Data Visualization Analysis

David

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1 Introduction

This tutorial is designed to help you learn data visualization analysis by providing simple and useful information in a way that is easy to follow and understand.

2 Preparation

In order to draw a chart, we need to include the required packages for visualization and dataset. For example, ggplot2 package is for drawing charts and gcookbook is for using pg_mean dataset, but in the next section we will use more dataset from gcookbook.

```
library(ggplot2)
library(gcookbook)
```

3 Bar chart

In this section, we will draw a bar chart using pg_mean dataset. The dataset has two columns: group, weight.

pg_mean

```
group weight
1 ctrl 5.032
2 trtl 4.661
3 trt2 5.526
```

This dataset compares the weight across three groups:

- ctrl: Control group (baseline, weight = 5.032).
 trt1: Treatment 1 group (weight = 4.661).
- trt2: Treatment 2 group (weight = 5.526).

Below graph initializes a ggplot with the dataset pg_mean.

```
aes(x = group, y = weight) specifies the aesthetics:
```

- x = group: Assign the group variable to the x-axis (categorical data, such as ctrl, trt1, trt2).
- y = weight: Assign the weight variable to the y-axis (numerical data).

geom_col():

- Adds a column geometry to the plot.
- geom_col() creates bars where the height of each bar corresponds to the value of weight for each group.

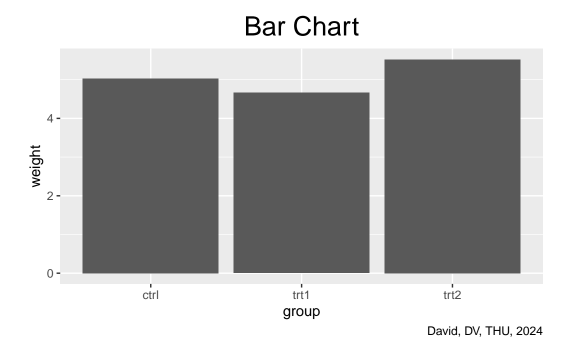
labs() function in ggplot2 is used to customize labels in a plot.

- title: To specifies the main title of the plot.
- caption: To Add a additional information.
- x: To change x label name.
- y: To change y label name.

theme() function in ggplot2 is used to customize the appearance of a plot.

- plot.title: To control the main title.
- element_text(): To customize the appearance of text elements in a plot.
 - hjust: To adjust the title position.
 - size: To control size of the title.

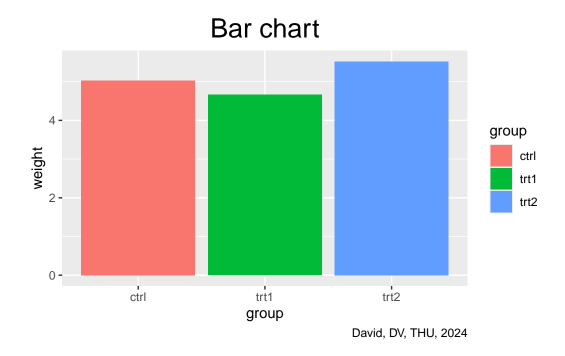
```
ggplot(pg_mean, aes(x = group, y = weight)) +
geom_col() +
labs(title = "Bar Chart", caption = 'David, DV, THU, 2024') +
theme(plot.title = element_text(hjust = 0.5, size = 20))
```



4 Bar chart with color

To add color in the bar chart you can add a function fill in the ggplot() function.

fill = group: Assign to color the bar of the group bar.



5 Line chart

In this section, we will draw a line chart using BOD dataset. The dataset has two columns: Time, demand

BOD

	Time	demand
1	1	8.3
2	2	10.3
3	3	19.0
4	4	16.0
5	5	15.6
6	7	19.8

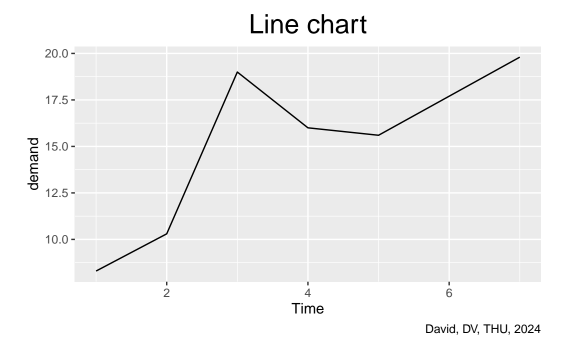
This dataset compares the data across two groups:

- Time: Represents the time in days at which the biochemical oxygen demand is measured.
- demand: Represents the biochemical oxygen demand (BOD) in milligrams per liter (mg/L).

geom_line():

- Used to add lines to a plot, typically to visualize trends or relationships between data points in sequential or continuous data.
- It connects data points in the order of their x-values.

```
ggplot(BOD, aes(x = Time, y = demand)) +
  geom_line() +
  labs(title = "Line chart",
      caption = "David, DV, THU, 2024") +
  theme(plot.title = element_text(hjust = 0.5, size = 20))
```



6 Histogram

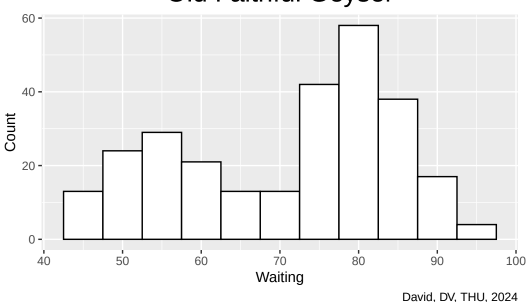
In this section, we will draw a histogram using faithful dataset. Our goal is to count waiting in the dataset.

geom_histogram(): To visualize the frequency distribution of a continuous variable.

- binwidth: To set the range of one bar.
- fill: To give a color to the bar.
- colour: To give color to the border of the bar.

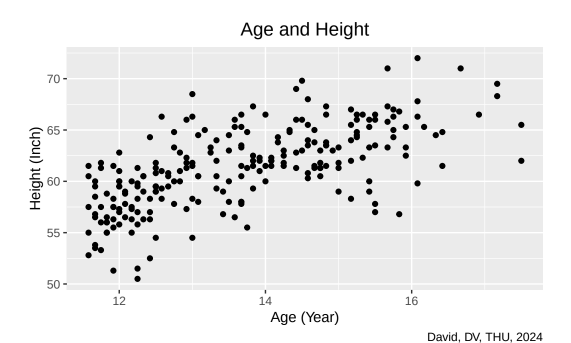
```
ggplot(faithful, aes(x = waiting)) +
labs(title = "Old Faithful Geyser",
    x = "Waiting",
    y = "Count",
    caption = "David, DV, THU, 2024") +
theme(plot.title = element_text(hjust = 0.5, size = 20)) +
geom_histogram(binwidth = 5, fill = "white", colour = "black")
```

Old Faithful Geyser



7 Correlation chart

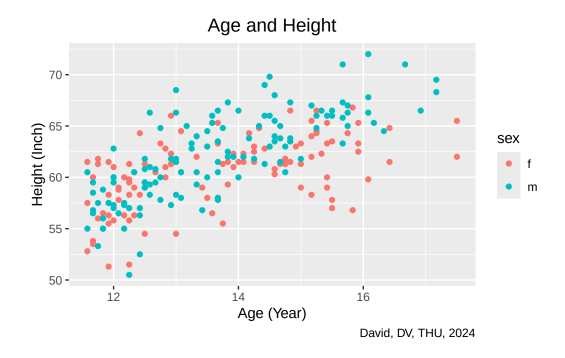
In this section, we will draw correlation chart using heightweight dataset. Our goal is to make a correlation between two variables. In this case, we correlate ageYear and heightIn by using geom_point.



8 Correlation chart: Color by group

To add color in the chart we can add color function to ggplot function.

