READ ME

- How to use this guide
 - Currently the best way to search through this guide is to press CMD+F (if on Mac) or Ctrl+F (if on Windows/Linux) and to search for the following keywords
 - snippets, examples, python, plt, sns, statistics, terminology alert, sparsity, ETL, read_csv, get_dummies, probability, keras, tflearn, impute, onehotencod, dummy v, PCA, Hive, CNN, ensemble, validation, memory, usage, Hadoop, MAE, RSME, confusion, grid, function, matplotlib, plot, seaborn, facet, violin, histogram, bar, box, anomaly, count, label, title, libraries, sklearn., keras., tensor, from import, subplotting, regression, regressor, linear, logistic, SVM, clustering, classification, KNN, k-m, boosting, xgb, decision tree, random forest, bagging, deep, learning, convolution, dense, bias, variance, skew, heat, drop, probability, pandas, preprocessing, skew, correction, subsetting, dataframe, for i, apply, column, row, data engineering, concatenate, aggregate, terminology, important, feature, index, unique, value, naive bay, scaling, AB, A/B, predict, classifier, model, X train, X, y val, y test, error, deviation, accurate, metric, melt, loc, iloc, .column, NaN, missing, lambda, np.expm1, np.log1p, split, CART, gradient, Series, Index, list, array, convert, lightgbm, xgboost, catboost, reinforcement, rule, selection, gaussian, numba, timeit, dendogram, t test, z test, f test, matched pair test, anova, edge, bootstrap, ANOVA, F1 score, AWS, standard error, standard dev, degrees of freedom

STATISTICS

- Terminology Alert
 - X_bar is commonly used for *sample* mean, whereas μ is commonly used to represent the true population mean
 - Variance and Std Deviation I Why divide by n-1? by zedstatistics https://www.youtube.com/watch?v=wpY9o_OyxoQ (great video)
 - Note, to calculate variance if we have an entire population and no average
 - Population Variance
 - σ^2 = summation($(X \mu)^2$) / (N) # note population mean μ = summation(X)/n
 - Whereas, if we DO NOT have the population mean μ, we have to rely on using X_bar (a sample mean, typically given in problem statement) and use
 - Sample Variance
 - $s^2 = summation((X X_bar)^2)/(n 1)$

Probability

- Probability of rolling three dice without getting a six
 - To obtain the probability that at least one 6 is rolled in the three tosses
 - 1- (5/6)^3 Note, the probability of rolling a 6 with one dice on one roll is 1/6.
- Probability of rolling double sixes twice in a roll (rolling two dices at a time, two times)
 - Since each dice is being rolled independently
 - $(1/6)^1 * (1/6)^1 * 1/6)^1 * (1/6)^1 = (1/6)^4 = 0.00077$
- Probability of rolling one dice twice and getting a six both times
 - $(1/6)^1 * (1/6)^1 = (1/6)^2 = 1/36$
- https://sciencing.com/calculate-dice-probabilities-5858157.html

Statistics

- Statistics Fundamentals: Population Parameters by StatQuest (https://www.youtube.com/watch?
 - v=vikkiwjQqfU&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=4)
 - Since we rarely, if ever, have enough time to measure every single thing in a population distribution
 - for example, lets imagine that instead of having a population distribution created with 240 billion cells of Gene X, we only have a population distribution of 5 cells
 - We will use these 5 measurements to estimate the population parameters. The reason why we want the population parameters is to ensure results drawn from our experiment are reproducible.
 - In other words, if someone else measures the gene in 5 different liver cells, then they will get 5 different measurements. However, the new measurements will come from the same population.
 - Insights can then be derived from the population. One example insight might be, what is the probability of observing more than 30 mRNA transcripts in a single cell? We will apply this question to both of the two experiments and future experiments.
 - Note, on the screen, Josh is showing is showing a normal distribution with the number of mRNA transcripts from Gene X in 5 different liver cells
 - Blue line (Gene X) ranging from 0-40 mRNA with 5 green dots representing our samples
 - This green curve represents the population of our data (eg, one csv with data we are told to model)
 - Instead of describing the first 5 measurements we made, we want to actually *estimate* the population

parameters (eg, population mean, population std. deviation) and use them as the basis for the results

- Note, in the onscreen example, Josh is showing the *true* population mean is 20 and the *true* population standard deviation is 10.
 Again, these parameters are the *true* parameters of the dataset. Think of this as the data that we have been given to predict/model.
- We see from the first 5 measurements we did the *estimated* population mean is 17.6 and the *estimated* population std deviation is 10.1
 - Terminology alert: This is the training set
- Now when we repeat the experiment the estimated population mean is 19.2 and the estimated standard deviation is 12.7
 - Terminology alert: This is the testing set
- Thus, each time we do the experiment we get different estimates of the population parameters.
- And both sets of *estimates* are different
- Let's say we only had 2 measurements in the training set instead of 5.
 - The estimated population mean is 11 and the estimated population std deviation is 11.3
 - Note, the *true* population mean is 20 and the *true* population standard deviation is 10
- Let's say we only had 3 measurements in the training set instead of 5.
 - The estimated population mean is 15.3 and the estimated population std deviation is 11
- Let's say we only had 10 measurements in the training set instead of 5.
 - The *estimated* population mean is 19 and the *estimated* population std deviation is 10.5
- Therefore, this means that the more data we have, the more *confidence* we can have in the accuracy of the estimates.
- This is why it's important to choose the appropriate "test_size" for when we split our data for machine learning algorithms
- One of the main goals in statistics is quantifying how much confidence we can have in population estimates.

- Specifically, statisticians often calculate pvalues and confidence intervals to quantify the confidence in the estimated parameters.
- And like we just saw, generally speaking, the more data, the more confidence we have in the estimates
- Going back to the two replicate experiments (training data vs testing data). Even though these two experiments resulted in different estimates for the population mean and population standard deviation, we can use statistics to quantify our confidence in how different they are.
 - In this case, a p-value, or, alternatively, a confidence interval, would tell us that while the estimates are different, they are not significantly different.
 - That means, the results from the first experiment (training) should not be significantly different from the results generated from the second experiment (testing).
 - <u>In conclusion, we can generate results</u> that are reproducible in future experiments.
- StatQuest: Confidence Intervals (https://www.youtube.com/watch? v=TqOeMYtOc1w&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=19)
 - Many people misunderstand confidence intervals, but that's only because they didn't learn about bootstrapping first
 - Here is a bootstrap refresher
 - Imagine if we weighed a bunch of female mice
 - In the video, Josh is currently showing a 1D line ranging from 15-35. On the line, are 12 sample of mice who have been weighed. Note, we didn't weigh every single female mouse on the planet, just 12.
 - We can calculate the mean of the 12 mice.
 Note, the sampled mean is not the mean of all the mice on the entire planet.
 - This is where bootstrapping comes in. We can use it to get a better estimate of the global mean of all the female mice on the entire planet.
 - Since we already have the mean from the first sample we collected we can implement bootstrapping.

- Step one —> We randomly select 12 weights from the original sample (global mice population, duplicates are ok)
 - First. In the video, Josh is now showing another 1D line with 12 new randomly chosen mouse samples directly underneath the first 1D line. However, in this sample there are a few duplicate mice weights, requiring us to stack the markers on top of one another.
 - Note, there is something called sampling width replacement that can happen here. In the second 1D line we se that there are *two* furthest left samples fall directly beneath the *one* furthest left sample on the first line. Also, to the right of the *one* furthest left sample is another sample a few weights away. We then notice that there isn't a weight down at this value on the second 1D. There is a gap. We call thing sampling width replacement.
- Step two —> We calculate the mean of the random sample (the second 1D line)
- Repeat steps one and 2 until we have calculated a lot of means (> 10,000)
 - Note, in the video Josh is now displaying a new 1D line with all of the means plotted on the number line. There is a heavily dense region right in the middle of the line around the middle, ~25. Appears to be normally distributed.
 - Thats all there is to bootstrapping
- Lets look at what a confidence interval is now (typically 95% confidence intervals are used)
 - A 95% confidence interval is just an

- interval that covers 95% of the means.
- For example, 95% of the bootstrapped means fall in between 21 and 31.
- That's all a confidence interval is!
- Now that we know what a confidence interval is, let's look at why they are useful
 - Confidence intervals are useful because they are statistical tests that can be performed visually
 - For our example, because the interval covers 95% of the means, we know that anything outside of it occurs less than 5% of the time.
 - That is to say, the p-value of anything outside of the confidence interval < 0.05 (and thus, significantly different)
 - Here is an example of a visual statistic test
 - What is the p-value that the "true" mean of all female mice, not just in our sample, weights are < 20?
 - Looking at our bootstrapped means, we see the highlighted region is outside of the 95% confidence interval which contains 95% of the means, we know that the probability that the "true" mean is in this area has to be < 0.05.
 - The p-value is < 0.05. This is unlikely and we say there is a statistically significant difference between this sample relative to the other samples on the bootstrapped line/plot.
 - Here is another example (comparing two samples)
 - Female Mice vs Male Mice
 - In the video, Josh is still showing the female mice bootstrapped means on a 1D plot. Again, the bootstrapped means

- appear to be normally distributed around a mean of 25.
- Directly underneath the female plot, Josh is showing a new plot with bootstrapped means of male mice. However, here the mice are normally distributed but the mean on the male mice plot falls on the upper-end value on the female mice plot
- In other words, the righttail of the female mice distribution crosses over the left tale of the male mice distribution.
 - IMPORTANT: Only the tails overlap here. The 95% confidence intervals on both plots do not overlap. Therefore, we know there is a statistically significant difference in the weights of female and male mice. We know the p-vale is < 0.05 just by looking at this picture!
 - HOWEVER:
 There is one caveat. What if the confidence intervals overlap a little (some p-value > 0.05)? If they overlap, there is still a chance that the means

are significantly different from each other, so, in this case, we still have to do our t-test! When they don't overlap, we don't have to and can be rest assured that is a statistically significant difference between the two means.

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ANOVA and related

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ANOVA and related

- Hypothesis Testing (Critical Value Approach)
 - https://newonlinecourses.science.psu.edu/statprogram/ reviews/statistical-concepts/hypothesis-testing/critical-valueapproach(great read)
- https://keydifferences.com/difference-between-t-test-and-f-test.html#targetText=The%20difference%20between%20t%2Dtest%20and%20f%2Dtest%20can%20be,is%20small%20is%20t%2Dte st.&targetText=In%20contrast%2C%20f%2Dtest%20is,to%20comp are%20two%20population%20variances and <a href="https://brandalyzer.blog/2010/12/05/difference-between-z-test-f-test-and-t-test/#targetText=A%20z%2Dtest%20is%20used,population%20standard%20deviation%20or%20not.&targetText=An%20F%2Dtest%20is%20used%20to%20compare%202%20populations'%20variances. (great read)
 - Note, the following methods are generally used to validate our data. Just because we have a p-value lower than our acceptance criteria doesnt mean there is conclusive evidence that something significant is going on.
 - T test is usually performed after a p-value results in a number around our acceptance criteria.
 - For example, a p-value of 0.07 when our acceptance criteria is 0.05 (see StatQuest:

Confidence Intervals by Josh Starmer (~4:00 https://www.youtube.com/watch? v=5Z9OIYA8He8&list=PLblh5JKOoLUK0FLuz wntyYI10UQFUhsY9&index=20) and StatQuickie: Thresholds for Significance by Josh Starmer (FANTASTIC short video https://www.youtube.com/watch? v=KEofcJ1tfkl&list=PLblh5JKOoLUK0FLuzwnt yYI10UQFUhsY9&index=21)

- Also, note that just because we calculate small p-value (eg, 0.03) under our acceptance criteria (eg, 0.05), it doesn't mean that we have explained our data. We also generally want to have a good r-squared and use additional methods (eg, t-test, z-rest, etc) with our model that correlates and validates the data, respectively. We want to be able to explain the data.
- Another big note....extraordinary claims need extraordinary data! We don't want a p-value of 0.047 with an acceptance criteria of 0.05. If we were to go publish an extraordinary claim our p-value better be crazy small.
- T test is used to compare two related samples (good for small # of observations < ~30)
 - StatQuickie: Which t test to use (https:// www.youtube.com/watch?v=nnBJeb_Iq8&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9& index=22)
 - There are two main type:
 - Paired Data
 - Are useful if we have "before and after" measurement taken from the same test subject
 - For ex, if we have a group of people who are fixing to go in a clinical trial for blood pressure, we can measure their blood pressure before taking a certain blood pressure medication and then measure their blood pressure after they have taken the medication.
 - A pair of "before and after" measurements for a test subject

- Unpaired Data
 - Occurs when we have two different samples from a population.
 - For ex, if we measured height for one group of people (Group A), and measured the heights of another group of people (Group B)
 - There are two subcategories of unpaired t-tests
 - One assumes the variance of height measurements in Group A is equal to the variance of height measurements in Group B
 - The other one assumes the opposite, the variances differ (Josh Starmer recommends going with this one as it is a more conservative approach. Therefore, if the data can pass this more conservative t-test, it will mean the data is rock solid. This is my general recommendation.)
- Should you use a one-tail / one-sided t-test or a two-tail / two-sided t-test?
 - A two-sided t-test
 - Let's go back to our example with Group A and Group B height measurements. A two-tail t-test for instance will test to see if Group A is significantly higher than Group B or if Group A is

significantly lower than Group B. It is agnostic to the t-test, it doesn't know if Group A should be higher or lower than Group B (same for vice-versa.

- A one-sided t-test
 - Is a lot less conservative. It requires the user (us) to know ahead of time which one is higher.
- IMPORTANT: Generally, especially in academic journals, we always want the data to speak for itself. Therefore, the two-sided t-test is the better test to go with since it is slightly more conservative and lets the data speak for itself.
- T-test is a univariate hypothesis test, that is applied when standard deviation is not known and the sample size is small. T-statistic follows Student t-distribution, under null hypothesis.
 - https://www.youtube.com/watch?
 v=pTmLQvMM-1M (great video on Student's t-test)
 - t value = signal / noise = diff.
 Between group means / variability of groups
- A t-test is used for testing the mean of one population against a standard or comparing the means of two populations if you do not know the populations' standard deviation and when you have a limited sample (n < 30). If you know the populations' standard deviation, you may use a z-test.
 - Examples
 - Measuring the average diameter of shafts from a certain machine when you have a small sample.
- Z test (good for larger # of observations)
 - A z-test is used for testing the mean of a population versus a standard, or comparing the means of two populations, with large (n ≥ 30) samples whether you know the population standard deviation or not. It is also used for testing the proportion of some characteristic versus a standard proportion, or comparing the proportions of two populations.
 - Examples
 - Comparing the average engineering salaries of men versus women.

