortality_Flag - Percent: 0.16 %
# compare pre-pandemic and post-emergence phases
agepv = wilcox.test(pre_pt$Admit_Age_In_Days/30.44, post_pt$Admit_Age_In_Days/30.44, paired = FALSE)$p.
print("Age p-value:")
## [1] "Age p-value:"
agepv
## [1] 0
agecatpv = chisq.test(rbind(pre_pt_cat$age, post_pt_cat$age))$p.value
print("Age group p-value:")
## [1] "Age group p-value:"
agecatpv
## [1] O
sexpv = fisher.test(rbind(table(pre_pt$Gender_Title)[1:2], table(post_pt$Gender_Title)[1:2]))$p.value
print("Sex p-value:")
## [1] "Sex p-value:"
sexpv
## [1] 0.001180125
ethnpv = fisher.test(rbind(table(pre_pt$Ethnicity_Title)[1:2], table(post_pt$Ethnicity_Title)[1:2]))$p.
print("Ethnicity p-value:")
## [1] "Ethnicity p-value:"
ethnpv
```

```
racepv = chisq.test(rbind(pre_pt_cat$race, post_pt_cat$race))$p.value
print("Race p-value:")
## [1] "Race p-value:"
racepv
flupv = fisher.test(rbind(table(pre_pt$flu), table(post_pt$flu)))$p.value
print("Flu p-value:")
## [1] "Flu p-value:"
flupv
print("SC2 p-value:")
## [1] "SC2 p-value:"
sc2pv = fisher.test(rbind(table(pre_pt\$sc2), table(post_pt\$sc2)))\$p.value
sc2pv
## [1] 0
cccpv = fisher.test(rbind(table(pre_pt$Complex_Chronic_Condition_Flag), table(post_pt$Complex_Chronic_C
print("CCC p-value:")
## [1] "CCC p-value:"
cccpv
## [1] 0
# Fisher's for each age group's CCC distribution
cc_pv <- list()</pre>
age_groups <- unique(as.character(pt$Age_Group))</pre>
for (age_group in age_groups) {
 table_pre <- table(pre_pt$Complex_Chronic_Condition_Flag[pre_pt$Age_Group == age_group])
 table_post <- table(post_pt$Complex_Chronic_Condition_Flag[post_pt$Age_Group == age_group])
 pvalue <- fisher.test(rbind(table_pre, table_post))$p.value</pre>
 cc_pv[[age_group]] <- pvalue}</pre>
for (i in 1:length(age_groups)) {
 cat("Age stratified CCC:", age_groups[i], "p-value:", cc_pv[[i]], "\n")}
```

```
## Age stratified CCC: 0-3 months p-value: 0.5015325
## Age stratified CCC: 3-6 months p-value: 0.00000000000000000000001500192
## Age stratified CCC: 2-4 years p-value: 0
## Age stratified CCC: 5-17 years p-value: 0
prempv = fisher.test(rbind(table(pre_pt$Premature_And_Neonatal_Flag), table(post_pt$Premature_And_Neona
print("Premature & neonatal CCC p-value:")
## [1] "Premature & neonatal CCC p-value:"
prempv
## [1] 0.0000000000000000001076395
ptypepv = chisq.test(rbind(unlist(pre_pt_cat$ptype), unlist(post_pt_cat$ptype)))$p.value
print("Patient type p-value:")
## [1] "Patient type p-value:"
ptypepv
## [1] 0
testpv = chisq.test(rbind(unlist(pre_pt_cat$test), unlist(post_pt_cat$test)))$p.value
print("Test type p-value:")
## [1] "Test type p-value:"
testpv
## [1] 0
admpv = fisher.test(rbind(table(pre_pt$ED_Dispo[pre_pt$ED_entry == 1]),
                      table(post_pt$ED_Dispo[post_pt$ED_entry == 1])))$p.value
print("ED admission p-value:")
## [1] "ED admission p-value:"
admpv
## [1] 0
lospv = wilcox.test(pre_pt$Length_Of_Stay[pre_pt$Patient_Type_Title != "ED Visit"],
                 post_pt$Length_Of_Stay[post_pt$Patient_Type_Title != "ED Visit"], paired = FALSE)$p
print("Length of stay p-value:")
## [1] "Length of stay p-value:"
```

