

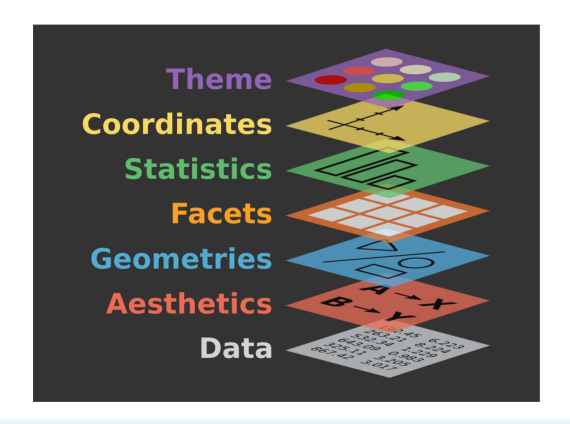
Data visualization

Part II

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02/25/2025

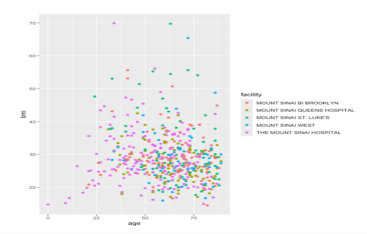
Let's recap



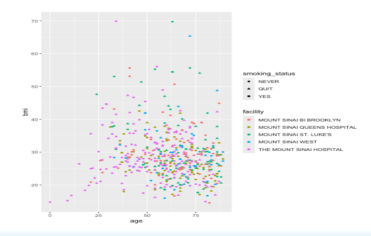
Let's recap

- What geometry would you use for plotting two numerical variables?
- What geometry would you use for plotting categorical vs continuous variables?
- How would you include a third variable in the plot?

Plotting num vs num vs cat



Plotting num vs num vs cat vs cat

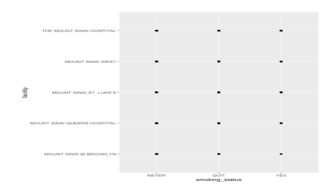


Your turn!

- Create a plot of bmi vs age
- Color by ethnicity
- Add shapes by sex

Plotting cat vs cat?

• How would you compare smoking_status vs facility?



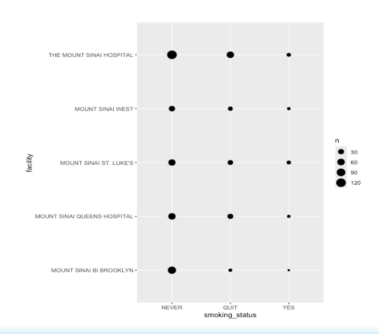
Plotting cat vs cat vs num

• First create a summarized tibble

```
count smoking <- sinai covid %>%
  group by(facility) %>%
  count(smoking status)
count smoking[1:3,]
## # A tibble: 3 × 3
## # Groups: facility [1]
                             smoking_status
##
   facility
##
    <chr>
                             <chr>
                                            <int>
## 1 MOUNT SINAI BI BROOKLYN NEVER
                                               78
## 2 MOUNT SINAI BI BROOKLYN OUIT
## 3 MOUNT SINAI BI BROOKLYN YES
```

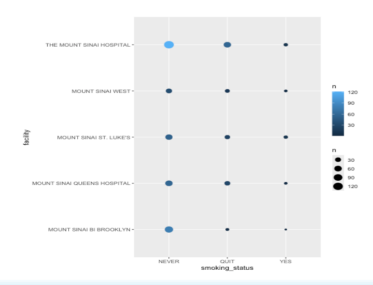
• Plot

```
ggplot(count_smoking,
    aes(x = smoking_status,
    y = facility,
    size = n)) +
geom_point()
```



Plotting cat vs cat vs num

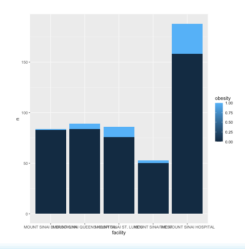
```
ggplot(count_smoking,
    aes(x = smoking_status,
    y = facility,
    size = n, color = n)) +
  geom_point()
```



Let's practice

- Count the number of patients with obesity (0 and 1) per facility.
- Create a dots plot, mapping the size of the dots to the obesity counts.