- Comparing the fraction defectives from 2 production lines.
- F test is used to test the equality of two populations
 - F-test is statistical test, that determines the equality of the variances of the two normal populations. Fstatistic follows Snedecor f-distribution, under null hypothesis. Comparing two population variances.
 - Examples
 - Comparing the variability of bolt diameters from two machines.

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- Matched pair test
 - Matched pair test is used to compare the means before and after something is done to the samples. A t-test is often used because the samples are often small. However, a z-test is used when the samples are large. The variable is the difference between the before and after measurements.
 - Examples
 - The average weight of subjects before and after following a diet for 6 weeks

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- F Statistic/Critical and F Value
 - https://www.statisticshowto.datasciencecentral.com/probabilityand-statistics/f-statistic-value-test/#ANOVA (great read)
 - "An F statistic is a value you get when you run an ANOVA test or a regression analysis to find out if the means between two populations are significantly different. It's similar to a T statistic from a T-Test; A-T test will tell you if a single variable is statistically significant and an F test will tell you if a group of variables are jointly significant."

F Distribution

- https://www.statisticshowto.datasciencecentral.com/ probability-and-statistics/f-statistic-value-test/#ANOVA(great read, see graph in F distribution section)
 - "The F Distribution is a probability distribution of the F Statistic. In other words, it's a distribution of all possible values of the f statistic."
 - "The F distribution is related to chi-square, because the f distribution is the ratio of two chi-square distributions with degrees of freedom v1 and v2 (note: each chi-square is first been divided by its degrees of freedom). Each curve depends on the degrees of freedom in the numerator (dfn) and the denominator

(dfd). These depend upon your sample characteristics."

- "For example, in a simple one-way ANOVA between-groups,"
 - Dfn = a 1
 - dfd = N a
- where
 - a = the number of groups
 - n = the total number of subjects in the experiment
- Terminology alert: The degrees of freedom in the denominator (dfd) is also referred to as the degrees of freedom error (dfe).
- Chi-squared
 - https://www.statisticshowto.datasciencecentral.com/ probability-and-statistics/chi-square/ (great read)
 - What is a Chi-squared Test?
 - Two types (both use "chi-square statistic" and "chi-square distribution" for different purposes)
 - Chi-square goodness of fit test
 - Determines if a sample data matches a population (see Goodness of Fit Test for more info https:// www.statisticshowto.datascience central.com/goodness-of-fit-test/)
 - Chi-square test for independence
 - Compares two variables in a contingency table to see if they are related. In a more general sense, it tests to see whether distributions of categorical variables differ from each other.
 - A very small chi square test statistic means that our observed data fits our expected data extremely well. In other words, there is a relationship.
 - A very large chi square test statistic means that the data does not fit very well. In other words, there isn't a relationship.
 - What is a chi-square statistic?
 - $(chi_c)^2 = summation((O_i E_i)^2 / E_i)$

- where the subscript c are the degrees of freedom, O is our observed value (eg, markers on a scatter plot), and E is our expected value from our model, and i for every "ith" position
 - Note, it's very rare that we'll ever want to actually use this formula to find a chi-square by hand.
- A low value chi-square means there is a high correlation between our two sets of data. In theory, if our observed and expected values were equal ("no difference") then chi-square would be zero
 - Note, deciding whether a chisquare test statistic is large enough to indicate a statistically significant difference isn't as easy it seems.
- One Way ANOVA (https://www.youtube.com/watch?v=q48uKU_KWas)
 - The ultimate goal of the one-way ANOVA is to compare the means at at least three conditions
 - The first step is to always state the null hypotheses and the alpha level, all of this is done apriori
 - The null hypotheses states there are no significant relationships/correlations in the data. In other words, they all have same same mean.
 - Example
 - 123 (column id's)
 - 122
 - 242
 - 524

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- Step 1
 - Null hypothesis (all means are equal, nothing significant in the data): $\mu_1 = \mu_2 = \mu_3$
 - Alternative hypothesis: We are going to hypothesize that there is at least one difference among the means.
 - Alpha level = 0.05 # commonly used
- Step 2
 - Determine degrees of freedom between groups (df_between) by taking the conditions/features in our study (k) and

subtracting one

- df_between = k 1
- df between = 3 1 = 2
- Terminology alert: "Degrees of freedom between" is also referred to as the "degrees of freedom in the numerator"
- Note, the between part means we have different subjects/cases/ features in each group
- Determine degrees of freedom within (df_within) by taking the total amount of values we have (N) and subtracting the number of conditions/features (k)
 - df within = N k
 - df within = 9 3 = 6
 - Terminology alert: "Degrees of freedom within" is also referred to as the "degrees of freedom in the denominator"
- Determine total degrees of freedom by adding df_between and df_within
 - df total = 8
- Determine F_critical (aka F_stat)
 - Use F distribution table or calculator to calculate F_critical
 - $F_{critical} = 5.14$
 - Note.
 - https://
 www.statisticshowto.datascie
 ncecentral.com/probabilityand-statistics/f-statisticvalue-test/#ANOVA
 - "A calculator will certainly give you a fast answer. But with many scenarios in statistics, you will look at a range of possibilities and a table is much better for visualizing a large number of probabilities at the same time."

- Analysis of Variance
 - Calculate mean for each condition

-
$$\mu_1 = (1+2+5)/3 = 2.67$$

$$\mu_2 = (2+4+2)/3 = 2.67$$

-
$$\mu_3 = (2+2+4)/3 = 3.00$$

- Calculate grand mean of data
 - μ _total = summation(3x3 matrix)/count(3x3 matrix) = 25/9 = 2.78

-
$$\mu_{\text{total}} =$$
 (2.67+2.67+3.00)/3 = 25/9 = 2.78

 Calculate sum of squares total (SS_total)

$$= (1 - 2.78)^2 + (2 - 2.78)^2 + (5 - 2.78)^2 + etc$$

 Calculate sum of squares within (SS_within)

$$(2 - 3.00)^2 + (4 - 3.00)^2$$

- Step 4
 - Calculate the mean square between / aka the variance between
 - μ_square_between = SS_between / df_between = 0.23 / 2
 - μ_square_between = 0.12
 - Calculate the mean square within / aka the variance within
 - μ_square_within = SS_within /

 $df_within = 13.34 / 6$

μ_square_within = 2.22

- Step 5
 - Calculate F value
 - F_value = μ_square_between / μ_square_within = 0.12 / 2.22
 - F value = 0.05
 - Now, let's compare F_value with F_critical
 - Here, the F_value (0.05) is smaller than F_critical (5.14)
 - So this supports the null hypothesis, meaning there is no significant difference between the three groups.
 - However, if the data set proved to be significant (aka we rejected the null hypothesis), we should also consider the p value. The p value is determined by the F statistic and is the probability the results could have happened by chance.
 - In other words, we would need to look into using the F critical/statistic in COMBINATION (not comparing) with the p-value to determine if ALL of the variables in the dataset are significant. The F critical/statistic is just comparing the joint effect of all the variables together.

- What is a p value?
 - https://www.statisticshowto.datasciencecentral.com/p-value/ (great read)
 - Note, in the examples below we are calculating a two-sided p-value since we are considering equally or rarer possibilities (part 2 and part 3 below).
 - Example 1 (Simple)
 - StatQuest: P Values, clearly explained
 - https://www.youtube.com/watch?v=5Z9OIYA8He8 (see ~05:20)
 - What is the p-value of getting two heads in a row

when flipping a coin?

- HH, HT, TH, TT
- Part 1
 - The first part of a p-value is the probability that random chance generated the data
 - The probability of flipping a coin two times and getting heads on each toss $-> (1/2)^1 * (1/2)^1 =$ $(1/2)^2 = 0.25$
- Part 2
 - The second part of a p-value is to add anything else in the outcome that has equal probability.
 - The probability of getting two tails on both coin flips is the same as getting two heads $-> (1/2)^1$ * $(1/2)^1 = (1/2)^2 = 0.25$
- Part 3
 - The last part of a p-value is to add on anything that is rarer than what was observed
 - This part would be equal to zero since there aren't any rarer outcomes than two heads or two tails —> 0
- Total p-value
 - herefore, the p-value for HH is 0.25+0.25+0 = 0.50
- To summarize this example
 - The probability of getting HH is 0.25
 - The p-value for getting HH is 0.5
 - In this case, the probability and p-value are not equal
- Example 2 (Simple, ~same as Example 1)
 - https://www.youtube.com/watch?v=5Z9OIYA8He8 (see ~07:00)
 - What is the p-value of getting five heads in a row when flipping a coin five times?
 - see video for all possible combinations and better visualization
 - Part 1
 - The first part of a p-value is the probability that random chance generated the data
 - $(1/2)^5 = 1/32 = 0.03125$
 - Part 2

- The second part of a p-value is to add anything else in the outcome that has equal probability.
 - $(1/2)^5 = 1/32 = 0.03125$
- Part 3
 - The last part of a p-value is to add on anything that is rarer than what was observed
 - This part would be equal to zero since there aren't any rarer outcomes than five heads or five tails —> 0
- Total p-value,
 - Therefore, the p-value for HHHHH is 0.03125+0.03125+0 = **0.0625**
- To summarize this example
 - The probability of getting HHHHH is 0.03125
 - The p-value for getting HHHHH is 0.0625
 - In this case, the probability and p-value are not equal
- Example 3 (intermediate)
 - https://www.youtube.com/watch?v=5Z9OIYA8He8 (see ~09:45)
 - What is the p-value of getting four heads and one tail when flipping a coin five times? Order doesn't matter.
 - see video for all possible combinations and better visualization
 - Part 1
 - The first part of a p-value is the probability that random chance generated the data
 - since there are 5 possibilities of getting four heads
 - out of 32 total possible outcomes
 - 5/32 = 0.15625
 - Part 2
 - The second part of a p-value is to add anything else in the outcome that has equal probability.
 - since there are 5 possibilities of getting four tails
 - out of 32 total possible outcomes
 - -5/32 = 0.15625
 - Part 3

- The last part of a p-value is to add on anything that is rarer than what was observed
 - Here, there are two outcomes that are more rare than a combination of four heads and one tail. They are —> HHHHH and TTTTT
 - We then add the probability of both rarer outcomes
 - -1/32 + 1/32 = 0.0625
- Total p-value
 - Therefore, the p-value for a combination of four heads and one tail is is 0.15625 + 0.15625 + 0.0625 = **0.375**
- Example 4 (advanced)
 - https://www.youtube.com/watch?v=5Z9OIYA8He8 (see ~11:00)
 - What about if we were measuring height instead? It was easy to write out all possible outcomes for coin tosses, but what if we wanted to calculate the probabilities for heights instead? We use the "density" function instead.
 - Let's look at the distribution/density of height measurements in women between 15 and 49 years old in 1996
 - The area under the curve indicates the probability that a person will have a height within a range of possible values.
 - In this example, 95% of the area under the curve is between 142 cm and 169 cm, indicating that most women are between those two values.
 - In other words, there is a 95% probability that each time we measure a woman's height, their height will be between 142 cm and 169 cm.
 - Also, there is a 2.5% probability that each time we measure a woman's height, their height be less than 142 cm.
 - There is also a 2.5% probability that each time we measure a woman's height, their height be greater than 169 cm.
 - What is the p-value for someone who is 142 cm tall? (See ~12:00)
 - Part 1
 - The first part of a p-value is the probability that random chance

- generated the data
- The area for people 142 cm or shorter
 -> 2.5%
- Part 2
 - The second part of a p-value is to add anything else in the outcome that has equal probability.
 - The area for people 169 cm or taller -> 2.5%
- Part 3
 - The last part of a p-value is to add on anything that is rarer than what was observed
 - Here nothing is rarer -> 0%
- Total p-value
 - 5.0%
- What's the p-value for someone who is between 155.4 and 156 cm tall between someone is 142 cm tall and 169 cm tall (at the middle of the distribution)? (See ~13:48)
 - Note: The probability of someone being between 155.4 and 156 cm is only 0.04 or 4%.
 The area is pretty small!
 - Part 1
 - Note: The first part of calculating a pvalue is the probability of the event of interest
 - Therefore -> 4.0 %
 - Part 2
 - The second part of a p-value is to add anything else in the outcome that has equal probability.
 - We see the left side of the distribution from 142 cm to 155.4 cm is 48%
 - We see the right side of the distribution from 156 cm and 169 cm is also 48%
 - Therefore, 48% + 48% = 96%
 - Part 3
 - The last part of a p-value is to add on anything that is rarer than what was observed
 - Here nothing is rarer -> 0%
 - Total p-value
 - 100% or 1
 - In conclusion, this means that there is nothing special about measuring someone who has the average height

- even though that in particular is rare
 Also, in this example, the probability of
 measuring someone between 155.4 and
 156 cm tall is tiny (0.04 or 4.0%), but
 the p-value is huge (1 or 100%)
- In other words, this is also conclusive that there is nothing significant about measuring someones who has an average height.
- StatQuest: One or Two Tailed P-Values by Josh Starmer (https://www.youtube.com/watch?v=bsZGt-caXO4&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=23) great
- The Binomial Distribution and Test, Clearly Explained!!! by Josh Starmer (https://www.youtube.com/watch?v=J8jNoF-K8E8&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=24) fantastic video to see how we can relate this to real business problems
 - Usually when people talk about the binomial distribution, they talk about flipping a coin
 - But who really cares about flipping coins?
 - What folks really want to know is whether or not people like
 Orange Fanta more than Grape Fanta
 - Which flavor reigns supreme? Or are they both equally loved (null hypothesis)??
 - Scenario
 - Let's say we take a sample of 7 people. From the sample, 4 people like Orange Fanta and 3 people like Purple Fanta. Is this one sample enough to be confident that "most people like Orange Fanta"? Or could it be that people, in general, don't have a preference and these results are just due to random chance (aka null hypothesis) and a small sample size?
 - Let's say we take another survey/sample of 7 people, and 3 people say they like Orange Fanta and 4 people say they like Purple Fanta.
 - To get to the bottom of this mystery, we need to get a sense of what to expect if there is no preference. Then we determine if our survey results fit those expectations. If not, then we can reject the idea that both Fanta are loved equally (aka we reject the null hypothesis)
 - The binomial distribution will tell us what to expect if there is no preference.
 - Note, the binomial distribution model follows the following formula of what to expect when

there is no preference

- $pr(x \mid n, p) = ((n!)/((x!)*(n-x)!)) * (p^x) * (1-p)^(n-x)$
- If the model is a poor fit, we will reject the idea that both flavors are loved equally (aka we reject the null hypothesis)
- Example
 - Assume I asked 3 people if they liked
 Orange Fanta more than Grape Fanta
 - The first two say they like Orange Fanta and the last one says Grape Fanta.
 - Note, there is a 50% chance a person will say either orange or grape
 - Now, we can calculate the probability of the first two people randomly choosing orange and the third person randomly choosing grape
 - Assuming there is no preference between the two
 - The probability of the first person selecting orange is
 - 0.5
 - The probability of the first two people selecting orange is
 - 0.5 * 0.5 =**0.25**
 - The probability of the first two people preferring orange and the third person preferring grape is
 - 0.5 * 0.5 * 0.5 =**0.125**
 - NOTE: 0.125 is the probability of the first two people saying they prefer orange and the third person saying they prefer grape.
 - IT IS NOT the probability that any 2 out of 3 people would prefer orange.
 - It could have just as easily been that the first person said they preferred grape.
 - Note, all of the following combinations are equally as likely
 - purple, orange, orange = 0.125
 - orange, purple, orange = 0.125
 - orange, orange, purple = 0.125
 - To get the probability that any 2 out of 3 people prefer Orange Fanta we have to sum up the probability of all possible

combinations:

- 0.125 + 0.125 + 0.125 = 0.375
- Therefore, the probability of randomly selecting three people and two of them saying they prefer Orange Fanta and one saying Purple Fanta is 0.375
- Alternatively, we could of done the math using this formula:
 - $pr(x \mid n, p) = ((n!)/((x!)*(n-x)!)) * (p^x) * (1-p)^(n-x)$
 - x = number of people who preferred Orange Fanta =2
 - n = total number of people we asked = 3
 - Note, "x I n" also means "n

 x". Therefore, it also means the total number of people minus the number of people who preferred Orange Fanta equals the number of people who said they prefer Grape Fanta
 - p = probability that someone will pick Orange Fanta = 0.5
 - Note, the probability that someone might pick Grape Fanta is "1 - p"
 - Together, the expression says "The probability of x (the number of people who say they prefer Orange Fanta), given n, the number of people we asked, and p (the probability of picking Orange Fanta)..."
 - $pr(x \mid n, p) = 0.375$
- Lets go back to our original question. If 4 people say they like Orange Fanta and 3 people say they like Grape Fane, can we conclude that people in general prefer Orange Fanta?
 - $pr(x \mid n, p) = ((n!)/((x!)*(n-x)!)) * (p^x) * (1-p)^(n-x)$
 - where x = 4, n = 7, and p = 0.5
 - pr(x | n, p) = 0.273

- In other words, 0.273 is the probability that 4 of 7 people would randomly prefer Orange Fanta
- Note, when you use a binomial distribution to calculate a p-value, it's called a Binomial Test
 - So what's the p-value for 4 of 7 people preferring Orange Fanta?
 - Note, the p-value is the probability of the observed data (4 of 7 people prefer Orange Fanta), plus the probabilities of all other possibilities that are equally likely and rarer (see ~11:30 in the video, https://www.youtube.com/watch?v=J8jNoF-K8E8&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=24)
 - The conclusion for the p-value saying that 4 out of 7 people stating that they prefer Orange Fanta is 1
 - Therefore, this means that we support the null-hypothesis. Both flavors are equally loved. For there to be strong evidence in saying one is preferred over the other, the p-value would need to be lower than some acceptance criteria (eg, α = 0.05)
- StatQuickie: Standard Deviation vs Standard Error by Josh Starmer (https://www.youtube.com/watch? v=A82brFpdr9g&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=13)
 - The standard deviation of the means is called the standard error.
 - For example, imagine we took 5 different samples where there are 5 measurements/observations per sample. The sample mean and sample standard deviation can be calculated for each sample. If we were then to take the 5 means, and plot them on a number line and calculate the standard deviation. We would get the standard error.
- Alpha Value
 - https://www.statisticshowto.datasciencecentral.com/p-value/ (see P Value vs Alpha level section)
 - "Alpha levels are controlled by the researcher and are related to confidence levels. You get an alpha level by subtracting your confidence level from 100%. For example, if you want to be 98 percent confident in your research, the

alpha level would be 2% (100% - 98%). When you run the hypothesis test, the test will give you a value for p. Compare that value to your chosen alpha level. For example, let's say you chose an alpha level of 5% (0.05). If the results from the test give you:"

- "A small p (≤ 0.05), reject the null hypothesis. This is strong evidence that the null hypothesis is invalid."
- "A large p (> 0.05) means the alternate hypothesis is weak, so you do not reject the null."
- Multicollinearity (occurs when independent variables in a regression model are correlated.)
 - <u>https://towardsdatascience.com/multicollinearity-in-data-science-c5f6c0fe6edf</u>
- Bias and Variance Tradeoff (Great)
 - https://www.youtube.com/watch?v=EuBBz3bI-aA
 - The first thing we do before start looking bias and variance is to split out data into a training and test set
 - Example
 - imagine viewing a scatter plot with data in a log10 shape and showing 80% of the markers/ data as blue and the remaining 20% as green. The blue dots are the training set and the green dots are the testing set

The inability for a machine learning method (like linear regression) to capture the true relationship is called bias

- Example

 Think about trying to fit a linear line to a training set with log10 type scatter data

- Because the linear line can't be curved like the "true" relationship, we say it has a relatively large amount of bias. In other words, the inability for a machine learning method (like linear regression) to capture the true relationship is called bias
- Example
 - Another machine learning method, may try to fit a squiggly line (zig-zag type shape) to the training set. The squiggly line can handle to the arc of a true relationship with log10 type data so we can it would have a relatively low amount of bias compared to trying trying to fit a straight line to log10 type data.
 - To quantify this, we can compare the sum of squares between the squiggly line model and straight line model. But

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note, the squiggly line model fits the scatter plot so well, that the sum of squares will be zero! The squiggly line wins here at modeling the training set data.

- Lets now look at the testing set and calculate the sum of squares
- The straight line wins here! It's sum of squares is lower than that of the squiggly line fine

Variance

- Example
 - Since the squiggly line model fitted the training set so well, but failed to fit the testing set well, we say it has a relatively large amount of variance. In other words, the difference in fits between the data sets is called variance.
 - In summary, the squiggly line had low bias with the training and test data since it is flexible and can adapt to the log10 shaped data. But...the squiggly line has high variability/variance because it results in a vastly different sums of squares for different data sets. In other words, it's hard to predict how well the squiggly line will perform with future data sets. It might do well sometimes, and other times it might do terribly.
 - In contrast, the straight line has relatively high balance when trying to capture the true relationship of the log10 shaped data. But.... The straight line has relatively low variability/ variance because the sums of squares are very similar between different datasets. In other words, the straight line might only give good predictions, and not *great* predictions. But they will be consistently good predictions
 - Terminology alert:
 - If the model fits the training set better than the test set we call this overfitting.

Overall

- Example
 - In machine learning, the ideal algorithm has low bias and can accurately model the true relationship (imagine a model fitting a training set that has log10 type shape data). Also, the ideal model will have low variability, by producing consistently predictions across

different data sets (imagine a representation of the same model but now with testing data. The model will ideally fit it the same. The sums of squares are about equal between data sets if it has low variability.) An ideal model for your data to create is done by finding the sweet spot between a simple model (think about the straight line fitting the training set, ~high bias) and complex model (think about the squiggly line model fitting the training set perfectly, zero bias)

- Terminology alert:
 - Three commonly used methods for finding the sweet spot between simple and complicated models are:

 regularization, boosting, and bagging

- Finance
 - Black-Scholes model
 - Definitions:
 - Risk
 - How do you measure Risk
 - Stocks
 - Portfolio
- Difference between linear and logistic regression
 - https://techdifferences.com/difference-between-linear-and-logistic-regression.html

SLANG TERMS/TERMINOLOGY

- Long Data
 - Bunch of columns, little rows
- Wide Data
 - Bunch of rows, little columns

MATHEMATICS

- Common Functions
 - Sigmoid

- Threshold
- Rectifier
- Hyperbolic tangent
- Softmax Function
- Cross-Entropy
- Common Distributions
 - Gaussian
- The Applications of Matrices
 - https://www.youtube.com/watch?v=rowWM-MijXU

LOOK UP / SORT LATER

- Python
 - Error Handing!
 - OpenCV
 - Looping through files in a directory
 - sparsity (can come up a lot in NLP, we want to avoid it)
 - https://realpython.com/python-modules-packages/
- Watch Later
 - YouTube
 - Image Classifier using VGG16 Model (Great playlist to go through by deeplizard)
 - https://www.youtube.com/watch?
 v=oDHpqu52sol&list=PLZbbT5o_s2xrwRnXk_yCPtnqqo4_u 2YGL&index=13
- Big Data Tools
 - Hadoop
 - Spark
 - H2O
 - CDSW
 - Domino Labs
 - HIVE
 - PIG
 - and unstructured data
 - etc.
 - CPLEX, IBM-DOC & AnyLogic
 - Developing and/or applying linear, mixed integer, stochastic programming to solve demand planning, supply chain, production scheduling
 - Discrete Event Simulation, Factor Analysis, Genetic Algorithms, Bayesian Probability Models, Hidden Markov Models and Sensitivity Analysis. (Cotiviti)
 - Amazon AWS machine learning technologies such as Amazon SageMaker, TensorFlow, PyTorch, Keras, Amazon Transcribe, Amazon Textract
- Matrices

- scipy.sparse matrices vs numpy matrices?
- Feature Selection
 - Heatmaps
- Leaf-wise vs Level-wise Decision Tree Algorithms
 - Leaf-Wise
 - lightgbm
- Omnichannel
- How to Implement a Real World Usecase or Project for Data Science
 - https://www.youtube.com/watch?v=Ur6tpb0elkY
 - Do Kaggle Competitions
 - Also learn Django and Flask to actually be able to implement models into web applications
 - Learn deployment techniques
 - Pipelines

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- XGBoost, Lightgbm, and CatBoost
 - https://www.youtube.com/watch?v=V5158Oug4W8
 - https://towardsdatascience.com/catboost-vs-light-gbm-vs-xgboost-5f93620723db
 - these are are gradient boosting algorithms for decision trees
 - xgboost
 - https://xgboost.readthedocs.io/en/latest/tutorials/index.html
 - loss function term
 - regularization term
 - xgboost has show to work very welling practice
 - LightGBM
 - written by Microsoft
 - much faster than xqboost
 - works about the same
 - CatBoost
 - great if you have a lot of categorical variables
 - uses something called mean target encoding (https://www.youtube.com/watch?v=V5158Oug4W8, see around ~13:00)
- L1 Regularization (Lasso Regression), L2 Regularization (Ridge Regression), Gaussian Prior
- Dendograms, Agglomerative Hierarchical Clustering
 - https://www.datanovia.com/en/lessons/agglomerative-hierarchicalclustering/#dendrogram
- fit_transform # ignores dependent var, https://stackoverflow.com/questions/ 24458645/label-encoding-across-multiple-columns-in-scikit-learn

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- GridSearchCV vs random search
 - learning rate
 - https://medium.com/datadriveninvestor/an-introduction-to-grid-search-ff57adcc0998

- Precision, Recall, and F1 Score
 - https://machinelearningmastery.com/how-to-calculate-precision-recall-f1-and-more-for-deep-learning-models/
 - https://scikit-learn.org/stable/auto_examples/model_selection/ plot_precision_recall.html (great read)
 - Precision (P) is defined as the number of true positives (T_p) over the number of true positives plus the number of false positives (F_p)

-
$$P = T_p / (T_p + F_p)$$

- Recall (R) is defined as the number of true positives (T_p) over the number of true positives plus the number of false negatives (F_n)

-
$$R = T_p / (T_p + F_n)$$

- Precision and Recall are also related to the F1 score

$$-$$
 F1 = 2 * ((P * R) / (P + R))

- Note that the precision may not decrease with recall. The definition
 of precision shows that lowering the threshold of a classifier may
 increase the denominator, by increasing the number of results
 returned. If the threshold was previously set too high, the new
 results may all be true positives, which will increase precision.
 - For example,

-
$$P = 10 / (10 + 1) = 10/11$$

- If the previous threshold was about right or too low, further lowering the threshold will introduce false positives, decreasing precision.
 - For example,

-
$$P = 8 / (8 + 3) = 8/11$$

- Recall is defined as (see formula for R above), where T_p + F_n does not depend on the classifier threshold. This means that lowering the classifier threshold may increase recall, by increasing the number of true positive results. It is also possible that lowering the threshold may leave recall unchanged, while the precision fluctuates.
- Precision-recall curves are typically used in binary classification to study the output of a classifier. In order to extend the precisionrecall curve and average precision to multi-class or multi-label classification, it is necessary to binarize the output. One curve can be drawn per label, but one can also draw a precision-recall curve by considering each element of the label indicator matrix as a binary prediction (micro-averaging).
- f1 score
 - https://scikit-learn.org/stable/modules/generated/ sklearn.metrics.f1_score.html
- .ffill
 - https://www.geeksforgeeks.org/python-pandas-dataframe-ffill/
- DecisionTreeRegressor
- BaggingRegressor
- RandomForestRegressor

- ExtraTreesRegressor
- AdaBoostRegressor
- GradientBoostingRegressor
- XGBRegressor
- KerasRegressor
- How to determine how many n_estimators we need for most of the regressors
- Memory Usage Feature in Jupyter Notebook
- Survival Analysis
 - https://stats.idre.ucla.edu/stata/seminars/stata-survival/
- Data Engineering
 - Goals of a Data Engineer
 - Developing data pipelines (most undergrad courses don't teach you how to do this)
 - Taking data from an operational system and moving it to something so it can be used by analysts and data scientists
 - Manage tables and data sets for analysts and data scientists
 - Design with the product in mind
 - What data is being tracked, what questions are the users going to be asking while they are using the product (for example, dashboards)
 - Skills of a Data Engineer
 - Data Modeling
 - Relational Databases
 - Note, databases prefer data in long format
 - Data Warehouses
 - Automation
 - Timing (isolated task/function at a specific time routinely)
 - Dependencies (for example, we want to make sure task/ table A comes before task/table B)
 - Failures (you want to make sure you capture your failures like a notification system (email, conbon?)
 - ETL Development (there is another type called ELT, essentially Data Pipeline, means Extract Transformation Label)
 - For Example, taking data from an operational system such as Workday (an HR system) or MySQL (if its a product), and moving this data into a data warehouse. The old school way is to use SQL Server, Oracle databases, etc as the data warehouse. The more modern way is Hadoop, Redshift, etc.
 - Also in ETL development, data engineers can also be in charged of implementing an analytical data layer on the data to aggregate it so it can be easily fed to dashboards.
 - Product Understanding (data is our product, we want to understand what the data scientists want)
 - Dashboards
 - Datasets
 - Data engineers develop data pipelines to

- improve ease of data access
- produce analytical layers for dashboards
- clean up data (eliminate duplicate data, etc)
 - we want data scientists to know that every observation they get is that its accurate/valid)
- Tools Used
 - Pipeline Framework (SSIS (GUI, hard to customize with Python), Infromatica, Airflow (AirBnB's version of a data pipeline management system, easy to customize)), Spark
 - SQL
 - It is easier to hire someone who knows SQL instead of map/reduce in Hadoop, Hive, etc
 - Python, Powershell, Linux and/or Java (Java is for map/ reduce stuff)
 - Tableau (heavily used, easiest), D3.js (fun and customizable, but painful if you are not Java script fan), SSRS, etc.
- How does a Data Engineer make an impact?
 - Creating optimized pipelines
 - influencing
 - designing
 - maintaining
- Note, Business Intelligence and Data Engineering are now more of the same.

