Group 4 Report

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TidyR data setup

The data is composed of two datasets with other lapping patients (see ID), exam.dataset.txt consists of PCR rdts for COVID19 during the 2020 pandemic, data includes information on patients including clinic, gender, test results week of pandemic, and exam_joindata.txt containing endpoint titer data from some of the patients (information can be found in the files codebook_exam_data.html, exam.descr.md). Data files were joined and processed as per conventions of tidyR for later visulization and analysis.

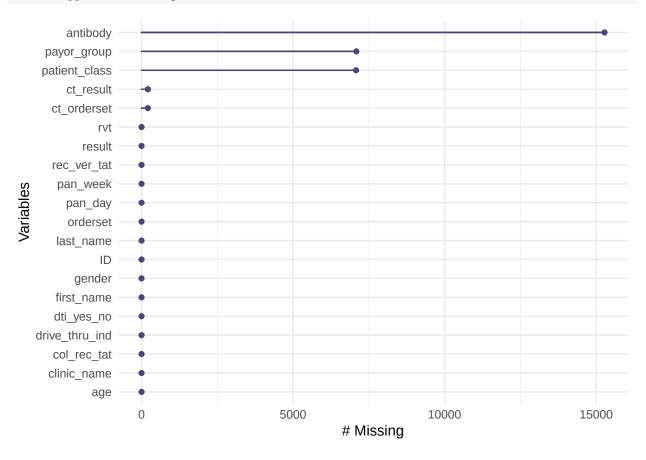
An overview of the data follows

summary(complete_data)

```
##
          ID
                                                        first_name
                                           gender
                           age
##
    Min.
                                0.00
                                        female:7832
                                                       Length: 15524
                     Min.
                     1st Qu.:
##
    1st Qu.: 2330
                                2.00
                                        male :7692
                                                       Class : character
    Median: 5268
                     Median:
                                9.00
                                                       Mode
                                                             :character
##
    Mean
            : 5571
                     Mean
                             : 14.19
##
    3rd Qu.: 8636
                     3rd Qu.: 18.00
            :12346
##
                             :138.00
    Max.
                     Max.
##
     last_name
##
                                    clinic name
                                                         result
                                                                      drive_thru_ind
##
    Length: 15524
                         clinical lab
                                           :7500
                                                    invalid:
                                                                301
                                                                      0:7537
                         emergency dept
                                           :3413
                                                                      1:7987
##
    Class : character
                                                    negative:14358
##
    Mode :character
                        oncology day hosp: 533
                                                    positive:
                                           : 294
##
                         nicu
##
                         laboratory
                                           : 270
##
                         picu
                                           : 261
##
                         (Other)
                                           :3253
##
      ct_result
                         orderset
                                                     payor_group
##
            :14.05
                             :0.0000
                                        commercial
                                                            :3726
    Min.
                     Min.
##
    1st Qu.:45.00
                     1st Qu.:0.0000
                                        government
                                                            :3644
    Median :45.00
##
                     Median :1.0000
                                        unassigned
                                                            : 733
##
    Mean
            :44.12
                             :0.6952
                                        self pay
                                                             216
                     Mean
##
    3rd Qu.:45.00
                     3rd Qu.:1.0000
                                        medical assistance:
                                                              84
##
    Max.
            :45.00
                             :1.0000
                                        (Other)
                                                              34
                     Max.
    NA's
            :209
##
                                        NA's
                                                            :7087
                                      pan_day
##
                  patient class
                                                      rec ver tat
##
    inpatient
                          :3438
                                  Min.
                                          : 4.00
                                                            :-18.600
                                                     Min.
##
    emergency
                          :1378
                                  1st Qu.: 38.00
                                                     1st Qu.: 4.000
    not applicable
                          :1096
                                  Median: 65.00
                                                     Median :
                                                               5.000
##
##
    outpatient
                          : 973
                                  Mean
                                          : 63.21
                                                     Mean
                                                               5.639
##
    recurring outpatient: 795
                                  3rd Qu.: 87.00
                                                     3rd Qu.:
                                                               6.200
##
    (Other)
                          : 767
                                  Max.
                                          :107.00
                                                     Max.
                                                             :218.200
    NA's
                          :7077
##
```

```
pan_week
##
     col_rec_tat
                                                              dti_yes_no
                           rvt
##
                0.00
                       Length: 15524
                                                : 0.5714
                                                             Length: 15524
    Min.
         :
                                           Min.
##
    1st Qu.:
                0.70
                       Class : character
                                           1st Qu.: 5.4286
                                                              Class : character
                1.90
                                           Median : 9.2857
##
   Median :
                       Mode :character
                                                             Mode :character
##
    Mean
                7.22
                                           Mean
                                                  : 9.0298
                3.60
##
                                           3rd Qu.:12.4286
    3rd Qu.:
##
   Max.
           :61370.20
                                           Max.
                                                  :15.2857
##
##
     ct_orderset
                      antibody
   Min. : 0.0
                           : 21.48
##
                   Min.
   1st Qu.: 0.0
                   1st Qu.: 63.30
   Median:45.0
                   Median :103.52
##
                          :109.99
##
    Mean
           :30.6
                   Mean
##
    3rd Qu.:45.0
                   3rd Qu.:156.95
##
   Max.
           :45.0
                   Max.
                           :199.89
##
   NA's
           :209
                   NA's
                           :15275
```