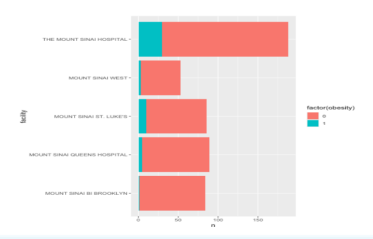
An alternative plot



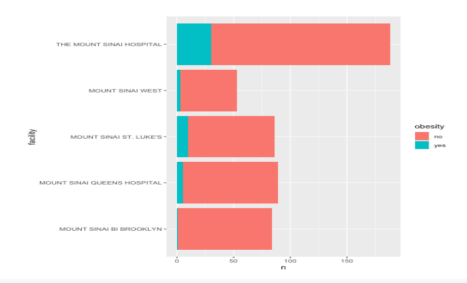
How do we fix the label?

• Let's treat the variable Obesity as factor

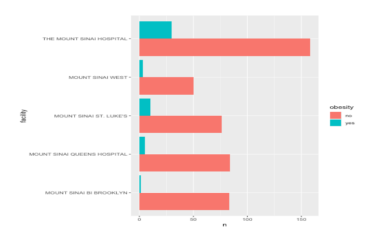
```
ggplot(ob_count,
    aes(x = n,
        y = facility,
        fill = factor(obesity))) +
    geom_col()
```



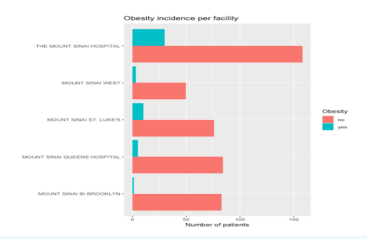
Alternatively, replace the values



Splitting the bars



Adding titles



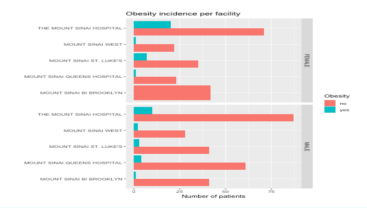
Your turn

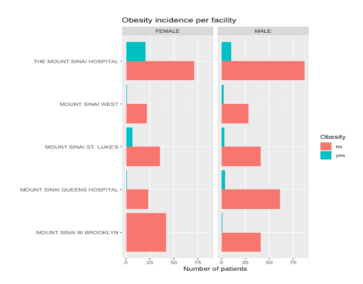
- Create a summarizing tibble with the number of patients per ethnicity and asthma status.
- Use the summarized tibble to create a bar plot, using a position dodge and coloring by asthma status.
- Add a title to the plot, modify the axis titles, and change the legend title.

Facets

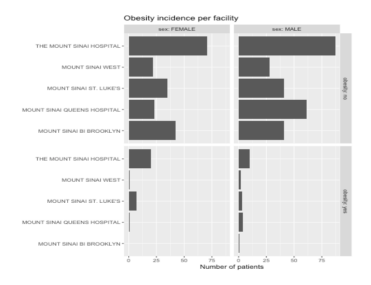
```
ob count <- sinai covid %>%
  group_by(facility, sex) %>%
  count(obesity) %>%
  mutate(obesity = case_when(obesity == 0 ~ "no",
                             obesity == 1 \sim "yes"))
ob count [1:3,]
## # A tibble: 3 × 4
## # Groups: facility, sex [2]
   facility
                                    obesity
                             sex
    <chr>
                             <chr>
                                    <chr> <int>
## 1 MOUNT SINAI BI BROOKLYN FEMALE no
                                               42
## 2 MOUNT SINAI BI BROOKLYN MALE
                                              41
                                    no
## 3 MOUNT SINAI BI BROOKLYN MALE
                                    yes
```

Facet grid





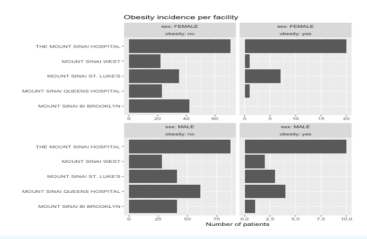
```
ggplot(ob_count,
    aes(x = n, y = facility)) +
geom_col() +
labs(title = "Obesity incidence per facility",
    x = "Number of patients",
    y = "") +
facet_grid(rows = vars(obesity),
    cols = vars(sex),
    labeller = label_both)
```



Let's practice!

- Count the number of patients depending on their smoking status per ethnic group and asthma status.
- Represent the data on a column plot, color them by asthma status.
- Add a plot title, axis title and change the legend title to remove the underscores.
- Split the plots in columns and rows by smoking and asthma status.