Undersampling vs Oversampling

GENERAL

- Great Reads
 - LinkedIn post by NABIH IBRAHIM BAWAZIR
 - Here are some essential math/stats for #DataScience

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- Theory

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- 1. Linear Algebra (Matrices, Vectors, Eigenvalues/Eigenvectors, Linear Transformations) https://lnkd.in/gMzkkup
- 2. Basic Calculus (Derivatives & Integrals) https://lnkd.in/gDg4Nsz

- 3. Optimization (Gradient Algorithms & Objective Functions) https://lnkd.in/g_e9sJu
- 4. Inferential Statistics (Distributions, CLT, Hypothesis Testing, Erors, ANOVA, Chi-Square, T-Test)- https://lnkd.in/gbh3aRj
- 5. Probability Theory (Random Variables, Types of Distributions, Sampling, CI) - https://lnkd.in/gf6q8FN
- 6. Graph Theory (Trees, Nodes, Edges) https://lnkd.in/gYUgBhA
- 7. Data Structures (Algorithms, Big-O, Sorting, Time Complexity) https://lnkd.in/gHZEw3d

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- If you want to apply Machine Learning on Business

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- 1. Data Science Process https://lnkd.in/fMHtxYP
- 2. Data Visualization in Business https://lnkd.in/fYUCzgC
- 3. Understand How to answer Why https://lnkd.in/f396Dqg
- 4. Know ML Key Terminology https://lnkd.in/fCihY9W
- 5. Understand ML Implementation https://lnkd.in/f5aUbBM
- 6. ML Applications on Marketing https://lnkd.in/fUDGAQW
- and Retail https://lnkd.in/fihPTJf

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- Definitely brush up on these concepts and you'll be great.

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- Overfitting occurs when there is greater accuracy seen with the training set than the test set
- Anything model that is built with X_train and y_train is considered supervised learning. Any model but with just X_train is considered unsupervised learning.
- Features are column-wise data

- Supervised Learning vs Unsupervised Learning vs Reinforcement Learning (note, I need to un indent this over)
 - The main difference between the two types is that supervised learning is done using a ground truth, or in other words, we have prior knowledge of what the output values for our samples should be.
 - Supervised Learning
 - the goal of supervised learning is to learn a function that, given a sample of data and desired outputs, best approximates the relationship between input and output observable in the data.
 - Classification or Regression
 - Classification:
 - Regression:
 - Common Algorithms:
 - logistic regression
 - naive bayes
 - support vector machines
 - artificial neural networks
 - random forests
 - In both regression and classification, the goal is to find specific relationships or structure in the input data that allow us to effectively produce correct output data. Note that "correct" output is determined entirely from the training data, so while we do have a ground truth that our model will assume is true, it is not to say that data labels are always correct in real-world situations. Noisy, or incorrect, data labels will clearly reduce the effectiveness of your model. When conducting supervised learning, the main considerations are model complexity, and the biasvariance tradeoff. Note that both of these are interrelated.
 - When conducting supervised learning, the main considerations are model complexity, and the biasvariance tradeoff. Note that both of these are interrelated.
 - The bias-variance tradeoff also relates to model generalization. In any model, there is a balance between bias, which is the constant error term, and variance, which is the amount by which the error may vary between different training sets. So, high bias and low variance would be a model that is consistently wrong 20% of the time, whereas a low bias and high variance model would be a model that can be wrong anywhere from 5%-50% of the time, depending on the data used to train it. Note that bias and variance typically move in opposite directions of each other:

increasing bias will usually lead to lower variance, and vice versa. When making your model, your specific problem and the nature of your data should allow you to make an informed decision on where to fall on the bias-variance spectrum. Generally, increasing bias (and decreasing variance) results in models with relatively guaranteed baseline levels of performance, which may be critical in certain tasks. Additionally, in order to produce models that generalize well, the variance of your model should scale with the size and complexity of your training data — small, simple datasets should usually be learned with low-variance models, and large, complex data-sets will often require higher-variance models to fully learn the structure of the data.

- Unsupervised Learning
 - Does not have labeled outputs, so its goal is to infer the natural structure present within a set of data points
 - The most common tasks within unsupervised learning are clustering, representation learning, and density estimation. In all of these cases, we wish to learn the inherent structure of our data without using explicitlyprovided labels.
 - Common Algorithms:
 - K-Means Clustering
 - Principal Component Analysis (PCA)
 - Autoencoders
 - Some common use-cases for unsupervised learning are exploratory analysis (I thinkk, this is apriori) and dimensionality reduction (PCA, LDA)
- Reinforcement Learning

- Decision Tree Terminology
 - The very top of the tree is called the "Root Node" or just "The Root". It represents the entire population or sample, and this further gets divided into two or more homogeneous sets.
 - "Internal/Child Nodes" have arrows pointing to them and away from them
 - "Leaf Nodes" only have arrows pointing to them
 - "Splitting" is dividing the root node/sub node into different parts on the basis of some condition

- "Pruning" is the opposite of splitting, basically removing unwanted branches from the trees. In other words, pruning is a method of limiting tree depth to reduce overfitting in decision trees. There are two types of pruning:
 - Pre-pruning: A decision tree involves setting the parameters of a decision tree before building it. For example, setting the maximum tree depth, maximum number of terminal nodes, minimum samples for a node split (controls the size of the resultant terminal nodes), maximum number of features
 - Post-pruning: To post-prune, validate the performance of the model on a test set. Afterwards, cut back splits that seem to result from overfitting noise in the training set. Pruning these splits dampens the noise in the data set.
 - Post-pruning may result in overfitting the model and is currently not available in Pythons sklearn, but its available in R
- "Branch/SubTree" is formed by splitting the tree/node
- "Entropy" & "Information Gain"
 - https://homes.cs.washington.edu/~shapiro/EE596/notes/ InfoGain.pdf
 - https://www.youtube.com/watch?v=qDcl-FRnwSU&t=235s (great example, 27:00, 34:00)
 - Entropy
 - Common way to measure impurity. This value is used in the information gain formula.
 - Information Gain
 - Measures the reduction in entropy. Decides which attribute should be selected as the decision node. We select the attribute with the maximum gain to be the root node.
 - Also, in other words, which ever feature has the lowest Gini Impurity will be the one that provides the greatest information gain. Therefore, this feature will be selected as the node and so forth for the remaining nodes.
- Important decision trees can be constructed using binary data, continuous data, and categorical/ranked data:
 - Example with binary data (3 independent vars, 1 dependent var, all binary data (0,1))
 - See @03:25 https://www.youtube.com/watch?v=7VeUPuFGJHk
 - To begin a decision tree, you calculate the Gini impurity using each independent var with the dependent var one at a time and keep track of all the true pos, false pos, false neg, true neg. Which ever feature shows the lowest *total* Gini impurity is the one selected to start the root node. Next, the *subsection* Gini impurity's that were created for the "true pos, false pos" block and "false neg, true neg" block are the splits for that node. Now let's start with going down the left side of

the root node (the true pos, false pos) side. We must compare the Gini impurity found in this section to the Gini impurities of the other features (note, these features have to fall under the left side of the of the root node too). If a smaller Gini impurity is found in one of the other features, then it becomes the node for this level of the tree. Therefore, in a sense, it overrides the "true pos, false pos" section of the root node.

- Example with continuous data (1 independent var (continuous data), 1 dependent var (binary data))
 - See @13:57 https://www.youtube.com/watch?
 v=7VeUPuFGJHk
 - To begin a decision tree with the independent var as continuous, you first have to sort the column data (eg, 125, 135, 155, and so forth (make sure the dependent vars stay with the appropriate ones)). Then you take an average between each observation (130, 145). From there, you then start calculating the *subsection* Gini Impurities, and then get the *total* Gini impurity. See the link directly above.
- Example with ranked/categorical data (1 independent var (categorical data), 1 dependent var (binary data))
 - See @15:25 https://www.youtube.com/watch?
 v=7VeUPuFGJHk
 - Need to revisit
- Advantages: Decision trees are easy to interpret. To build a decision tree requires little data preparation from the user - there is no need to normalize the data
- Disadvantages: Decision trees are likely to overfit noisy data. The probability of overfitting on noise increases as a tree gets deeper.
- Ensembles
 - Creating ensembles involves aggregating the results of different models. Ensemble decision trees are used in bagging and random forests while ensemble regression trees are used in boosting
- Bagging/Bootstrap aggregating
 - Bagging involves creating multiple decision trees each trained on a different bootstrap sample of the data. Because bootstrapping involves samples with replacement, some of the data in the sample is left out of each tree.
 - Consequently, the decision trees created are made using different samples which solves the problem of overfitting to the training sample. Ensembling decision trees in this way helps reduce the total error because variance of the model continues to decrease with each new tree added without an increase in the bias of the ensemble.
- Random Forest
 - A bag of decision trees that uses subspace sampling is referred to as a random forest. Only a selection of the features is considered at

- each node split which decor relates the trees in the forest.
- Another advantage of random forests is that they have an in-built validation mechanism. Because only a percentage of the data is used for each model, an out-of-bag error of the moles performance can be calculated using the 37% of the sample left out of each model

- Boosting

- Boosting involves aggregating a collection of weak learners(regression trees) to form a strong predictor. Ad boosted model is built over time adding a new tree into the model that minimizes the error by previous learners. This is done by fitting the end tree on the residuals of the previous trees.
- If it isn't clear this far, for many real-world applications a single decision tree is not a preferable classification as it is likely to overfit and generalize very poorly to new examples. However, an ensemble of decision or regression trees minimizes the overfitting disadvantage and these models become stellar, state of the art classification and regression algorithms.

Model Selection

- Regression
 - Cross-Validation
 - MAE
 - RSME
- Classification
 - Confusion Matrix
- Selecting the best model in scikit-learn using cross-validation
 - What is the drawback of using the train/test split procedure for model evaluation?
 - How does K-fold cross-validation overcome this limitation?
 - How can cross validation be used for selecting tuning parameters, choosing between models, and selecting features?
 - What are some possible improvements to cross-validation?
- K Fold Validation
 - This is used to over come importing just one random state and testing it. For example, if cv=10 is set in the cross_val_score model, 10 increments will be tested as the test set with the remaining data being used as the training set each time.

DATA PREPROCESSING

- Terminology
 - feature scaling/normalization/
 - Checking for NaN values in Pandas

- <u>https://stackoverflow.com/questions/29530232/how-to-check-if-any-value-is-nan-in-a-pandas-dataframe</u>
- Data Transformation
 - Why should we transform data when it is already clean?
 - Different features in the data set may have values in different ranges. For example, in an employee data set, the range of salary feature may lie from thousands to lakhs but the range of values of age feature will be in 20- 60. That means a column is more weighted compared to other.
 - Two most common methods for normalization:
 - Min-Max
 - Min- Max tries to get the values closer to mean. But when there are outliers in the data which are important and we don't want to loose their impact, we go with Z score normalization.
 - Z score
 - Skewness of data:
 - According to Wikipedia," In probability theory and statistics, skewness is a measure of the asymmetry of the probability distribution of a real-valued random variable about its mean."
 - Generally, if the skewness value lies above +1 or BELOW -1, data is highly skewed. If it lies between +0.5 to -0.5, it is moderately skewed. If the value is 0, then the data is symmetric
 - It is also important to make sure the absolute value of the skewness is greater than twice the std error value (04:45, https://www.youtube.com/watch?
 v=_c3dVTRIK9c)
 - NOTE, we want to be very careful when sampling data sets if the results are highly skewed. For example, in the fraud analysis I am going through https://www.kaggle.com/janiobachmann/credit-fraud-dealing-with-imbalanced-datasets there are 99.83% of cases that are not fraud and 0.17 % of cases that are. Therefore, if we did sampling with this dataset, we may not get any of the fraud cases in our training set.
 - Heat maps
 - are great for seeing correlated variables
 (<myDataFrameName1>.corr() —> with the Pandas
 dataframe, this gives a print out of all the correlation
 coefficients).
 - Copy a dataframe
 - temp3 = <myDataFrameName1>.copy() # our dataframe

- Data Frame Types

- Calling <myDataFrameName1>.info() will show us information related to the Dataframe
 - "Object" are typically string values
- Imputer
 - import Imputes module to handle missing data
- Encoding variables
 - Terminology
 - Same Thing
 - Dummy Encoding (if you are coming from the statistics field)
 - One Hot Encoding (if you are coming from the computer science or electrical engineering field)
 - eg, male and female into 1 and 0
 - eg, France, Spain, and Germany into 0, 2,and 1 (note, we have to be careful of the dummy variable trap here. We must use OneHotEncoder to create ONLY TWO new columns that show only 1 and 0's). We do not need three columns for each country. We need one minus for some reason. I think this has something to do with the degrees of freedom. We are avoiding what is called the Dummy Variable Trap.
- Feature scaling
 - 5.3.1.3. Scaling data with outliers
 - If your data contains many outliers, scaling using the mean and variance of the data is likely to not work very well. In these cases, you can use robust_scale and RobustScaler as drop-in replacements instead. They use more robust estimates for the center and range of your data. (see https://scikit-learn.org/stable/modules/preprocessing.html#scaling-data-with-outliers)
 - Feature extraction is very different from Feature selection: the former (feature extraction) consists in transforming arbitrary data, such as text or images, into numerical features usable for machine learning. The latter is a machine learning technique applied on these features.
 - Examples of Algorithms where Feature Scaling matters
 - K-Means uses the Euclidean distance measure here feature scaling matters.
 - K-Nearest-Neighbours also require feature scaling.
 - Principal Component Analysis (PCA): Tries to get the feature with maximum variance, here too feature scaling is required.
 - Gradient Descent: Calculation speed increase as Theta calculation becomes faster after feature scaling.
 - Note: Naive Bayes, Linear Discriminant Analysis, and Tree-Based models are not affected by feature scaling. In Short, any Algorithm which is Not Distance based is Not affected by Feature Scaling.