```
lospv
loscatpv = chisq.test(rbind(pre_pt_cat$los, post_pt_cat$los))$p.value
print("Length of stay (categorical) p-value:")
## [1] "Length of stay (categorical) p-value:"
loscatpv
icupv = fisher.test(rbind(table(pre_pt$nicu_icu[pre_pt$Patient_Type_Title != "ED Visit"]),
                   table(post_pt$nicu_icu[post_pt$Patient_Type_Title != "ED Visit"])))$p.value
print("ICU p-value:")
## [1] "ICU p-value:"
icupv
mechpv = fisher.test(rbind(table(pre_pt$Mechanical_Vent_Flag[pre_pt$Patient_Type_Title != "ED Visit"]),
                    table(post_pt$Mechanical_Vent_Flag[post_pt$Patient_Type_Title != "ED Visit"]
print("Mechanical ventilation p-value:")
## [1] "Mechanical ventilation p-value:"
mechpv
ecmopv = fisher.test(rbind(table(pre_pt$ECMO_Flag[pre_pt$Patient_Type_Title != "ED Visit"]),
                    table(post_pt$ECMO_Flag[post_pt$Patient_Type_Title != "ED Visit"])))$p.value
print("ECMO p-value:")
## [1] "ECMO p-value:"
ecmopv
## [1] 0.0004307284
deathpv = fisher.test(rbind(table(pre_pt$Discharge_Mortality_Flag[pre_pt$Patient_Type_Title != "ED Visi
                    table(post_pt$Discharge_Mortality_Flag[post_pt$Patient_Type_Title != "ED Vi
print("Death p-value:")
## [1] "Death p-value:"
```

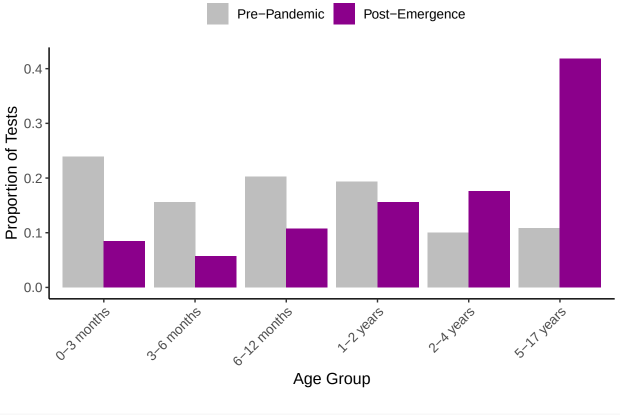
```
deathpv
```

```
## [1] 0.0003390302
```

```
rm(list = ls(pattern = "pv|tab"))
rm(i)
```

Plot the proportion of tested patients in each age group (Figure 2A).

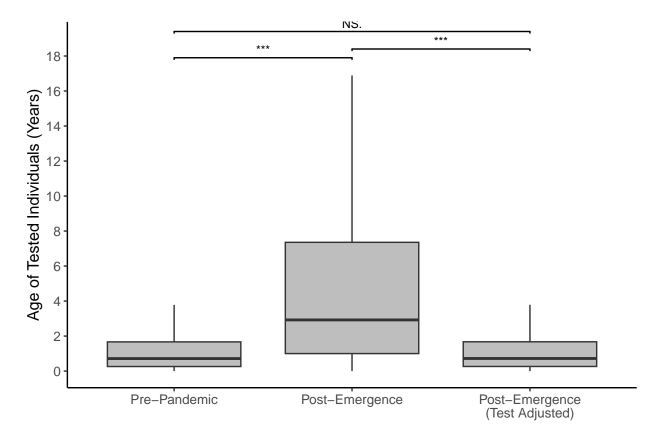
```
test_age = rbind(pre_ptest_cat$age/nrow(pre_ptest), post_ptest_cat$age/nrow(post_ptest))
rownames(test_age) <- c("Pre-Pandemic", "Post-Emergence")</pre>
# pivot the data to long format for plotting
test_age_long = gather(as.data.frame(test_age), key = "Age Group", value = "Proportion")
test age long$Group <- rownames(test age)</pre>
test_age_long$Group <- factor(test_age_long$Group, levels = c("Pre-Pandemic", "Post-Emergence"))
test_age_long$ Age Group` <- factor(test_age_long$ Age Group`, levels = c("0-3 months",
                             "3-6 months", "6-12 months", "1-2 years", "2-4 years", "5-17 years"))
# plot the proportions
fig2a \leftarrow ggplot(test_age_long, aes(x = `Age Group`, y = Proportion, fill = Group)) +
  geom_bar(stat = "identity", position = "dodge") +
 labs(title = "",
       x = "Age Group",
       y = "Proportion of Tests") +
  scale_fill_manual(values = c("grey", "darkmagenta")) +
  theme classic() +
  theme(axis.text.x = element_text(angle = 45, hjust = 1),
        plot.title = element_text(hjust = 0.5),
       legend.position = "top",
       text = element text(size = 12)) +
  guides(fill = guide legend(title = NULL))
fig2a
```



