# Covid19\_Impact\_on\_Hospitals\_visualization.R

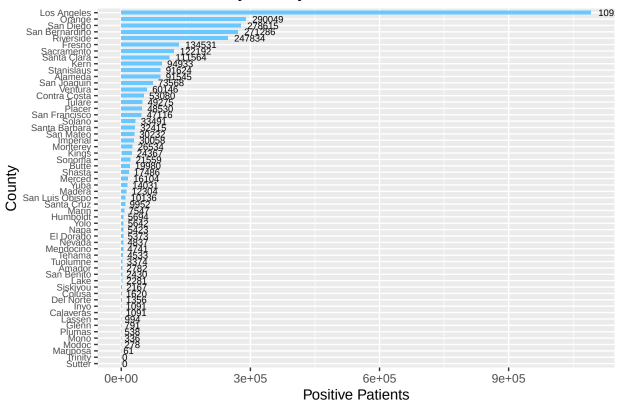
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#### 2022-01-21

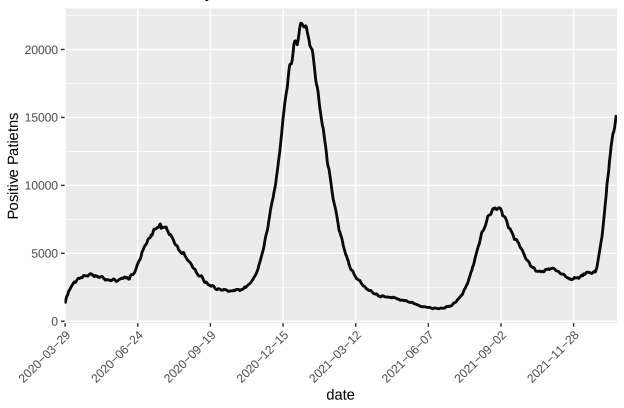
```
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(ggplot2)
library(scales)
setwd("C:/Users/dongj/Desktop/Covid_Cal/The_impact_on_hospitals_in_the_CA")
data <- read.csv("./dataset/covid19hospitalbycounty_011922.csv")</pre>
data_v1 <- read.csv("./dataset/covid19hospitalbycounty_011922.csv")</pre>
positive_patients_by_county<- aggregate(hospitalized_covid_confirmed_patients ~ county,</pre>
                                         data_v1,sum)
positive_patients_by_date<- aggregate(hospitalized_covid_confirmed_patients ~ todays_date,
                                         data_v1,sum)
suspected_patients_by_county<- aggregate(hospitalized_suspected_covid_patients ~ county,</pre>
                                         data v1, sum)
suspected_patients_by_date<- aggregate(hospitalized_suspected_covid_patients ~ todays_date,
                                       data v1, sum)
icu_positive_patients_by_county<- aggregate(icu_covid_confirmed_patients ~ county,</pre>
                                         data_v1,sum)
icu_positive_patients_by_date<- aggregate(icu_covid_confirmed_patients ~ todays_date,
                                       data_v1,sum)
```

```
positive_patients_by_county_graph <- ggplot(data=positive_patients_by_county,</pre>
                                         aes(x = hospitalized_covid_confirmed_patients,
                                             y = reorder(county,
                                                         hospitalized_covid_confirmed_patients,
                                                         sum))
                                    )+
                                     geom_bar(stat = "identity",
                                              width=.6,
                                              position = position_dodge(width = 0.5),
                                              fill = "#69c8ff")+
                                    labs(
                                       title="Positive Patients by County",
                                       x = "Positive Patients",
                                      y = "County"
                                    theme(axis.text.y = element_text(size = 7)) +
                                     geom_text(aes(label= hospitalized_covid_confirmed_patients),
                                               hjust = -0.2,
                                               size = 2.5,
                                               position = position_dodge(width = 1),
                                               inherit.aes = TRUE)
options(repr.plot.width = 14, repr.plot.height = 8)
positive_patients_by_county_graph
```