Good Reads

- https://stats.stackexchange.com/questions/189652/is-it-a-good-practiceto-always-scale-normalize-data-for-machine-learning
- https://stats.stackexchange.com/questions/342140/standardization-of-continuous-variables-in-binary-logistic-regression (eg, 2 regressors are

continuous and 2 are binary)

- <u>https://datascience.stackexchange.com/questions/20237/why-do-we-convert-skewed-data-into-a-normal-distribution</u>
- Parametric and Nonparametric: Demystifying the Terms
 - Parametric
 - We have to normally distributed datasets (not positively or negatively skewed)
 - Common transformations of this data include square root, cube root, and log10.
 - The cube root transformation involves converting x to x^(1/3). This is a fairly strong transformation with a substantial effect on distribution shape: but is weaker than the logarithm. It can be applied to negative and zero values too. Negatively skewed data.
 - The square root transformation, x^2 can only be applied to positive values only. Hence, observe the values of column before applying.
 - The logarithm transformation, x to log base 10 of x, or x to log base e of x (ln x), or x to log base 2 of x, is a strong transformation and can be used to reduce right skewness (positively skewed).
 - If tail is on the right as that of the second image in the figure, it is right skewed data. It is also called positive skewed data.
 - In order to fix a positively skewed distribution using a log10 distribution, the following assumptions have to be met:
 - no negative values, no zeroes, and the positively skewed (if you have negative values or zeros, see around ~08:00 mins, https://www.youtube.com/watch?v=_c3dVTRIK9c)
 - If the tail is to the left of data, then it is called left skewed data. It is also called negatively skewed data.
 - Similarly, to fix a negatively skewed distribution using a log10 distribution, the same conditions have to be met:
 - but this time we call the log10 function as
 - $\log 10(\max(x) 1 + x)$
- Resolving outliers
 - Anomaly Detection (best seen from boxplot)
 - Interquatile Range Method
- Aggregation of data

CLASSIFICATION

- Decision Tree Classification
 - graphical representation of all the possible solutions to a decision

- decisions are based on some conditions
- decisions made can be easily explained
 - Example
 - Independent variables: "Am I hungry?" "Do I have \$25?"
 - Data is in binary form (yes or no)
 - Dependent variable: "Outcome"
 - Data is "Go to restaurant" "Buy a hamburger" "Go to sleep" (this is a multiple classification problem)
- NOTE: When building a decision tree you first have to individually compare each independent variable to the dependent variable in order to determine which independent variable will begin the decision tree (classification problem, yes or no for all results for the independent and dependent variable, you keep track of which independent variables that has the lowest "Gini Impurity" (aka the one that is most pure) and you begin with that one at the top of the tree.. See https://www.youtube.com/watch?v=7VeUPuFGJHk&list=PLsCzljdLz2iR66dIIpz9E9veNj7wijPS7. I thinkkk you can also look at the true pos, false pos, true neg, false neg.
- Important: Decision trees for classification use Gini Impurity and/or Information Gain. Decision trees for regression use Standard Deviation Reduction (see http://saedsayad.com/decision_tree_reg.htm).
- Random Forest Classification
 - Builds multiple decision trees and merges them together
 - more accurate and stable prediction
 - random decision forests correct for decision trees' habit of overfitting to their training set
 - trained with the "bagging" method
- Naïve Bayes Classification
 - Classification technique based on Bayes' Theorem
 - Assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature
- K Nearest Neighbors Classifier
 - Stores all the available cases and classifies new cases based on a similarity measure
 - The "K" is the KNN algorithm is the nearest neighbors we wish to take a vote from
 - Example:
 - Imagine a scatter plot with real training data on it representing 5 classes. Each dot on the scatter plot is a color. Generally, the color dots in this training set data are grouped by similar color dots. So visually, you can see five classes. Now, I think the KNN algorithm builds the model by looking at super small segments of the plot and starts a computational radius going outward from each segment. So if K is specified to be "3." The computational radius will stop expanding for each segment when it hits three dots of the

same color and will give that segment of the plot the appropriate color.

from lightgbm import LightGBMClassifier

REGRESSION

Decision Tree Regression

- For example, when visualizing two independent variables on a scatter plot, a decision tree algorithm "splits" your dataset based off of information entropy (need to look up more) and based off of these splits is how the tree is made.
 - https://www.saedsayad.com/decision_tree_reg.htm (example)
- Important: Decision trees for classification use Gini Impurity and/or Information Gain. Decision trees for regression use Standard Deviation Reduction (see http://saedsayad.com/decision_tree_reg.htm).
- Random Forest Regression
 - a form of ensemble learning, ensemble learning is when you take the same algorithm multiple times to make your model much more powerful than the original. Gradient boosting is a form of ensemble learning.
 - Not very sensitive to hyperparameters (need to verify still)
- Support Vector Regression
 - Great video —> https://www.youtube.com/watch?v=Y6RRHw9uN9o&t=366s
 - Dog and cat example. X axis there is snout length and on the Y axis there is ear length. SVR algorithm looks at the outliers within the data set and applies a linear linear / nonlinear line according to them. These outliers are none as support vectors. Note, in the example in the video the data is separable between the cat and dog classes.
 - When using Support Vector algorithms it is often helpful to map your vectors to a higher dimension in order to fully (better?) separate the data. However, this is often computationally expensive.
 - See Kernel function or Kernel Trick. The dot product can be computed to project the vectors into a higher dimension.
- Logistic Regression
 - Note, logistic regression is probably better placed under CLASSIFICATION.
 - Great explanation of how to interpret the coefficients in a logistic or linear regression model (see ~02:48:00, https://www.youtube.com/watch?v=5rNu16O3YNE&list=PLsCzljdLz2iR9pnDlZkTqSvbUZqZenhck&index=5)
 - Has to do only with probability. Think about example of where a bank decides they should allow you to get a loan or not depending on your credit score, income, age, etc..

- Or think about an image data set of digits where the target variable is 0-9. You can do classification binary or multi class problems with LogisticRegression.
- Not very sensitive to hyperparameters (need to verify still)
- Linear Regression
 - Great explanation of how to interpret the coefficients in a logistic or linear regression model (see ~02:48:00, https://www.youtube.com/watch?v=5rNu16O3YNE&list=PLsCzljdLz2iR9pnDlZkTqSvbUZqZenhcK&index=5)

CLUSTERING:

- Hierarchial Clustering
- Clustering
 - K-Means
 - Very good for discovering categories or groups in your data that you may of not been able to recognize yourself.
 - Can work for multiple dimension problems
 - How it works (imagine a 2D scatter plot with a bunch of gray markers and we want to separate our data into 3 clusters (red, green, and blue)):
 - https://www.udemy.com/course/machinelearning/learn/lecture/5714416#overview (great example)
 - Step 1. Choose the number K of clusters
 - Step 2. Select **at random** K points. The centroids (they don't not necessarily have to be from or around your dataset).
 - Step 3. Assign each data point on the plot to the closest centroid that will in result form K clusters (generally in the form of Euclidean distance (there are other types, need to look up))
 - Step 4. Compute and place the new centroid of each cluster.
 - Step 5. Reassign each data point to the new closest centroid. If any reassignment took place, go back to Step 4, otherwise go to finish.
 - This is an iterative process.
 - The algorithm runs quickly by drawing a straight line connecting to each centroid and then drawing a perpendicular line from that line easily separate the data set and see which data points are closest (well, Im not sure if that's how the algorithm actually runs, but that is a great visual way to think about it, this would be really good to show on a technical presentation).

MACHINE LEARNING ALGORITHMS

- A/B Testing
 - Example, comparing which landing page of a website performs best.
 - A/A tests should be conducted first to make sure there's nothing wrong on the backend such as if the data is messy, sampling problem because not randomizing properly, or too much noise.
- KNeighborsRegressor (aka KNN)
 - https://scikit-learn.org/stable/auto_examples/neighbors/ plot_regression.html#sphx-glr-auto-examples-neighbors-plot-regressionpy
 - https://en.wikipedia.org/wiki/K-nearest_neighbors_algorithm (great explanation of the weights parameter)
- DecisionTreeRegressor (aka CART)
- SVM
 - C-Parameter:
 - https://stats.stackexchange.com/questions/31066/what-is-the-influence-of-c-in-svms-with-linear-kernel see Kent Munthe Caspersen answer
 - "In general, having few training instances and many attributes make it easier to make a linear separation of the data."
- xgboost
 - https://xgboost.readthedocs.io/en/latest/tutorials/index.html

ASSOCIATED RULE LEARNING

- Apriori Algorithm (think of grocery cart example —> diapers and beer relationship)
- FP Growth Model

REINFORCEMENT LEARNING

- Upper Confidence Bound Algorithm (REVISIT 2019-08-10 21:33:37, DONT UNDERSTAND FULLY)
 - Multi Armed Bandit Problem
- Thompson Sampling
 - Multi Armed Bandit Problem

- Important Terminology
 - Explore vs. Exploitation

METRICS

- Use the sklearn.metrics to create a confusion matrix to analyze true negatives, false negatives, true positives, and false positives
- See the section Model Selection & Boosting below for a better way to look at the performance of your models (GridSearchCV)
- https://www.theanalysisfactor.com/assessing-the-fit-of-regression-models/
 - Lower values of RMSE indicate better fit.

NATURAL LANGUAGE PROCESSING

- rule learning
- sparsity
- sparse matrices

DEEP LEARNING

- Important Terminology
 - Output values of the neural network can either be continuous (eg, price), binary (eg, yes/no), or categorical
 - weighted sum, activation function (sigmoid, threshold, rectifier, hyperbolic tangent)
 - Think about churn modeling problem (trying to predict which customers will leave the bank or not)

ARTIFICIAL NEURAL NETWORKS

NEURAL NETWORKS

- Additional Reading
 - https://stats.stackexchange.com/questions/154879/a-list-of-cost-functions-used-in-neural-networks-alongside-applications
- How do neural networks learn?
 - You can either hard code them or let them learn on their own. We are always trying to let the models learn on their own.
 - Basically the only thing we have control over in the programming are the weights
 - Batch gradient descent (think of the smooth parabolic cost function

curve with the ball bouncing back and forth trying to find the minimum)

- Stochastic gradient descent (this method is used to find the global minimum of a rough shaped parabolic cost function curve) this method is faster than batch (there is also a mini-batch method)
 - https://iamtrask.github.io/2015/07/27/python-network-part2/

Important Terminology

- Back propagation (the process of optimizing the cost function (aka adjusting the weights to find optimal value to agree with the known value))
- Normally the rectifier function is used for the development of the hidden layers and for the output layer the sigmoid function is used.

CONVOLUTIONAL NEURAL NETWORKS

- Think about the following images: duck or rabbit, man looking at you or away from you, and image of the girl with duplicate facial features (eyes, nose, mouth, eyebrows)
- Steps -> Convolution, Max Pooling, Flattening, and Full Connection
 - Convolution: The key take away of what convolution is, is to find features in your image by using a feature detector (think of a feature detector as a specific feature that distinguishes an image from the other images) and putting them into a feature map. Then from the feature map, it preserves the spatial relationships between pixels
 - Convolution (Cont.): ReLU Layer —> we introduce this Rectifier Linear Units algorithm to break up linearity.
 - Good read on convolution filters: <a href="http://www.roborealm.com/http://www.robo
 - Max Pooling: Looking for a specific feature of an image. For example, the tears/marks on a cheetah. It is one feature that makes the cheetah very distinct. Note, there are other forms of pooling (mean pooling, sum pooling, etc). In the example I was following on Udemy, the instructor mentioned that max pooling eliminates the need of unnecessary features. In the cheetah example, we were able to get rid of 75% of the information. Max pooling allows us to instantly see specific distinct regions. In summary, we are able to preserve distinct features, inducing spatial variance, and reducing the size of information and parameters (this helps us prevent over fitting)
 - http://scs.ryerson.ca/~aharley/vis/conv/flat.html
 - http://ais.uni-bonn.de/papers/icann2010_maxpool.pdf
 - http://scs.ryerson.ca/~aharley/vis/conv/
 - http://www.cs.cmu.edu/~aharley/
 - <u>Flattening</u>: For example, making a 3x3 feature map display vertically 1-9
 - Full Connection: The middle/inner hidden layers (these develop

- attributes that describe each output). The final inner layer nodes each pass on a vote/probability from 0.0 to 1.0 on whether the image presented is a cat or dog (for example)
- <u>SUMMARY</u>: https://adeshpande3.github.io/The-9-Deep-Learning-Papers-You-Need-To-Know-About.html
- Softmax & Cross-Entropy (Loss Function)
 - These functions behave similar to how the the mean square error (MSE) / cost function works when linear fitting
 - Classification Error, Mean Squared Error, Cross-Entropy
 - Cross-Entropy is typically the best value to look at to evaluate your neural network (note, cross-entropy involves a logarithmic function. Also, cross-entropy is only the best for classification problems)
- Image Preprocessing
 - https://keras.io/preprocessing/image/
 - To avoid overfitting, data augmentation is used.

DIMENSIONALITY REDUCTION

- Two Types
 - Feature Selection
 - Backward elimination, forward selection, bidirectional elimination, score comparison, and more (these topics were covered in the Regression lecture)
 - Feature Extraction
 - Principal Component Analysis (PCA)
 - see —> https://www.youtube.com/watch?v=_UVHneBUBW0
 and https://www.youtube.com/watch?v=FgakZw6K1QQ (great)
 and https://www.youtube.com/watch?v=HMOI_lkzW08 (NOTE: I think the cells are considered the classes/dependent variables and the genes are considered as the independent variables)
 - PCA is a form of data preprocessing I thinkk. It has nothing to do with the model/classifier. PCA is a method of compressing a lot of data into something that captures the essence of the original data. PCA reduces dimensions by focusing on the genes with the most variation. The business example I followed in the Udemy course involved analyzing 178 data observations (12 independent variables, 1 dependent variable (3 classes, each class represents a different wine)). The business was trying to predict based off the ingredients what type of wine the drink should be classified as. PCA was applied to reduce the independent variables to the ones that show the most variance. This also allows us to visualize the data in 2D or 3D better.
 - Linear Discriminant Analysis (LDA)

- see -> https://www.youtube.com/watch?v=azXCzI57Yfc (great)
- LDA is very similar to PCA. It is also used in the preprocessing step for pattern classification. Its goal is to project a dataset onto a lower-dimensional space. However, LDA differs because in addition to finding the component axises with LDA. In other words, we are interested in maximizing the separability between the two (or more) groups/categories so we can make the best decisions. We are interested in the axes that maximize the separation between classes (think of the diagram that shows gaussian distributions on both the x-axis and y-axis). The goal of LDA is to project a feature space (a dataset n-dimensional samples) onto a small subspace k(where k is less than or equal to n-1) while maintaining the class-discriminatory information.
 - Both PCA and LDA are linear transformation techniques used for dimensional reduction. PCA is described as unsupervised but LDA is supervised because of the relation to the dependent variable.
 - PCA: Component axes that maximize the variance
 - LDA: Maximizing the component axes for classseparation.
- Kernel PCA (this is a nonlinear reduction strategy)
 - This is for nonlinear problems. Note, the same linear model can be used with the normal PCA object but this time we must use an additional argument "rbf.' This accounts for the nonlinearity.