naniar::gg_miss_var(complete_data)

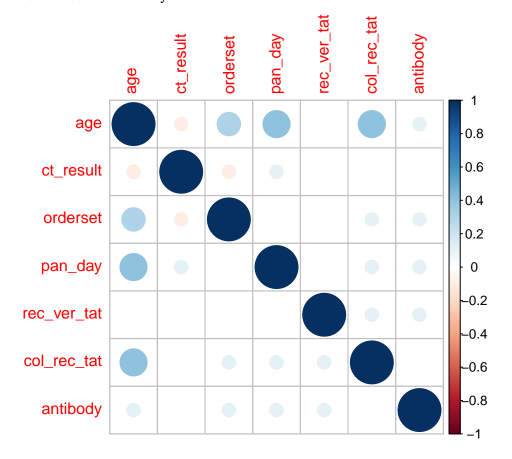


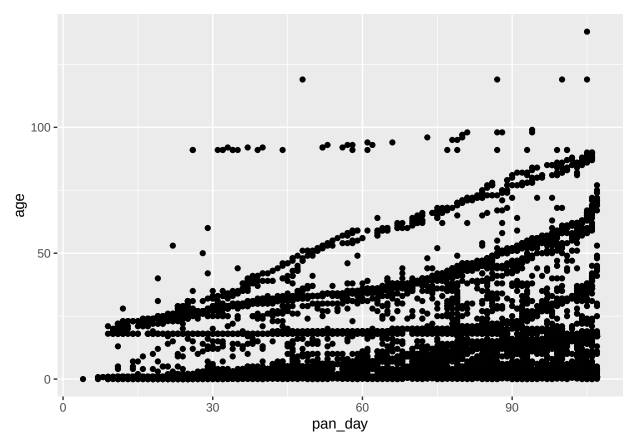
Descriptive plots

Visulization was done using a number of plots. Significant correlations are observiable with age, and orderset and days since the pandemic. Indicating early in the pandemic individuals seen in the clinic were usually young adults and children

```
## # A tibble: 7 x 8
    rowname
                       age ct_result orderset
                                                pan_day rec_ver_tat col_rec_tat
```