Facet wrap



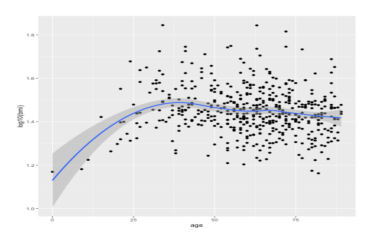
Let's practice!

• Create the previous plot again, but use facet_wrap instead of facet_grid.

Statistical transformations

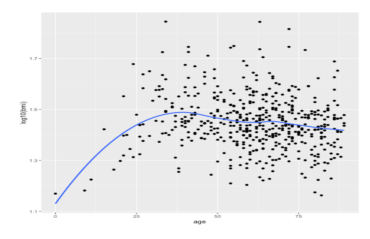
```
sinai_covid %>%
  ggplot(aes(x = age, y = log10(bmi))) +
  geom_point() +
  geom_smooth()
```

`geom_smooth()` using method = 'loess' and formula = 'y \sim x'



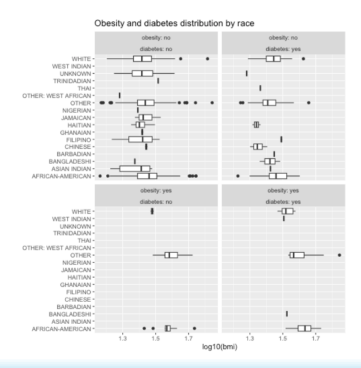
```
sinai_covid %>%
  ggplot(aes(x = age, y = log10(bmi))) +
  geom_point() +
  geom_smooth(se = FALSE)
```

`geom_smooth()` using method = 'loess' and formula = 'y \sim x'



Your turn!

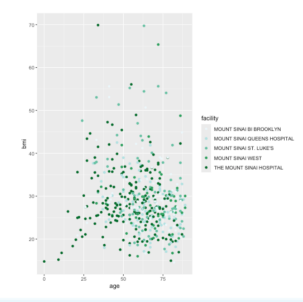
• Write the code to create the following plot:



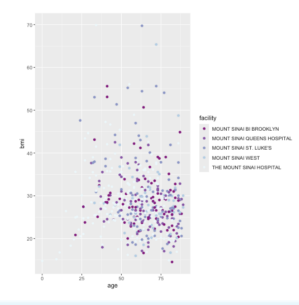
Possible Answer

Scales

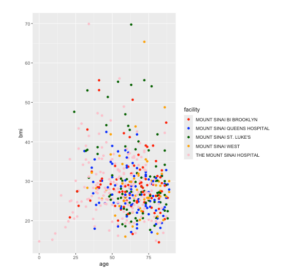
```
ggplot(sinai_covid,
        aes(x = age, y = bmi, color = facility)) +
   geom_point() +
   scale_color_brewer(palette = 2)
```



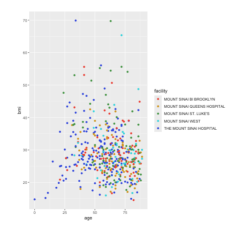
• Invert scale direction



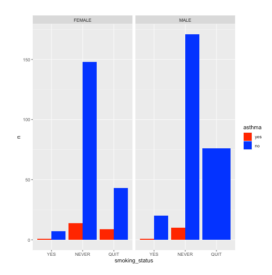
• Using the manual scale



• Using the manual scale. Look for "html color picker" on Google browser



Customized position

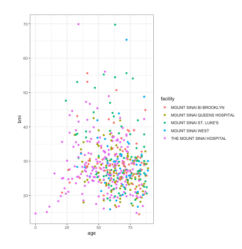


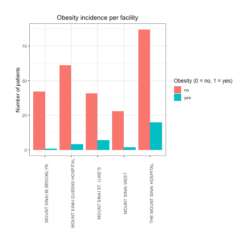
Your turn!

- Plot facility vs bmi.
- Color by diabetes.
- Change the order of facilities to place The Mount Sinai Hospital at the beginning of the x axis.
- Choose your favorite colors to modify the diabetes coloring.

Themes

```
ggplot(sinai_covid,
    aes(x = age, y = bmi,
        color = facility)) +
  geom_point() +
  theme_bw()
```





Your turn!

- Count the number of patients grouping by smoking status, ethnic group and asthma status.
- Plot the number of patients by ethnic group using vertical bars. Color the bars by asthma status.
- Add a plot title, axis titles and modify the legend title. Explore the available themes and use one.
- Modify the angle and size of the text of the axis. Split in several plots by smoking status.

Thanks!