# Positive Patients by County

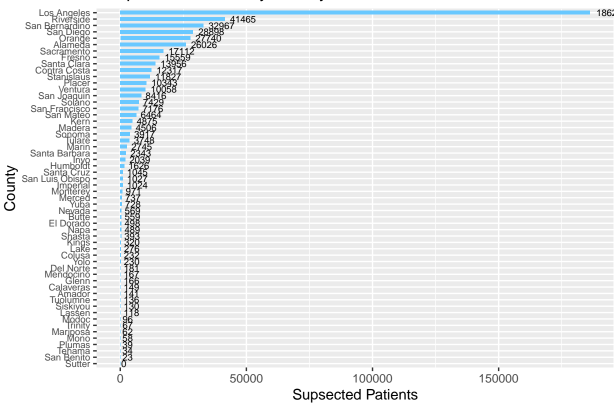


#### Positive Patients by Date

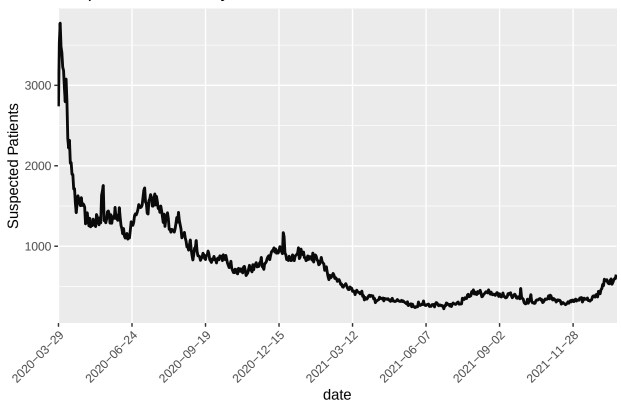


```
suspected_patients_by_county_graph <- ggplot(data=suspected_patients_by_county,</pre>
                                             aes(x = hospitalized_suspected_covid_patients,
                                                 y = reorder(county,
                                                             hospitalized_suspected_covid_patients,
                                                             sum))
                                       )+
                                       geom_bar(stat = "identity",
                                                width=.6,
                                                position = position_dodge(width = 0.5),
                                                fill = "#69c8ff")+
                                       labs(
                                         title="Suspected Patients by County",
                                         x = "Supsected Patients",
                                         y = "County"
                                       theme(axis.text.y = element_text(size = 7)) +
                                       geom_text(aes(label= hospitalized_suspected_covid_patients),
                                                 hjust = -0.2,
                                                 size = 2.5,
                                                 position = position_dodge(width = 1),
                                                 inherit.aes = TRUE)
options(repr.plot.width = 14, repr.plot.height = 8)
suspected_patients_by_county_graph
```

# Suspected Patients by County

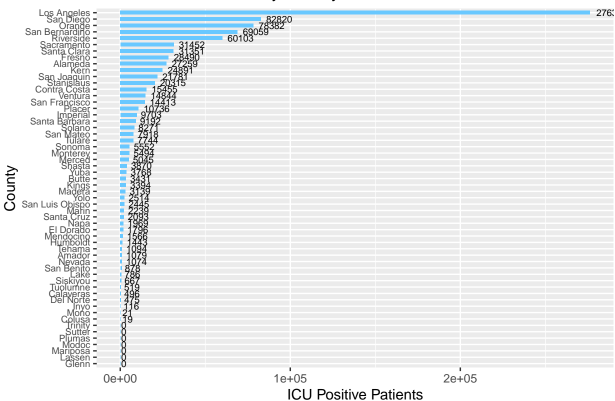


# Suspected Patients by Date

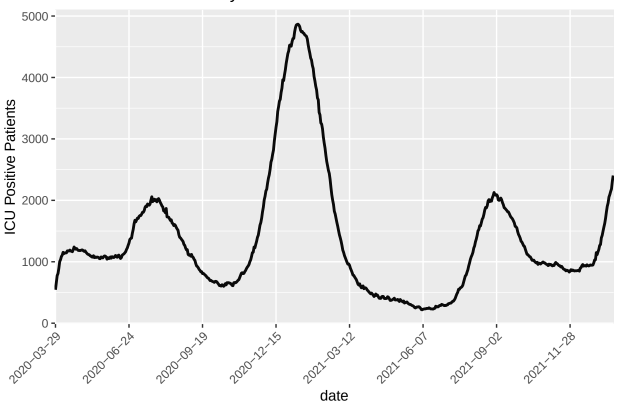


```
icu_positive_patients_by_county_graph <- ggplot(data=icu_positive_patients_by_county,</pre>
                                              aes(x = icu_covid_confirmed_patients,
                                                  y = reorder(county,
                                                              icu_covid_confirmed_patients,
                                                              sum))
                                         )+
                                         geom_bar(stat = "identity",
                                                  width=.6,
                                                  position = position_dodge(width = 0.5),
                                                  fill = "#69c8ff")+
                                        labs(
                                           title="ICU Positive Patients by County",
                                           x = "ICU Positive Patients",
                                           y = "County"
                                         theme(axis.text.y = element_text(size = 7)) +
                                         geom_text(aes(label= icu_covid_confirmed_patients),
                                                   hjust = -0.2,
                                                   size = 2.5,
                                                   position = position_dodge(width = 1),
                                                   inherit.aes = TRUE)
options(repr.plot.width = 14, repr.plot.height = 8)
icu_positive_patients_by_county_graph
```