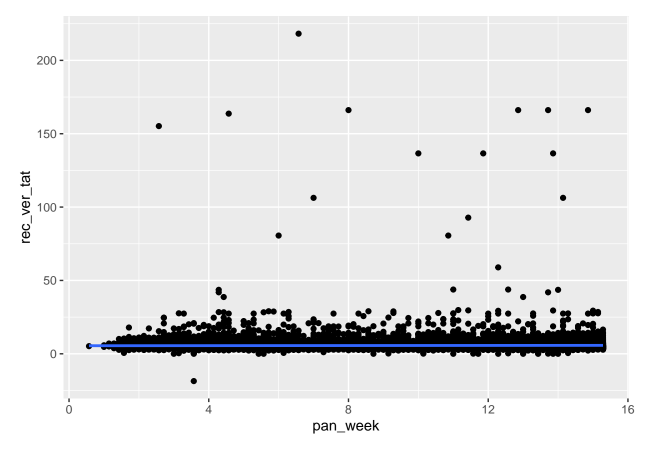
##	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>		
##	1 age	0	4.54e-17	9.34e-151	3.95e-323	0.0666	0.844		
## :	2 ct_result	4.54e- 17	0	3 e- 11	7.86e- 1	0.0829	0.849		
## 3	3 orderset	9.34e-151	3 e-11	0	7.38e- 1	0.000108	0.158		
## 4	4 pan_day	3.95e-323	7.86e- 1	7.38e- 1	0	0.125	0.809		
## !	5 rec_ver_tat	6.66e- 2	8.29e- 2	1.08e- 4	1.25e- 1	0	0.854		
## (6 col_rec_tat	8.44e- 1	8.49e- 1	1.58e- 1	8.09e- 1	0.854	0		
## '	7 antibody	4.16e- 2	9.53e- 1	2.89e- 1	4.05e- 2	0.0704	0.623		
## # i 1 more variable: antibody <dbl></dbl>									





Suprisingly over the course of the pandemic there was no indication that the wait time from collection of patient sample to PCR test result was reduced.

##	# A	tibble:	15 x 4		
##		pan_week	mean_value	std	<pre>'median(rec_ver_tat)'</pre>
##		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
##	1	1	5.51	1.57	5.3
##	2	2	5.27	2.00	4.8
##	3	3	5.62	5.44	5
##	4	4	5.50	3.19	5
##	5	5	5.57	5.07	5
##	6	6	5.69	3.61	5.1
##	7	7	5.73	7.81	4.9
##	8	8	5.66	5.97	4.9
##	9	9	5.48	2.59	4.9
##	10	10	5.65	4.68	5
##	11	11	5.73	4.17	5
##	12	12	5.57	4.73	4.9
##	13	13	5.59	5.22	4.9
##	14	14	5.82	6.88	5
##	15	15	5.81	5.62	5
##	'ge	om_smootl	h()' using	formula	a = 'y ~ x'

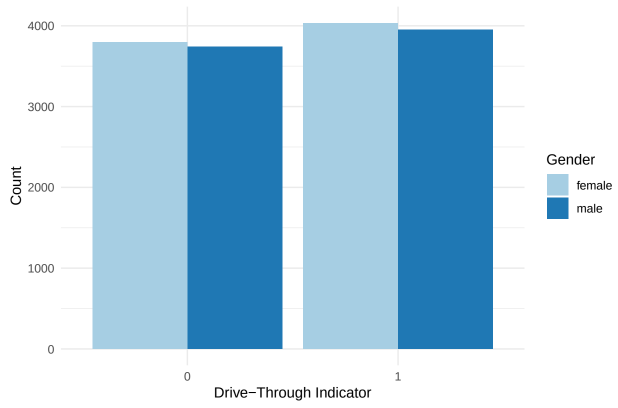


We observe slightly more females than males in the cleaned data set. The bar plot below indicates that more people from both genders took the test at the drive-through than those who did not. However, from observation, the difference between genders is slightly larger among those who took the test at the drive-through than those who did not. To examine the relationship between gender and drive-through test participation, we conducted a Chi-squared test. With a p-value of 0.8728, we fail to reject the null hypothesis, suggesting no significant association exists between gender and drive-through test participation in this data set.

```
dt_and_gender <- table(complete_data$gender, complete_data$drive_thru_ind)
# Create dt_and_gender as a data frame from the table
dt_and_gender_df <- as.data.frame(dt_and_gender)</pre>
print(chisq.test(dt_and_gender))
##
##
   Pearson's Chi-squared test with Yates' continuity correction
##
## data: dt_and_gender
## X-squared = 0.02564, df = 1, p-value = 0.8728
# Rename columns for clarity
colnames(dt_and_gender_df) <- c("Gender", "Drive_Through", "Count")</pre>
# plot using ggplot
ggplot(dt_and_gender_df, aes(x = Drive_Through, y = Count, fill = Gender)) +
  geom_bar(stat = "identity", position = "dodge") +
  labs(title = "Gender Distribution of Drive-Through Test Participants",
       x = "Drive-Through Indicator",
```

```
y = "Count") +
scale_fill_manual(values = c("#a6cee3", "#1f78b4")) + # Custom colors
theme_minimal()
```