MODEL SELECTION & BOOSTING

- Two Types
 - Cross Vaidation (multiple methods)
 - https://www.youtube.com/watch?v=L_dQrZZjGDg
 - k-Fold Cross Validation
 - https://www.youtube.com/watch?v=gJo0uNL-5Qw
 - Grid Search (GridSearchCV)
 - This technique involves grid search to find optimal hyperparameters. Note, hyperparameters are the parameters you specify when you build your model.
- Boosting
 - XGBoost
 - It is the most powerful implementation of gradient boosting
 - When using XGBoost, you don't have to use feature scaling,
- Mean Absolute Error (MAE) vs Root mean squared error (RMSE)

- For continuous dependent variables only
 - great read
 - <u>https://medium.com/human-in-a-machine-world/mae-and-rmse-which-metric-is-better-e60ac3bde13d</u>

PLOTS

- Violin plots
 - https://www.youtube.com/watch?v=M6Nu59Fsyyw
- Line plot
 - ax = sns.lineplot(x="pickup_hour", y="Trip_distance", legend="full", data=summary_wdays_avg_duration, estimator=None, hue="day of week")

STATISTICS

- Pearson Coefficient
- Skewness
 - measures the symmetry of a distribution
- Kurtosis
 - measures how peaked or flat a distribution is
- Standard Error
 - https://www.investopedia.com/terms/s/standard-error.asp
- Standard Deviation
 - https://www.investopedia.com/terms/s/standarddeviation.asp
- Homoscedasticity
 - <u>https://datascience.stackexchange.com/questions/20237/why-do-we-convert-skewed-data-into-a-normal-distribution</u>
 - The errors your model commits should have the same variance, i.e. you should ensure the linear regression does not make small errors for low values of X and big errors for higher values of X. In other words, the difference between what you predict \hat{Y} and the true values Y should be constant. You can ensure that by making sure that Y follows a Gaussian distribution. (The proof is highly mathematical.)

- Commonly used functions
 - list()
 - set()
 - https://realpython.com/python-sets/
 - see the "Operators vs Methods" section (the or operator)
 - that structure comes up a lot while getting unique elements for encoding with both a training and test set
 - len()
 - range()
 - append()
 - apply()
- Syntax
 - ebola_long["variable"].str.split("_", expand=True)
 - to view the syntax better, you can rewrite as
 - (ebola_long["variable"]
 - .str
 - .split("_", expand=True)
- Functions
- Libraries
 - tflearn
 - keras
 - numba
 - Tries to make all the computations numpy does really fast
 - seaborn
 - matplotlib.pyplot
 - Anatomy of a figure
 - https://matplotlib.org/3.1.1/gallery/showcase/anatomy.html

NUMPY

- Common functions
 - np.log1p() # fixing skewed data
 - np.expm1() # done after np.log1p
 - np.linspace

PANDAS

- Dataframe
- Dataframe
 - Terminology
 - Mean the same thing

- Example 1
 - <myDataFrameName1>["<myFeatureName1>"] and <myDataFrameName1>.<myFeatureName1>
- There are three parts to a dataframe
 - <myDataFrameName1>.columns (feature names)
 - <myDataFrameName1>.index (row names)
 - <myDataFrameName1>.values (body values, this is good for extracting data from the Dataframe to use the data in another library that may just use numpy arrays)
- Dataframe information
 - <myDataFrameName1>.info()
 - "object" are typically string values
 - <myDataFrameName1>.types
 - If you extract a single column of data from a DataFrame set that new data then becomes a Series object
- aggregate()

Q&A STACKEXCHANGE

https://datascience.stackexchange.com/questions/26776/how-to-calculate-ideal-decision-tree-depth-without-overfitting

https://stackoverflow.com/questions/34162443/why-do-many-examples-use-fig-ax-plt-subplots-in-matplotlib-pyplot-python

https://stackoverflow.com/questions/3584805/in-matplotlib-what-does-the-argument-mean-in-fig-add-subplot111

https://www.geeksforgeeks.org/python-pandas-dataframe-ix/

https://stackoverflow.com/questions/31593201/how-are-iloc-ix-and-loc-different

https://stats.stackexchange.com/questions/56302/what-are-good-rmse-values

https://stackoverflow.com/questions/22409855/randomforestclassifier-vs-extratreesclassifier-in-scikit-learn

RANDOM SOURCES

https://www.kdnuggets.com/2017/10/edge-analytics.html https://medium.com/@livewithai/the-significance-of-edge-cases-and-the-cost-of-imperfection-as-it-pertains-to-ai-adoption-dc1cebeef72c

GREAT YOUTUBE CHANNELS

- Data School (https://www.youtube.com/channel/UCnVzApLJE2ljPZSeQylSEyg)
- codebasics
 - https://www.youtube.com/watch?v=gJo0uNL-5Qw (Machine Learning Tutorial Python 12 K Fold Cross Validation)
 - https://github.com/codebasics/py (see all repositories)

MEDIUM

 https://towardsdatascience.com/understanding-input-and-output-shapes-inconvolution-network-keras-f143923d56ca

----INTERVIEW

PROBLEM SOLVING TIPS

- Classification Problem
 - Typically its good to test logistic regression vs random forest classifier to see which one performs best. If the random forest classifier performs best, then we should look into boosting with either xgboost, lightgbm, or catboost

Overall process (from Allstate Severity Example):

- Data statistics
 - Shape
 - Peek
 - Description
 - Skew
- Transformation
 - Correction of skew

- Data Interaction
 - Correlation
 - Scatter plot
- Data Visualization
 - Box and density plots
 - Grouping of one hot encoded attributes
- Data Preparation
 - One hot encoding of categorical data
 - Train-test split
- Evaluation, Prediction, and Analysis
 - Linear Regression (Linear algo)
 - Ridge Regression (Linear algo)
 - LASSO Linear Regression (Linear algo)
 - Elastic Net Regression (Linear algo)
 - KNN (non-linear algo)
 - CART (non-linear algo)
 - SVM (Non-linear algo)
 - Bagged Decision Trees (Bagging)
 - Random Forest (Bagging)
 - Extra Trees (Bagging)
 - AdaBoost (Boosting)
 - Stochastic Gradient Boosting (Boosting)
 - MLP (Deep Learning)
 - XGBoost
 - Make Predictions

##########################

CODE SNIPPETS

-

- Jupyter Notebook / Markdown
 - Github
 - This is a self-exercise notebook that was created while following along the Natural Language Processing section in the following Udemy course by SuperDataScience:
 - This is a self-exercise notebook that was created while following along in the following YouTube video by codebasics:
- Python
 - Important
 - There seems to be a lot of issues with copying snippets over from TextEdit to Jupyter. It's something to do with single quotes. Change them to double quotes.
 - Snippet Template Names
 - <myDataFrameName1>
 - \diamond

- <mvFeatureName1> <myFeatureName1_categorical> <myFeatureName1 continuous> <myVarName1> <mvModelName1> <myXTestName1> <myFunctionName1> <myInt1> <myIndexName1> **Pandas** Info # this is really useful when trying to pandas. version get help from Google or StackOverflow [] # single brackets are for specifying if you are in rows or columns [[]] # inner brackets are typically used when you want to implement a list/multiple things [["year', "date']] # example <myDataFrameName1>["<myFeatureName1>"] is the same thing as <myDataFrameName1>.<myFeatureName1> Set Options pd.set_option('display.max_columns', 999) # allows us to see all columns in Jupyter notebook Converting Data Types <myDataFrameName1>[<myFeatureNameList1>] = <myDataFrameName1>[<myFeatureNameList1>].apply(pd.t o numeric, errors='coerce') Commonly used timeit example %%timeit # line 1 in Jupyter cell <myFunctionName1>(<myDataFrameName1>["<myFeature" Name1>"].values, <myDataFrameName1>["<myFeatureName2>"].values) # line 2 in Jupyter cell
 - print(train_X.shape)
 - print(type(train_X.shape))
 - Commonly used libraries
 - import pandas as pd
 - import numpy as np
 - import matplotlib.pyplot as plt

import warnings warnings.filterwarnings('ignore') import tensorflow as tf tf.logging.set_verbosity(tf.logging.ERROR) # To surpress any TensorFlow warnings. from sklearn.preprocessing import LabelEncoder, OneHotEncoder from sklearn.model_selection import train_test_split from sklearn.metrics import mean absolute error from sklearn.linear_model import LinearRegression from sklearn.neighbors import KNeighborsRegressor # KNN (Non-linear Algo) from sklearn.tree import DecisionTreeRegressor # CART from sklearn.svm import SVR from sklearn.ensemble import BaggingRegressor from sklearn.ensemble import RandomForestRegressor from sklearn.ensemble import ExtraTreesRegressor from sklearn.ensemble import AdaBoostRegressor from sklearn.ensemble import GradientBoostingRegressor from xgboost import XGBRegressor import re # used often for NLP, removes and replaces characters from keras.models import Sequential from keras.layers import Convolution2D from keras.layers import MaxPooling2D from keras.layers import Flatten from keras.layers import Dense from keras.preprocessing.image import ImageDataGenerator https://keras.io/preprocessing/image/ #Import libraries for deep learning from keras.wrappers.scikit_learn import KerasRegressor from keras.models import Sequential from keras.layers import Dense # Optimize using dropout and decay

import seaborn as sns

```
from keras.constraints import maxnorm
Encoding
      pandas
            Example 1
                  <myDataFrameName1> = sns.load_dataset("tips")
                  <myDataFrameName1>.head()
                  <myDataFrameName2> =
                  pd.get_dummies(<myDataFrameName1>,
                  drop_first=True)
                  <myDataFrameName2>.head()
      sklearn
            Example 1
                  from sklearn.preprocessing import LabelEncoder,
                  OneHotEncoder
                  <myDataFrameName1> = pd.read_csv("train.csv")
                  <myDataFrameName2> = pd.read_csv("test.csv")
                  cols = <myDataFrameName1>.columns
                  split = <myIntOfCategoricalColumnsToEncode>
                  labels = []
                  # Making sure we account for all of the unique
                  variables that show up in both the training and test set
                  provided.
```

from keras.optimizers import SGD from keras.layers import Dropout

```
# For instance, this ensures we dont run into any
unforeseen variables when going from the training set
to test set.
for i in range(0, split):
   train = <myDataFrameName1>[cols[i]].unique()
   test = <myDataFrameName2>[cols[i]].unique()
   labels.append(list(set(train) | set(test))) #note the
OR operator! (See how sets work https://
realpython.com/python-sets/)
print("labels %s" % labels)
cats = []
for i in range(0, split):
  label_encoder = LabelEncoder() # Label Encode
  label encoder.fit(labels[i])
  feature =
label_encoder.transform(dataset_train.iloc[:,i])
  feature = feature.reshape(dataset_train.shape[0],
1)
  onehot encoder =
OneHotEncoder(sparse=False,n_values=len(labels[i])
      # One Hot Encode
  feature = onehot encoder.fit transform(feature)
  cats.append(feature)
print("########")
print("List of 1D array of cats--> %s" % cats)
print("#########")
# Make a 2D array from a list of 1D arrays
encoded_cats = np.column_stack(cats)
print("#########")
print("List of 2D array of cats--> %s" % encoded_cats)
print("#########")
print("########")
print(encoded_cats.shape) # These are the
categorical variables we just encoded. Now, we just
need to concatenate it with the continuous vars.
```

- print("##########")
- _
- #Concatenate encoded attributes with continuous attributes
- <myDataFrameName3> =
 np.concatenate((encoded_cats,dataset_train.iloc[:,spli
 t:].values),axis=1)

_

- del cats
- del dataset train
- del encoded_cats

-

- Data Set Details
 - <myDataFrameName1>.info() # data info. ex: datatypes, size etc.
 - <myDataFrameName1>.dtypes
 - <myDataFrameName1>.describe()
 - <myDataFrameName1>.skew() # Values close to 0 show less skew.
 - <myDataFrameName1>.shape
 - Column Details
 - <myVarName1> =
 <myDataFrameName1>["<myFeatureName1>'].min()
 # minimum trip distance, same for max()
 - <myVarName2> =
 <myDataFrameName1>["<myFeatureName1>'].value
 _counts() # counts unique occurrences
 - <myVarName2>.head(10)
 - Searching for NaN Values
 - <u>https://stackoverflow.com/questions/29530232/how-to-check-if-any-value-is-nan-in-a-pandas-dataframe</u>
 - <myDataFrameName1>.isnull().sum()
 (searches for which features contain NaN values)
 - Renaming columns
 - To see a random sample of data
 - <myDataFrameName1>.sample(30)
 - Return names of columns where a certain character shows up
 <myDataFrameName1>.columns[<myDataFrameName1>.isi

n(['?']).any()]

- Alternative method of setting index of DataFrame (set_index)

- https://pandas.pydata.org/pandas-docs/version/0.22/generated/ pandas.DataFrame.set_index.html Shuffling the DataFrame <myDataFrameName1> = <myDataFrameName1>.sample(frac=1).reset index(drop=True) Reading In Data **CSV** Example 1 df = pd.read_csv("green_tripdata_2015-09.csv", low memory=False, parse dates=["lpep pickup datetime"]) # the parse_dates argument is turning "lpep_pickup_datetime" from an object type into a datetime64 type **TSV** Example 1 dataset = pd.read csv("Restaurant Reviews.tsv", delimiter="\t") Converting data types series and numpy array <myDataFrameName1>["<myFeatureName1>"] # this gives a series <myDataFrameName1>["<myFeatureName1>"].values this gives a numpy array Creating a DataFrame Manually <myDataFrameName1> = pd.DataFrame({ "<myFeatureName1>": [10, 20, 30], "<myFeatureName2>": [20, 30, 40] }) Operations on a DataFrame By Feature (this is called Broadcasting) <myDataFrameName1>["<myFeatureName1>"] ** 2 # this squares every row Adding rows <myDataFrameName1>["<myFeatureName1>"] + <myDataFrameName1>["<myFeatureName2>"] Copy a data frame <myDataFrameName2> = <myDataFrameName1>.copy() # our dataframe Subsetting a data frame By columns
- - Into a series
 - <myDataFrameName2> = <myDataFrameName1>["<myFeatureName1>"]
 - <myDataFrameName2>.head()