We can add different color to determine different group by adding what group we want to differentiate. (color = sex)



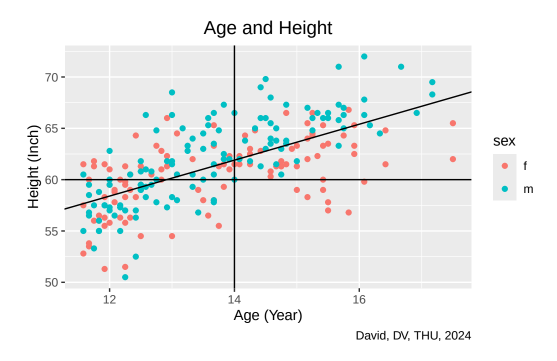
9 Correlation chart: Add lines

We can use these function to add lines in our graph:

- geom_vline(): To add a vertical line.
 - yintercept =: To determine coordinate where we want to add a horizontal line in y line.
- geom_hline(): To add a horizontal line.
 - xintercept =: To determine coordinate where we want to add a vertical line in x line.
- geom_abline(): To add a line according to our decision.
 - intercept: To determine coordinate where we want to add a line.
 - slope: To define the slop of the line.

```
ggplot(heightweight, aes(x = ageYear, y = heightIn, colour = sex)) +
  geom_point() +
  geom_hline(yintercept = 60) +
  geom_vline(xintercept = 14) +
  geom_abline(intercept = 37.4, slope = 1.75) +
```

```
labs(title = "Age and Height",
    x = 'Age (Year)',
    y = "Height (Inch)",
    caption = "David, DV, THU, 2024") +
theme(plot.title = element_text(hjust = 0.5, size = 14))
```



10 Correlation Chart: Change the text ticks of labels

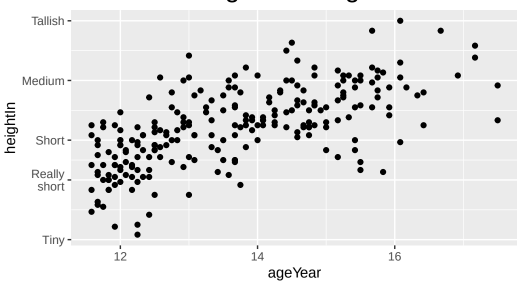
Now we want to change the text of tick labels by using heightwight dataset.

- breaks: Specifying the locations of the tick marks and labels on the y-axis (breaks = seq(min, max, by = interval)). This gives you more control over the spacing and readability of the axis labels.
- labels: Customizing the text of the axis labels (labels = function(x) ...). This allows for formatting, unit specification, or abbreviation of the labels.

```
ggplot(heightweight, aes(x = ageYear, y = heightIn)) +
  geom_point() +
  scale_y_continuous(
    breaks = c(50, 56, 60, 66, 72),
    labels = c("Tiny", "Really\nshort", "Short", "Medium", "Tallish")
```

```
) +
labs(title = "Age VS Height", caption = 'David, DV, THU, 2024') +
theme(plot.title = element_text(hjust = 0.5, size = 20))
```





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11 Multigroup histogram

To make a double histogram in a single chart, firstly we can add MASS to our library(), so we can use facet_grid() function.

```
library(MASS)
```

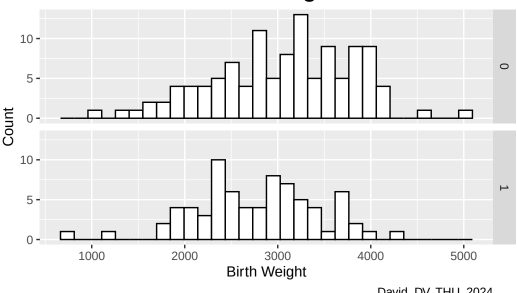
In this section, we will draw a histogram using birthwt dataset. Our goal is to count smoking and nonsmoking in the dataset.

facet_grid(): To determine whether the histogram wants to split horizontally or vertically.

- facet_grid(smoke ~ .): To split the histogram horizontal.
- facet_grid(. ~ smoke): To split the histogram vertical.

```
ggplot(birthwt, aes(x = bwt)) +
  labs(title = "Birth Weight",
      x = "Birth Weight",
     y = "Count",
      caption = "David, DV, THU, 2024") +
 theme(plot.title = element_text(hjust = 0.5, size = 20)) +
  geom_histogram(fill = "white", colour = "black") +
 facet_grid(smoke ~ .)
```

Birth Weight



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12 Multigroup Histogram: Change the name of histogram

To change the name of the histogram we can use recode_factor(), but first we must input tidyverse in our library().