Gender Distribution of Drive-Through Test Participants

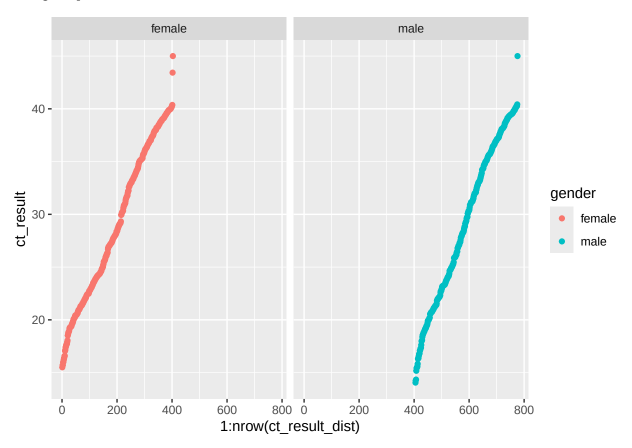


A comparison of cycle threshold values between men and women shows a similar distribution in the number of reaction cycles. However, males showed the lowest cycle threshold value compared to females. Despite this, the majority of threshold values exceeded 40 cycles for both genders. The p-value of 0.272 from the one-way ANOVA test means we fail to reject the null hypothesis, suggesting that gender does not affect the cycle threshold value.

```
ct_result_cat <- cut(complete_data$ct_result, c(10, 20, 30, 40, 50))
summary(ct_result_cat)
## (10,20] (20,30] (30,40] (40,50]
                                       NA's
##
        92
               346
                        415
                              14462
                                        209
ct_result_dist <- complete_data %>%
  group_by(gender) %>%
  count(ct_result)
ggplot(ct_result_dist,
         x = 1:nrow(ct_result_dist),
         y = ct_result,
         group = gender,
         color = gender)
) +
```

```
geom_point() +
facet_wrap(vars(gender))
```

Warning: Removed 2 rows containing missing values or values outside the scale range
('geom_point()').



ct_result_gender_anova <- aov(complete_data\$ct_result ~ complete_data\$gender, data = complete_data\$
summary(ct_result_gender_anova)</pre>

```
## Complete_data$gender 1 19 19.11 1.206 0.272 ## Residuals 15313 242691 15.85 ## 209 observations deleted due to missingness
```

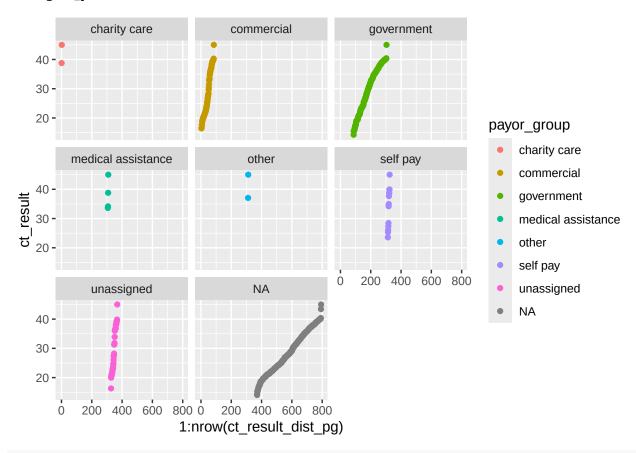
The distribution of patients by payor group shows that the most common groups were commercial and government, with a significant proportion of patients listed as NA or unassigned. Despite this, a similar pattern was observed between government and commercial patients. In contrast, the minimum cycle threshold value for charity care, medical assistance, and other self-pay patients was higher than for the former groups. The one-way ANOVA test result is highly significant, indicating that the cycle threshold value is influenced by the payor group.

```
ct_result_dist_pg <- complete_data %>%
  group_by(payor_group) %>%
  count(ct_result)

ggplot(
  ct_result_dist_pg,
```

```
aes(
    x = 1:nrow(ct_result_dist_pg),
    y = ct_result,
    group = payor_group,
    color = payor_group
    )
) +
geom_point() +
facet_wrap(vars(payor_group))
```