- Into a dataframe

 - <myDataFrameName2>.head() # note if we did a single column it would be a series
- By rows
 - Example 1
 - <myDataFrameName1>.loc[[2,0]] # label based indexing, note the algorithm isn't actually looking for the index. It is doing string matching instead.
 - loc is typically used
 - Example 2
 - <myDataFrameName1>.iloc[[0, 1, -1]] # another example of positional indexing, note this prints the first two rows and the last row of the data frame
- Range Selection
 - By Columns
 - Example 1

<myDataFrameName1>[["<myFeatureName1>","<myFeatureName2>"]]

- Example 2
 - # Get the column names
 - <myColumnNames1> =
 <myDataFrameName1>.columns[<myIndexInt
 1>:<myIndexInt2>]
 - # Then get the data columns

<myDataFrameName1>[<myColumnNames1>]

- By Rows
 - Example 1

<myDataFrameName1>.iloc[<myIndexInt1>:<
myIndexInt2>]

- By Columns and Rows
 - Example 1
 - <myIndexName1> =
 <myDataFrameName1>.columns[<myInt1>:<m
 yInt2>] # returns an Index of column names
 - <myDataFrameSeriesName1> =
 <myDataFrameName1>[<myIndexName1>] #
 returns a data frame series
 - <myDataFrameName2> = <myDataFrameSeriesName1>.iloc[<myInt3>:< myInt4>] # returns data frame subsetted by rows and columns

- Example 2
 - <myDataFrameName2> =
 <myDataFrameName1>[<myDataFrameName
 1>.columns[<myInt1>:<myInt2>]].iloc[<myInt3>
 :<myInt4>]
- Filtering dataframe
 - df.loc[(df["smoker"]=="No")&(df["total_bill"] >= 10)]
 # Filter rows by smoker == "No" and total_bill >= 10
 - df (["smoker","day","time"])["total_bill"].mean() # What is the average total_bill for each value of smoker, day, and time?
 - df.groupby(["smoker","day","time"])["total_bill"].mean().reset_index()
 # To get back to a Dataframe
 - df.loc[df["year"]==1967,["year", "pop"]]
 - df.loc[(df["year"]==1967) & (df["pop"] > 1_000_000),["year", "pop"]]
 # Note: Python has a neat feature where it ignores underscores in integers to help visualize.
 - $df.loc[(df["year"]==1967) | (df["pop"] > 1_000_000),["year", "pop"]]$
 - df[billboard_melt["track"] == "Loser"]
 - df.groupby(["year"])["lifeExp"].mean()
 - The aggregate function in the numpy library can do the same thing
 - df.groupby(["year"])["lifeExp"].agg(np.mean)
 - df.groupby(["year"])["lifeExp"].agg(np.std)
 - df.groupby(["year", "continent"])[["lifeExp",
 "gdpPercap"]].agg(np.mean) # note, the created
 Dataframe becomes wonky when using a list
- Fixing Skewed Data
 - #log1p function applies log(1+x) to all elements of the column
 - <myDataFrame1>["<myFeatureName1>"] =
 np.log1p(<myDataFrame1>["<myFeatureName1>"])
- Unskewing Skewed Data
 - <myVarName1> =
 np.expm1(<myModelName1>.predict(<myXTest1>))

-

- Tidy data (cleaning data)
 - http://vita.had.co.nz/papers/tidy-data.pdf (first three sections are a good read)
 - Criteria 1 (what we don"t want): Column headers are values, not variable names
 - aka going from wide data to long data
 - pew_long = pd.melt(pew, id_vars="religion")
 # doesn't touch the religion column, and condenses the rest of the data frame into a "variable" and "value" column

- pew_long = pd.melt(pew, id_vars="religion", var_name="income", value_name="count") # renames the variable and value heading.
- billboard_melt = pd.melt(billboard,
 id_vars=["year","artist","track","time","date.ente
 red"], var_name="week", value_name="rating")
 # many column example
- billboard_melt = pd.melt(billboard,
 id_vars=["year","artist","track","time","date.ente
 red"], var_name="week",
 value_name="rating").groupby("artist")
 ["rating'].mean()
- billboard_melt = pd.melt(billboard,
 id_vars=["year","artist","track","time","date.ente
 red"], var_name="week",
 value_name="rating").groupby("artist")
 ["rating'].mean().reset_index()
- Criteria 2 (what we don"t want): Multiple variables are stored in one column. #random keywords: parse
 - variable_split = ebola_long["variable"].str.split("_")
 # Note, the ".str." is called an accessor. There is a commonly used ".dt." accessor too.
 - ebola_long["status"] = variable_split.str.get(0)
 - ebola_long["country"] = variable_split.str.get(1)
 - or, you can do the above code all in one line with
 - ebola_long[["status", "country"]] =
 (ebola_long["cd_country"].str.split("_",
 expand=True)) #this is also an
 example of concatenating
- Criteria 3 (what we don"t want): Variables are stored in both rows and columns
 - Note, # if we see a lot of rows with repeated values, this is a symptom of this type of problem.
 - weather_melt = pd.melt(weather, id_vars=["id","year",
 "month", "element"], var_name="day",
 value_name="temp")
 - weather_tidy =
 weather_melt.pivot_table(index=["id","year",
 "month","day"], columns="element",values="temp")
 # note, pivot_table is the opposite of melt. Also, by
 default, this drops NaN values.
 - weather_tidy
 - weather_tidy.reset_index()
- Criteria 4 (what we don"t want): Multiple types of observational units are stored in the same table / This is also called "normalization"

- billboard_melt.loc[billboard_melt["track"] == "Loser"]
- billboard_songs = billboard_melt[["year", "artist",
 "track", "time"]]
- billboard_songs = billboard_songs.drop_duplicates()
- billboard_songs["id"] = range(len(billboard_songs))
- billboard_ratings =
 billboard_melt.merge(billboard_songs, on = ["year", "artist", "track", "time"]) # merges billboard_melt (left) and billboard_songs (right)
 - billboard_ratings = billboard_ratings[["id",
 "date.entered","week","rating"]]
- Criteria 5 (what we don't want): A single observational unit is stored in multiple tables
- Joining Tables/Datasets
 - billboard_songs["id"] = range(len(billboard_songs))
 - billboard_ratings = billboard_melt.merge(billboard_songs, on =
 ["year", "artist", "track", "time']) # merges billboard_melt (left) and billboard_songs (right)
 - billboard_ratings = billboard_ratings[["id",
 "date.entered","week","rating"]]
 - billboard ratings.head()
- Saving to csv
 - billboard_songs.to_csv("billboard_songs.csv", index=False)
- Concatenating (similar to joining)
 - X = np.concatenate((X_train, X_test), axis=0)
- Functions
 - We use what is called an assert statement to test a function. This is the basis of unit testing.
 - Example
 - assert my_sq(4) == 16 # if you run this code nothing happens, but if you change "16" to "15" the code will crash
 - Broadcasting
 - df["a"] ** 2 # this squares every row in the column labeled a
 - With a data series (specific to one feature/column)
 - Example 1
 - def my_sq(x)
 - return x ** 2
 - df["a"].apply(my_sq)
 - Example 2
 - def my_sq(x, e)
 - return x ** e
 - df["a"].apply(my_sq, e)
 - With a data frame (applies a function to each entire column of a dataframe)
 - Example 1 # the apply statement here will print all

data out in each column of the dataframe

- def print_me(x):
- return x
- df.apply(print_me)
- Example 2 # Get the mean of every column
 - def avg_apply(col):
 - return np.mean(col)
 - df.apply(avg_apply)
- Example 3 # Get the mean of every column
 - def avg_apply(col):
 - x = col[0]
 - y = col[1]
 - $\qquad z = col[2]$
 - return (x+y+z)/3
 - df.apply(avg_apply) # or we could do df.apply(avg_apply, axis="columns") to get the average of each row
- Example 4 # A more realistic example
 - @np.vectorize # Important step. We have to vectorize our functions so it can pass rows one at a time. We write this at the top right before the function.
 - def avg_2_mod(x, y):
 - if (x==20):
 - return np.NaN
 - else:
 - return (x + y) / 2
 - avg_2_mod(df["a"], df["b"]) #if we use number we have to do avg_2_mod(df["a"].values, df["b"].values)
- Example 5
 - def extract population(rate):
 - population = rate.split("/")[1]
 - return population
 - # if we want to return population as an integer we can just return it as int(population)
 - assert extract_population("123/456") == "456" # quick way to check to make sure the function won't fail # random keywords: unit testing
 - tbl3["population"] =
 tbl3["rate"].apply(extract_population)
 # to apply the function to the column and create a new column called "population"

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Splitting into train and test set

```
(MAKE sure X and y are data frames! Not np.ndarray's! It keeps
X_train, etc. as data frames so you can still run X_train.head() after
splitting)
      X = df \mod el
                                      # entire data frame
      y = df_model.tip_percent # tip_percent (dependent /
      target var)
      from sklearn.model_selection import train_test_split
      X train, X test, y train, y test = train test split(
         X, y, test_size=0.2, random_state=seed)
Example 2
      # Getting the number of rows and columns
       r, c = <myDataFrameName1>.shape
      # Creating an array which has indexes of columns
      i_cols = []
      for i in range(0, c-1):
         i_cols.append(i)
      # Y is the target column, X has the rest
      X = \exp DataFrameName1>[:, 0:(c-1)]
      y = \langle myDataFrameName1 \rangle [:, (c-1)]
      del <myDataFrameName1>
      # Validation chunk size
      val size = 0.1
      # Using a common seed in all experiments so that same
       chunk is used for validation
      seed = 0
      # Splitting the data
      from sklearn.model_selection import train_test_split
      X_train, X_test, y_train, y_test = train_test_split(
         X, y, test_size=val_size, random_state=seed)
      del X
      del y
      # All features
      X \text{ all} = []
```

IMPORTANT: Split into X_train, X_test, y_train, y_test like this

```
# List of combinations
comb = []
# Dictionary to store the Mean Absolute Error for all
algorithms
mae = []
#Scoring parameter
from sklearn.metrics import mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all)
              # A list of all the columns along with dummy
vars
```

- Grabbing specific feature values
 - df["Trip_distance"][df["Trip_distance"]<50]
- Duplicating time column, then striping hour field of it, and putting the hour value into a new column

 - df["pickup_hour"] = df["pickup"].apply(lambda x: x.hour)
- Creating a new feature from an existing date time column (datetime64 type):
 - X_train["pickup_day"] = X_train["pickup_datetime"].dt.day
- This puts, the average trip distance of every pickup hour into a new column
 - df["avg_trip_dist_per_pickup_hour"] =

df[["Trip_distance","pickup_hour"]].groupby("pickup_hour").mean()

- List Unique Values In A pandas Column
 - https://chrisalbon.com/python/data_wrangling/ pandas_list_unique_values_in_column/
- select-pandas-dataframe-rows-and-columns-using-iloc-loc-and-ix
 - https://www.shanelynn.ie/select-pandas-dataframe-rows-and-columns-using-iloc-loc-and-ix/
- Groups two features, and provides the average of a third feature, and stores it into a dataframe
 - summary_wdays_avg_duration =
 pd.DataFrame(df.groupby(["day_of_week","pickup_hour"])
 ["Trip_distance"].mean()) (line 1)
 - summary_wdays_avg_duration (line 2)
 - summary_wdays_avg_duration.reset_index(inplace = True)
 (resets the index for the data frame / makes one?)
 - summary_wdays_avg_duration["unit"]=1 (makes a new column with all one's)
- Drop row or drop column
 - train_X = X_train.drop(["Tip_amount", "tip_percent","log_tip_percent"], axis=1) # how to drop multiple columns
 - https://chrisalbon.com/python/data_wrangling/ pandas_dropping_column_and_rows/
 - df = df[df["Tip_amount"] > 0]
 removes all instances in dataframe where Tip_amount <= 0</pre>
 - Calculations with two columns
 - df["tip_percent"] = df["Tip_amount"]/df["Total_amount"] # calculate tip percentage
 - df["tip_percent"] = df["tip_percent"].apply(lambda x: x * 100)
 # multiply by 100 to get the %
 - Python Pandas: Drop columns in DataFrame by label Names or by Index Positions
 - https://thispointer.com/python-pandas-drop-columns-indataframe-by-label-names-or-by-index-positions/
 - Drop Column
 - Example 1
 - <myDataFrameName1>.drop('<myFeatureName1>',
 axis=1, inplace=True)
 - Drop Index
 - See "Drop Column"
- Good To Know's When Plotting
 - Always pay attention to what your objects are returning while looking at the documentation for matplotlib, seaborn, etc.
- Preparing data to plot with matplotlib, etc (note all features that go into the plot arguments have the size of "(some#,)")
 - grouped_df = pd.DataFrame(df.groupby(["pickup_hour", "Airport"])
 ["tip_percent"].aggregate(np.mean).reset_index())
- Quick plotting with Pandas

Example 1 <myDataFrameName1>.<myFeatureName1 categorical>.pl ot(kind="bar") Example 2 <myDataFrameName1>.<myFeatureName1 continuous>.pl ot(kind="hist") Heatmaps https://docs.scipy.org/doc/numpy/reference/generated/ numpy.zeros like.html https://docs.scipy.org/doc/numpy/reference/generated/ numpy.triu_indices.html#numpy.triu_indices Example 1 (shows one heat map) sns.set(style="white") # Generate a large random dataset df model copy = <myDataFrameName1>.copy() # our dataframe # Compute the correlation matrix corr = df model copy.corr() # corr calculation # Generate a mask for the upper triangle. Note, we only want to show the relevant part # of the heat map mask = np.zeros like(corr, dtype=np.bool) mask[np.triu_indices_from(mask)] = True # Generate a custom diverging colormap cmap = sns.diverging_palette(220, 10, as_cmap=True) # Draw the heatmap with the mask and correct aspect ratio sns.heatmap(corr, mask=mask, cmap=cmap, vmax=.3, center=0. square=True, linewidths=.5, cbar_kws={"shrink": .5}) plt.show() Example 2 (shows two different heat maps, one before data skewing, and one after) # Make sure we use the subsample in our correlation $f_{1}(ax1, ax2) = plt.subplots(2, 1, figsize=(24,20))$ # Entire DataFrame corr = <myDataFrameName1>.corr()