# ICU Positive Patients by County

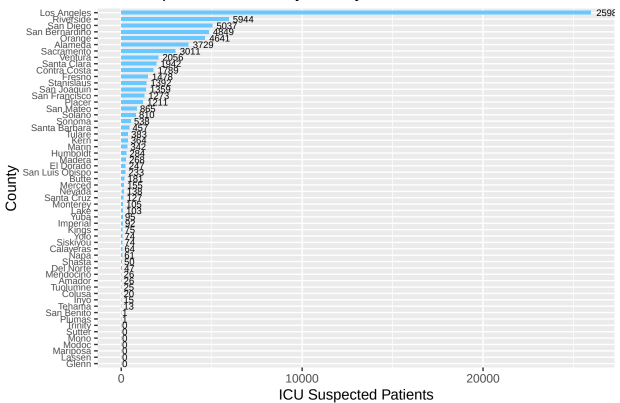


### ICU Positive Patients by Date

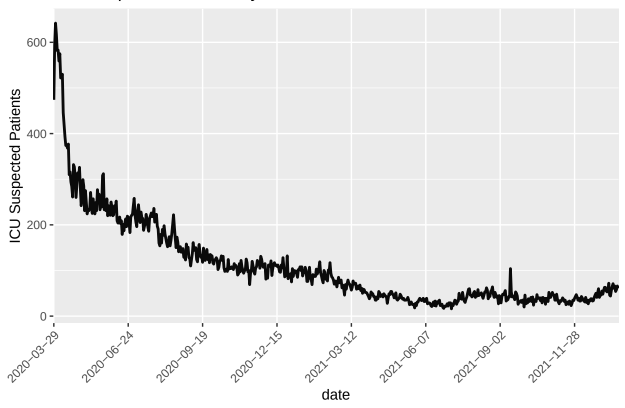


```
icu_suspected_patients_by_county_graph <- ggplot(data=icu_suspected_patients_by_county,</pre>
                                                 aes(x = icu_suspected_covid_patients,
                                                     y = reorder(county,
                                                                 icu_suspected_covid_patients,
                                                                 sum))
                                           )+
                                           geom_bar(stat = "identity",
                                                    width=.6,
                                                    position = position_dodge(width = 0.5),
                                                    fill = "#69c8ff")+
                                           labs(
                                             title="ICU Suspected Patients by County",
                                             x = "ICU Suspected Patients",
                                             y = "County"
                                           theme(axis.text.y = element_text(size = 7)) +
                                           geom_text(aes(label= icu_suspected_covid_patients),
                                                     hjust = -0.2,
                                                     size = 2.5,
                                                     position = position_dodge(width = 1),
                                                     inherit.aes = TRUE)
options(repr.plot.width = 14, repr.plot.height = 8)
icu_suspected_patients_by_county_graph
```

# ICU Suspected Patients by County

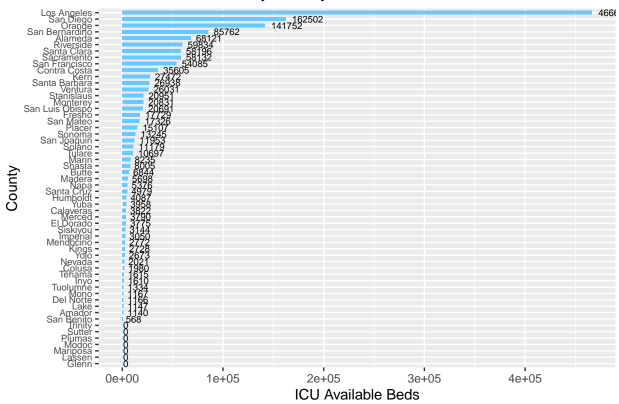


# ICU Suspected Patients by Date



```
icu_available_beds_by_county_graph <- ggplot(data=icu_available_beds_by_county,</pre>
                                                  aes(x = icu_available_beds,
                                                      y = reorder(county,
                                                                  icu_available_beds,
                                                                   sum))
                                    )+
                                       geom_bar(stat = "identity",
                                                width=.6,
                                                position = position_dodge(width = 0.5),
                                                fill = "#69c8ff")+
                                      labs(
                                         title="ICU Available Beds by County",
                                        x = "ICU Available Beds",
                                        y = "County"
                                       theme(axis.text.y = element_text(size = 7)) +
                                       geom_text(aes(label= icu_available_beds),
                                                 hjust = -0.2,
                                                 size = 2.5,
                                                 position = position_dodge(width = 1),
                                                 inherit.aes = TRUE)
options(repr.plot.width = 14, repr.plot.height = 8)
icu_available_beds_by_county_graph
```

# ICU Available Beds by County



# ICU Available Beds by Date