Warning: Removed 6 rows containing missing values or values outside the scale range
('geom_point()').



ct_result_pg_anova <- aov(complete_data\$ct_result ~ complete_data\$payor_group, data = complete_data)
summary(ct_result_pg_anova)</pre>

```
## Df Sum Sq Mean Sq F value Pr(>F)

## complete_data$payor_group 6 672 111.9 9.1 6.03e-10 ***

## Residuals 8296 102040 12.3

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

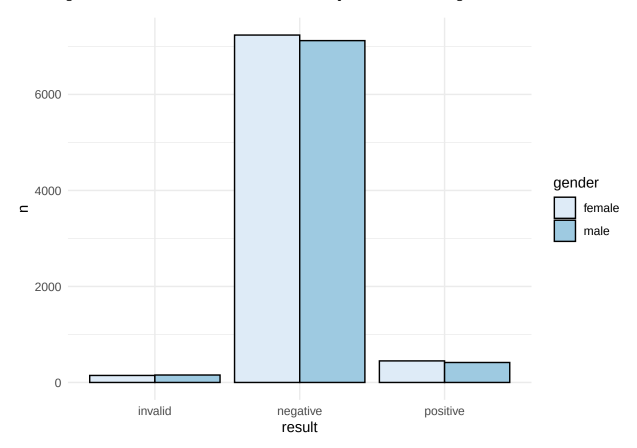
## 7221 observations deleted due to missingness
```

The distribution of results by gender shows a similar pattern for both, with women comprising a slightly larger proportion in each result category except for the invalid category. The chi-square test yielded a p-value of 0.5481, meaning that we fail to reject the null hypothesis. This suggests that gender does not influence the test result.

```
tests_by_gender <- complete_data %>%
  group_by(gender) %>%
  count(result)
tests_by_gender
## # A tibble: 6 x 3
## # Groups: gender [2]
    gender result
     <fct> <fct>
                     <int>
## 1 female invalid
                       146
## 2 female negative 7237
## 3 female positive
                       449
## 4 male
                       155
           invalid
## 5 male
           negative 7121
## 6 male
           positive
ggplot(data = tests_by_gender, aes(x = result, y = n, fill = gender)) +
  geom_bar(stat = "identity", color="black", position=position_dodge())+
  theme_minimal() +
  scale_fill_manual(values=c('#999999','#E69F00')) +
# Use brewer color palettes
  scale_fill_brewer(palette="Blues")
```

Scale for fill is already present.

Adding another scale for fill, which will replace the existing scale.



```
test_and_gender <- table(complete_data$gender, complete_data$result)</pre>
test_and_gender
##
##
            invalid negative positive
##
     female
                 146
                         7237
                                    449
##
     male
                 155
                         7121
                                    416
#chi-square test
test_and_gender_df <- as.data.frame(test_and_gender)</pre>
print(chisq.test(test_and_gender))
##
##
    Pearson's Chi-squared test
##
## data: test_and_gender
## X-squared = 1.2028, df = 2, p-value = 0.5481
```