```
rm(test_age, test_age_long)
```

Use bootstrapping to generate a post-emergence testing df with the age distribution (in 90-day bins) derived from the pre-pandemic testing df. Plot the pre-pandemic, post-emergence, and test-adjusted age distributions of tested patients (Supplementary Figure 2).

```
# match testing by 3-month periods of age
post_ptest$Age_Group <- as.character(floor(post_ptest$Admit_Age_In_Days / 90))</pre>
pre_ptest$Age_Group <- as.character(floor(pre_ptest$Admit_Age_In_Days / 90))</pre>
# bootstrapping post-emergence phase tests
boot_post = bootstrap_df_by_age(post_ptest, pre_ptest)
# plot the pre-pandemic, post-emergence, and test-adjusted age distributions
age_tests <- rbind(data.frame(Age = pre_ptest$Admit_Age_In_Days/365, Group = "Pre-Pandemic"),</pre>
                   data.frame(Age = post_ptest$Admit_Age_In_Days/365, Group = "Post-Emergence"),
                   data.frame(Age = boot_post$Admit_Age_In_Days/365, Group = "Post-Emergence\n(Test Adj
age_tests$Group = factor(age_tests$Group, levels = c("Pre-Pandemic", "Post-Emergence", "Post-Emergence")
sf2 <- ggplot(age_tests, aes(x = Group, y = Age)) +</pre>
  geom_boxplot(outlier.shape = NA, fill = "grey") +
  labs(x = "", y = "Age of Tested Individuals (Years)") +
  theme_classic() +
  coord_cartesian(ylim = c(0, 19)) +
  geom_signif(comparisons = list(c("Pre-Pandemic", "Post-Emergence"),
```



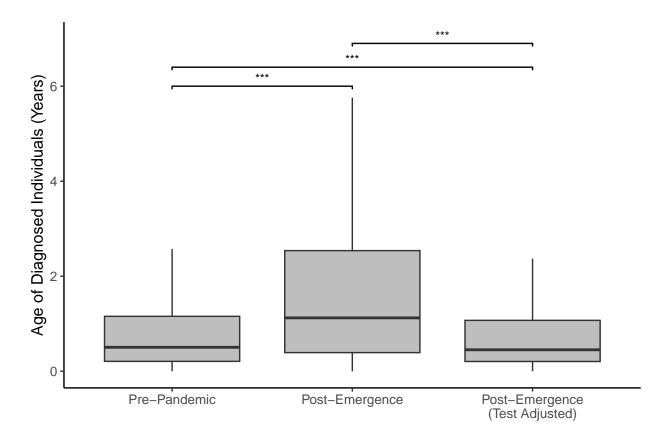
```
ggsave("figs/suppfig2.pdf", plot = sf2, width = 6, height = 6)
rm(age_tests, sf2)
```

Assess whether the distribution of patients diagnosed with RSV would have changed if the age of patients tested for RSV remained constant from the pre-pandemic to the post-emergence phase. Conduct 50 replicates of bootstrapping to ensure that the results are robust to sampling error.