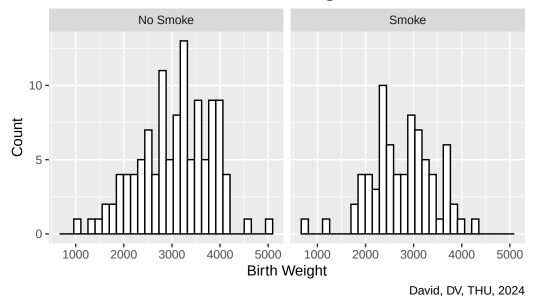
```
library(tidyverse)
```

We can create a new variable called birthwt_mod.

birthwt_mod\$smoke <- recode_factor(birthwt_mod\$smoke, '0' = 'No Smoke', '1' =</pre> 'Smoke'):

- We want to change the name in the smoke row.
 - By using \$ to mention the row name.
- Change 0 to No Smoke.
- change 1 to Smoke.

Birth Weight

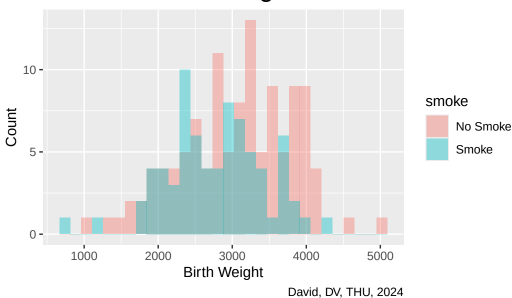


13 Multigroup Histogram: Add color

We can give color to the histogram by using fill in the ggplot function.

To make the both data become one graph we should add position = 'identity' and we can add aplha to adjust histogram density in the geom_histogram function.

Birth Weight



14 Multigroup Histogram: Add more than 2 histogram chart

In this section we will use functions from gridExtra library.

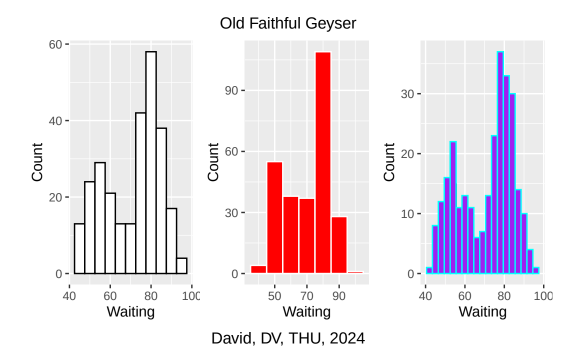
```
library(gridExtra)
```

To make a multigroup chart we need grid.arrange() function from the gridExtra package in R to arrange four plots (named plot1, plot2, plot3, and plot4) into a single figure.

We using faithful dataset.

grid.arrange(): To arrange a lot of charts into a single figure.

- ncol: To set how much charts in one row.
- top: To specifies the main title of a figure.
- bottom: To specifies the caption of a figure.



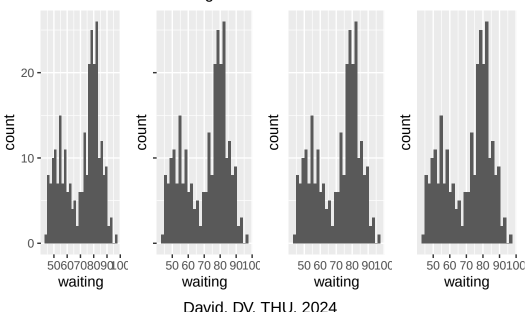
15 Multigroup Histogram: Remove tick marks

We can remove the tick marks in faithful dataset.

- To remove the tick marks, use theme(axis.ticks=element_blank()). This will remove the tick marks on both axes.
- To remove the tick marks, the labels, and the grid lines, set breaks to NULL

```
p1 \leftarrow ggplot(faithful, aes(x = waiting)) +
  geom_histogram() +
  theme(plot.title = element_text(hjust = 0.5, size = 12))
p2 \leftarrow ggplot(faithful, aes(x = waiting)) +
  geom_histogram() +
  theme(axis.text.y = element_blank())+
  theme(plot.title = element_text(hjust = 0.5, size = 12))
p3 \leftarrow ggplot(faithful, aes(x = waiting)) +
  geom_histogram() +
  theme(axis.ticks = element_blank(), axis.text.y = element_blank()) +
  theme(plot.title = element_text(hjust = 0.5, size = 12))
p4 \leftarrow ggplot(faithful, aes(x = waiting)) +
  geom_histogram() +
  scale_y_continuous(breaks = NULL) +
  theme(plot.title = element_text(hjust = 0.5, size = 12))
grid.arrange(p1, p2, p3, p4, ncol = 4,
             top = 'Removing Tick Marks and Labels',
             bottom = "David, DV, THU, 2024")
```