Statistics

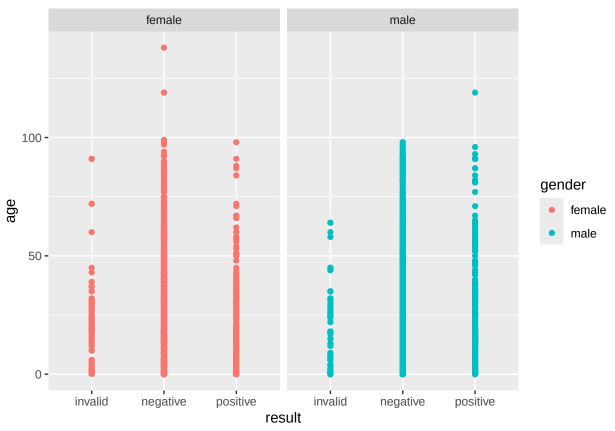
wilcox test indicated that the more tests from a single patient does not associate with increase positive test, in fact the opposite trend was noted.

```
## # A tibble: 1 x 4
##
     statistic p.value method
                                                                           alternative
##
         <dbl>
                   <dbl> <chr>
                                                                           <chr>>
       4979785 8.14e-11 Wilcoxon rank sum test with continuity correct~ two.sided
While drive-through use was associated with an increase in positive tests
##
##
    Pearson's Chi-squared test with simulated p-value (based on 2000
##
    replicates)
##
## data: positive_visits$n and positive_visits$positive
## X-squared = 44.749, df = NA, p-value = 0.01499
```

A visual inspection of the distribution suggests that age may be associated with the test result, as there appears to be a positive correlation between age distribution and test outcomes. However, the dataset is skewed toward younger individuals, so that the correlation may be due to the age structure. To determine whether this association holds statistically, we performed an ANOVA test. The results showed a significant p-value, indicating that we cannot rule out the possibility of an association between age and test result.

```
## complete data$result: invalid
##
     0-9 10-19 20-29 30-39 40-49 50-59 60-69 70-79 80-89 90-99
                                                                   100+
                                                                         NA's
##
     143
            60
                  49
                         29
                                8
                                       1
                                             4
                                                   2
                                                          0
                                                                2
                                                                      0
                                                                             3
##
## complete data$result: negative
     0-9 10-19 20-29 30-39 40-49 50-59 60-69 70-79 80-89 90-99
##
                                                                   100+
                                                                         NA's
  7325 3675
                 935
                        968
                              489
                                    370
                                           183
                                                  73
                                                        56
                                                               32
                                                                          248
```

```
## complete_data$result: positive
    0-9 10-19 20-29 30-39 40-49 50-59 60-69 70-79 80-89 90-99 100+ NA's
##
    296 278 85
                     99 22
                               30
                                     16 4 9 13
                                                        1 12
ggplot(complete_data,
      aes(
       x = result,
       y = age,
       group = gender,
        color = gender)
) +
 geom_point() +
 facet_wrap(vars(gender))
```



```
# ANOVA
result_age_aov <- aov(age ~ result, data = complete_data)
result_age_aov

## Call:
## aov(formula = age ~ result, data = complete_data)
##
## Terms:
## result Residuals
## Sum of Squares 22781 4189232
## Deg. of Freedom 2 15521
##</pre>
```

```
## Residual standard error: 16.42886
## Estimated effects may be unbalanced
summary(result_age_aov)
##
                       Sum Sq Mean Sq F value Pr(>F)
## result
                    2
                        22781
                                 11391
                                           42.2 <2e-16 ***
## Residuals
                15521 4189232
                                   270
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
The table below shows the cycle threshold (Ct) results and distribution of test outcomes. As seen earlier,
most individuals in this dataset have a Ct value between 40 and 50. Interestingly, this range also has the
lowest number of positive test results compared to those with lower Ct values. Additionally, the ANOVA
test shows a significant result, suggesting a difference in the distribution of Ct values across different result
outcome groups.
result_ct_table <- complete_data %>%
  mutate(ct_result_cat = cut(ct_result, breaks = c(10, 20, 30, 40, 50))) %>%
  count(result, ct_result_cat) %>%
  pivot_wider(names_from = ct_result_cat, values_from = n, values_fill = 0)
# View the table
print(result_ct_table)
## # A tibble: 3 x 6
                          'NA' '(30,40]' '(10,20]' '(20,30]'
##
              '(40,50]'
     result
     <fct>
                   <int> <int>
                                    <int>
                                               <int>
                                                          <int>
## 1 invalid
                     229
                             72
                                         0
                                                    0
                                                               0
                                                               0
## 2 negative
                   14213
                            132
                                        13
                                                    0
## 3 positive
                      20
                              5
                                       402
                                                  92
                                                            346
```

ct_result_ct_anova <- aov(complete_data\$ct_result ~ complete_data\$result, data = complete_data)