```
# isolate positive bootstrapped tests
  pt_post = boot_post %>% filter(Discharge_ID %in% pt$Discharge_ID)
  # append age range to df
  age_sim[nrow(age_sim)+1, ] <- c(quantile(pt_post$Admit_Age_In_Days))}
# check for consistency across replicates
summary(age_sim/30.44) # convert to months
##
         min
                                    median
                      q1
                                                       q3
                                                                      max
##
   Min.
           :0
                Min.
                       :2.431
                                Min.
                                       :5.355
                                                 Min.
                                                        :12.55
                                                                 Min.
                                                                        :214.4
                                                                 1st Qu.:215.3
   1st Qu.:0
                1st Qu.:2.431
                                1st Qu.:5.420
                                                 1st Qu.:12.68
## Median :0
               Median :2.464
                               Median :5.420
                                                 Median :12.71
                                                                 Median :215.7
## Mean
           :0
               Mean
                       :2.453
                               Mean
                                       :5.430
                                                 Mean
                                                        :12.71
                                                                 Mean
                                                                        :215.6
## 3rd Qu.:0
                3rd Qu.:2.464
                                3rd Qu.:5.453
                                                 3rd Qu.:12.75
                                                                 3rd Qu.:215.9
##
  Max.
           :0
                Max.
                       :2.497
                                Max.
                                       :5.453
                                                 Max.
                                                        :12.84
                                                                 Max.
                                                                         :215.9
rm(age_sim)
Plot the pre-pandemic, post-emergence, and test-adjusted age distributions of patients with a laboratory-
confirmed diagnosis of RSV (Figure 2B).
# create plotting df
age_adj_df <- rbind(data.frame(Age = (pre_pt %>% filter(Num_Tests > 0))$Admit_Age_In_Days/365, Group =
                    data.frame(Age = (post_pt %>% filter(Num_Tests > 0))$Admit_Age_In_Day/365, Group =
                    data.frame(Age = pt_post$Admit_Age_In_Day/365, Group = "Post-Emergence\n(Test Adjus
age_adj_df$Group = factor(age_adj_df$Group, levels = c("Pre-Pandemic", "Post-Emergence", "Post-Emergence")
# print summary statistics for each group
tapply(12*age_adj_df$Age, age_adj_df$Group, quantile) # convert to months
## $'Pre-Pandemic'
##
                     25%
                                50%
                                            75%
                                                      100%
           0%
##
     0.000000
                2.498630
                           6.049315 13.873973 215.736986
##
## $'Post-Emergence'
##
          0%
                   25%
                             50%
                                        75%
                                                 100%
##
     0.00000
               4.70137 13.47945 30.46849 216.09863
##
## $'Post-Emergence\n(Test Adjusted)'
##
           0%
                     25%
                                50%
                                            75%
                                                      100%
     0.000000
                           5.424658 12.854795 216.098630
                2.465753
# plot age distributions
fig2b <- ggplot(age_adj_df, aes(x = Group, y = Age)) +
  geom_boxplot(outlier.shape = NA, fill = "grey") +
  labs(x = "", y = "Age of Diagnosed Individuals (Years)") +
  theme_classic() +
  coord_cartesian(ylim = c(0, 7)) +
  geom_signif(comparisons = list(c("Pre-Pandemic", "Post-Emergence"),
                                  c("Pre-Pandemic", "Post-Emergence\n(Test Adjusted)"),
                                  c("Post-Emergence", "Post-Emergence\n(Test Adjusted)")),
```

test = "wilcox.test",

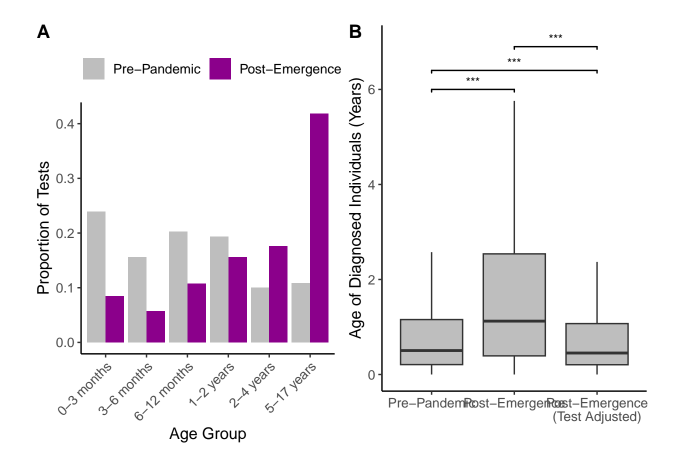
```
map_signif_level = TRUE,
    textsize = 3,
    y_position = c(5.1,5.5,6.0),
    tip_length = 0.003) +
theme(legend.position = "none",
    text = element_text(size = 12))
fig2b
```



```
rm(age_adj_df)
```

Generate Figure 2.

```
fig2 = plot_grid(fig2a, fig2b, nrow = 1, labels = c("A", "B"))
fig2
```



ggsave("figs/fig2.pdf", plot = fig2, width = 11, height = 5)