Removing Tick Marks and Labels



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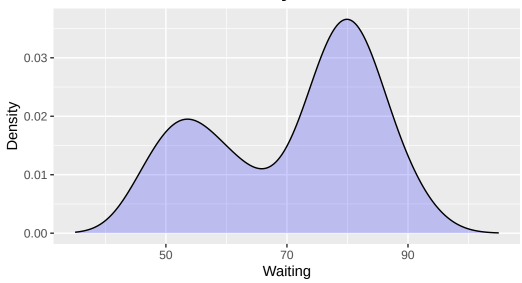
16 Density chart

In this section, we will make a density chart using faithful dataset.

We can use geom density to add a density chart - xlim(): To determine the x line range of density chart. - fill: To add a color to the density chart. - alpha: To determine color density in the chart

```
ggplot(faithful, aes(x = waiting)) +
 labs(title = "Density Chart",
      x = "Waiting",
      y = "Density",
      caption = "By David, Data Visualization Course, Tunghai University, 2024") +
 theme(plot.title = element_text(hjust = 0.5, size = 20)) +
 geom_density(fill = "blue", alpha = .2) +
 xlim(35, 105)
```

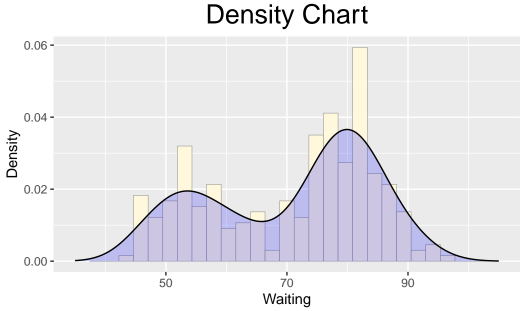




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17 Histogram and Density chart

To add a histogram chart in the density chart we can directly add <code>geom_histogram</code>. - fill: To add a color in the density or histogram chart. - <code>colour</code>: To change histogram border color. - <code>size</code>: To set a size of the histogram chart.

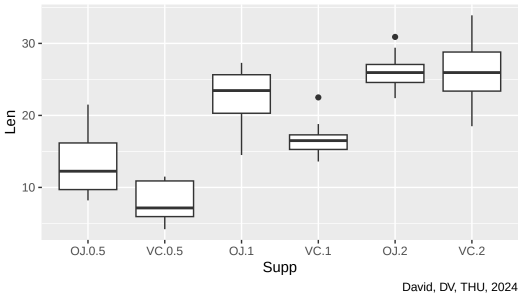


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18 Box plot

In this section, we will make a boxplot using geom_boxplot function in the ggplot. We will use ToothGrowth function.

Boxplot With Ggplot



19 Boxplot: Add labels and rotate labels

We use PlantGrowth dataset to try use these functions:

- Use scale_x_discrete() to change the text of the axis labels
- Use breaks = c() to break the axis labels
- Use labels = c() to add a name in axis labels
- axis.text.x = element_text(...): This part specifically targets the text elements of the x-axis. element text is a function that controls the formatting of text within the plot.
- angle = 30: This rotates the x-axis labels by 30 degrees. This is often useful when labels are long and overlapping.

```
pg_plot <- ggplot(PlantGrowth, aes(x = group, y = weight)) +
  geom_boxplot() +
  scale_x_discrete(
    breaks = c("ctrl", "trt1", "trt2"),
    labels = c("Control", "Treatment 1", "Treatment 2")
  theme(plot.title = element_text(hjust = 0.5, size = 16))
p1 <- pg_plot
```

Group VS Weight 6.0 -6.0 -6.0 -5.5 -5.5 **-**5.5 weight weight - 0.5 weight 4.5 -4.5 -4.0 -4.5 4.0 -3.5 -4.0 dronb Treatment 2 3.5 -Treatment 2 Treatment 1 3.5 -Contitoeatment 2 group group

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