- ax1.set_title("Imbalanced Correlation Matrix \n (don"t use for

20, ax=ax1)

sns.heatmap(corr, cmap="coolwarm_r", annot_kws={"size":

```
reference)", fontsize=14)
             sub_sample_corr =
             <myDataFrameName2 withSkewCorrection>.corr()
             sns.heatmap(sub_sample_corr, cmap="coolwarm_r",
             annot kws={"size":20}, ax=ax2)
             ax2.set_title("SubSample Correlation Matrix \n (use for
             reference)", fontsize=14)
             plt.show()
Violin Plots
      Example 1
             # We will visualize all the continuous attributes using Violin
             Plot - a combination of box and density plots
             # Creating a dataframe with only continuous features
             data = <myDataFrameSetForContinuousVarOnly>.iloc[:,
             116:] # Creating a dataframe with only continuous features
             cols=data.columns
             # Plot violin for all attributes in a 7x2 facetgrid
             n cols = 2
             n_rows = 7
             for i in range(n rows):
               fg, ax = plt.subplots(nrows=1,ncols=n_cols,figsize=(12,8))
               for j in range(n_cols):
                  sns.violinplot(y=cols[i*n_cols+j], data=dataset_train,
             ax=ax[j]
      Example 2
             # Single Violin Plot
             sns.violinplot(data=<myDataFrameName1>,
             y="<myFeatureName1 continuous>")
Subplotting (subplots)
```

Pair Plots

Example 1 (sns.pairplot) data = <myDataFrameSetForContinuousVarOnly>.iloc[:, 116:] # Creating a dataframe with only continuous features cols=data.columns Getting the names of all the columns # data corr = data.corr() Calculating Pearson coefficient for all combinations threshold = <myThresholdValue> # Setting the threshold to select only highly correlated attributes. Eq. 0.5. # corr_list = [] List of pairs along with correlation above threshold size = <myIntegerOfFeaturesToBeConsidered> # Eq. 15. # Searching for the highly correlated pairs for i in range(0, size): #for continuous features for j in range(i+1, size): if (data corr.iloc[i,j] >= threshold and data corr.iloc[i,j] < 1) or (data_corr.iloc[i,j] < 0 and data_corr.iloc[i,j] <= threshold): corr_list.append([data_corr.iloc[i,j],i,j]) # stores coefficient and appropriate column indexes # Sorting to show higher ones first s corr list = sorted(corr list, key=lambda x: -abs(x[0])). See key function, https://docs.python.org/3/howto/ sorting.html for v, i, j in s corr list: print("%s and %s = %.2f" % (cols[i],cols[j],v))# Scatter plot of all the highly correlated pairs for v, i, j in s corr list: sns.pairplot(dataset_train, size=6, x_vars=cols[i], y_vars=cols[j]) plt.show Boxplots Example 1 (box plots with hours) f, axes = plt.subplots(ncols=4, figsize=(20,4)) # Negative Correlations with our <myFeatureName1_categorical> (The lower our feature value the more likely it will be a fraud transaction) sns.boxplot(x="<myFeatureName1 categorical>",

y="<myFeatureName2 continuous>",

data=<myDataFrameName1>, palette=colors, ax=axes[0])

axes[0].set_title("<myFeatureName2_continuous> vs
 <myFeatureName1 categorical> Negative Correlation")

-

- sns.boxplot(x="<myFeatureName1_categorical>",
 y="<myFeatureName3_continuous>",
 data=<myDataFrameName1>, palette=colors, ax=axes[1])
- axes[1].set_title("<myFeatureName3_continuous> vs
 <myFeatureName1_categorical> Negative Correlation")

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- sns.boxplot(x="<myFeatureName1_categorical>",
 y="<myFeatureName4_continuous>",
 data=<myDataFrameName1>, palette=colors, ax=axes[2])
- axes[2].set_title("<myFeatureName4_continuous> vs
 <myFeatureName1_categorical> Negative Correlation")

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- sns.boxplot(x="<myFeatureName1_categorical>",
 y="<myFeatureName5_continuous>",
 data=<myDataFrameName1>, palette=colors, ax=axes[3])
- axes[3].set_title("<myFeatureName5_continuous> vs
 <myFeatureName1_categorical> Negative Correlation")

plt.show()

Histograms

- https://towardsdatascience.com/histograms-and-density-plots-in-python-f6bda88f5ac0
- Example 1 # a quick way to generate a histogram within Pandas (plot)

<myDataFrameName1>.<myFeatureName1_continuous>.pl
ot(kind="hist")

- Example 2 # with seaborn, good for continuous variables (distplot)

sns.distplot(<myDataFrameName1>.<myFeatureName1_continuous>) # this plots a histogram plot along with the kernel density shape

sns.distplot(<myDataFrameName1>.<myFeatureName1_co ntinuous>, kde=False) # plots only a histogram

- Scatter Plot (with Linear fits)
 - Example 1 # using seaborn, this shows a scatter plot and applies a linear fit
 - sns.Implot(x="<myFeatureName1_continuous>",
 y="<myFeatureName2_continuous>",
 data=<myDataFrameName1>) # note, sns.Implot returns
 back an entire figure whereas sns.regplot can be used for subplotting/in axes form (still a little confused, but overall I

think sns.regplot is typically used when subplotting)

- Example 2 # using seaborn, this shows the data as a scatter plot and colors the scattered data based on categories in myFeatureName3 (note, there are two linear fits applied to this plot if <myFeatureName3> is binary).
 - sns.lmplot(x="<myFeatureName1_continuous>", y="<myFeatureName2_continuous>", data=<myDataFrameName1>, hue="<myFeatureName3_categorical>")
- Example 3 # using seaborn, this shows the data as a scatter plot and colors the scattered data based on categories in myFeatureName3 (note, there are two linear fits applied to this plot if <myFeatureName3> is binary). The addition of the col argument <myFeatureName4> separates the data into two plots. This categorical var must match the categorical var in <myFeatureName3>
 - sns.Implot(x="<myFeatureName1_continuous>", y="<myFeatureName2_continuous>", data=<myDataFrameName1>, hue="<myFeatureName3_categorical>", col="<myFeatureName4_categorical>")
- Example 4 # similar to above, but including the row argument turns this into a FacetGrid type plot (think of an 8x2 grid, etc). This is a great way to plot using multiple vars. Note <myFeatureName3>, <myFeatureName4>, <myFeatureName5> can be binary, binary, multi categorical, respectively.
 - sns.Implot(x="<myFeatureName1_continuous>", y="<myFeatureName2_continuous>", data=<myDataFrameName1>, hue="<myFeatureName3_categorical>", col="<myFeatureName4_categorical>", row="<myFeatureName5_categorical>")
- Exampe 5
- FacetGrid / Subplotting
 - Scatter Plot
 - Example 1

- <someVar1> =
 sns.FacetGrid(<myDataFrameName1>,
 col="<myFeatureName1_categorical>",
 row="<myFeatureName2_categorical>",
 hue="<myFeatureName3_categorical>")

<someVar1>.map(plt.scatter,
 "<myFeatureName4_continuous>",
 "<myFeatureName5_continuous>") #think of
 "map()" similar to how "apply()" works. Here, map()

provides/applies the data to each subplot in the facet grid

- Histogram and Scatter Plot (with linear fit)
 - Example 1
 - fig, (ax1, ax2) = plt.subplots(1,2)

sns.distplot(<myDataFrameName1>.<myFeatureName1_continuous>, ax=ax1)

sns.regplot(x="<myFeatureName2_continuous>",y="<myFeatureName1_continuous>",
data=<myDataFrameName1>, ax=ax2) # note, this
seems very similar to sns.lmplot

- Bar Graph / Count Plot
 - Example 1 # a quick way to generate a bar plot within Pandas
 - cts =
 <myDataFrameName1>.<myFeatureName1_categorical>.va
 lue_counts
 - cts.plot(kind="bar")
 - Example 2 # with seaborn
 - Example 3 # FacetGrid type with seaborn

-

- n_rows = <myIntOfRows>

- n_cols = <myIntOfColumns>

- for i in range(n_rows):

fg, ax = plt.subplots(nrows=1,ncols=n_cols,figsize=(16,8))

for j in range(n_cols):

sns.countplot(x=cols[i*n_cols+j],
data=<myDataFrameName1_categorical>, ax=ax[j])

-

- Dendogram
 - Example 1

_

- Modeling
 - Regression
 - Linear
 - Linear Regression
 - Example 1

_

- from sklearn.linear_model import LinearRegression
- <myDataFrameName1> =
 sns.load_dataset("tips")
- <myDataFrameName1>.head()
- <myModel1> = LinearRegression()

<myModel1>.fit(X=<myDataFrameNam e1>[["<myFeatureName1>","<myFeatur eName2>"]],

y=<myDataFrameName1>["<myFeature Name3>"])

- <myModel1>.coef_
- <myModel1>.intercept_

_

- # LEARNING:
- # For every one dollar increase in total_bill, given that
- # the size remains the same, the tip increases by 9 cents.

- Example 2

_

- # Getting the number of rows and columns
- $r, c = \mbox{-myDataFrameName1>.shape}$

-

- # Creating an array which has indexes of columns
- i_cols = []

-

- for i in range(0, c-1):
- i_cols.append(i)

-

- # Y is the target column, X has the rest
- X = -myDataFrameName1>[:, 0:(c-1)]
- y = <myDataFrameName1>[:, (c-1)]

-

- # Validation chunk size
- val_size = 0.1

-

- # Using a common seed in all experiments so that same chunk is used for validation
- seed = 0

-

```
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X_all = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
              # A list of all the columns
print(X_all)
along with dummy vars
# LINEAR REGRESSION (Linear Algo)
from sklearn.linear_model import
LinearRegression
lin_reg = LinearRegression(n_jobs=-1) #
using all processors
algo = "LR"
# Accuracy of the model using all
```

from sklearn.model_selection import

```
features
             for name, i_cols_list in X_all:
               print(name)
               lin_reg.fit(X_train[:, i_cols_list],
             y_train) #fitting all features to the target
             column
               result =
             mean_absolute_error(np.expm1(y_test),
             np.expm1(lin_reg.predict(X_test[:,i_cols
             list])))
               mae.append(result)
               print(name + " %s" % result)
             comb.append(algo)
             print(comb)
             # #Plot the MAE of all combinations
             # fig, ax = plt.subplots()
             # plt.plot(mae)
             # #Set the tick names to names of
             combinations
             # ax.set_xticks(range(len(comb)))
             ax.set_xticklabels(comb,rotation='vertica
             # #Plot the accuracy for all combinations
             # plt.show()
Logistic Regression
      Example 1
             from sklearn.linear model import
             LogisticRegression
             <myDataFrameName1> =
             sns.load_dataset("titanic")
             <myDataFrameName1>.head()
             <myDataFrameName1>.info()
             <myDataFrameName1>.drop(["<myFeat
             ureName1>"], axis=1).head()
```

<myDataFrameName2> = <myDataFrameName1>[["<myFeatureN ame2>", "<myFeatureName3>", "<myFeatureName4>"]] <mvDataFrameName2>.head() <myDataFrameName3> = pd.get dummies(<myDataFrameName2 >, drop_first=True) <myDataFrameName3>.head() <myXTrain1> = <myDataFrameName3>.iloc[:,1:] <myXTrain1>.head() <myYTrain1> = <myDataFrameName3>.iloc[:,0] <myYTrain1>.head() <mvModel1> = LogisticRegression(random state=0, solver="lbfgs", multi_class="multinomial") <myModel1>.fit(<myXTrain1>, <mvYTrain1>) <myModel1>.coef Example 1 # Getting the number of rows and columns r, c = <myDataFrameName1>.shape # Creating an array which has indexes of columns i_cols = [] for i in range(0, c-1): i_cols.append(i) # Y is the target column, X has the rest $X = \exp DataFrameName1>[:, 0:(c-1)]$ y = <myDataFrameName1>[:, (c-1)] # Validation chunk size val size = 0.1# Using a common seed in all

Non-Linear

KNN

experiments so that same chunk is used

```
for validation
seed = 0
from sklearn.model_selection import
train test split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X \text{ all} = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all)
              # A list of all the columns
along with dummy vars
# KNN (Non-linear Algo)
# Evaluation of various combinations of
KNN
# Fitting Classifier to the Training set
from sklearn.neighbors import
KNeighborsRegressor
# https://scikit-learn.org/stable/modules/
```

```
generated/
sklearn.neighbors.KNeighborsRegressor
.html
# Add the N value to the below list if you
want to run the algo
n_list = np.array([]) #note, when the list
is empty, the algo doesnt run
With n_{list} = np.array([5])
All 1434.788795356784
['LR', 'KNN 5']
With n_{int} = np.array([2])
All 1526.7442802258656
['LR', 'KNN 2']
# we can use multiple values into n list
if we want to search for the optimal
n_neighbors. However, xgboost is usally
the best for parameter tuning.
for n neighbors in n list:
  # Setting the base model
  regressor =
KNeighborsRegressor(n_neighbors=n_n
eighbors,n_jobs=-1)
  algo = "KNN"
  #Accuracy of the model using all
features
  for name, i cols list in X all:
     regressor.fit(X_train[:, i_cols_list],
y_train) #fitting all features to the target
column
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(regressor.predict(X_test[:,i_c
ols_list])))
     mae.append(result)
```

```
print(name + " %s" % result)
               comb.append(algo + " %s" %
             n neighbors)
             print(comb)
             # since we know the outcome, we can
             skip the algorithm and append the result
             if (len(n_list)==0):
               mae.append(1527)
               comb.append("KNN" + " %s" % 2)
             ##Set figure size, this figure compares
             mae for all of the algorithms ran
             #plt.rc("figure", figsize=(25, 10))
             ##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set_xticks(range(len(comb)))
             #ax.set_xticklabels(comb,rotation='vertic
             ##Plot the accuracy for all combinations
             #plt.show()
             #Very high computation time
             #Best estimated performance is 1745
             for n=1
             # LEARNING:
             # KNN 5 performed the best. Lowest
             MAE.
Decision Tree (CART)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
```

```
# Creating an array which has indexes
of columns
i_cols = []
for i in range(0, c-1):
  i_cols.append(i)
# Y is the target column, X has the rest
X = \exp DataFrameName1>[:, 0:(c-1)]
y = <myDataFrameName1>[:, (c-1)]
# Validation chunk size
val_size = 0.1
# Using a common seed in all
experiments so that same chunk is used
for validation
seed = 0
from sklearn.model selection import
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test size=val size,
random_state=seed)
del X
del y
# All features
X_{all} = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
```

```
X_all.append([n, i_cols])
print(X_all) # A list of all the columns
along with dummy vars
# CART (Non-linear Algo)
#Evaluation of various combinations of
CART
#Import the library
from sklearn.tree import
DecisionTreeRegressor
#Add the max_depth value to the below
list if you want to run the algo
d list = np.array([])
for max_depth in d_list:
  #Set the base model
  model =
DecisionTreeRegressor(max_depth=ma
x_depth,random_state=seed)
  algo = "CART"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
    result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X_test[:,i_cols_
list])))
    mae.append(result)
    print(name + " %s" % result)
  comb.append(algo + " %s" %
max_depth)
# since we know the outcome, we can
```

```
skip the algorithm and append the result
if (len(d_list)==0):
  mae.append(1741)
  comb.append("CART" + " %s" % 5)
##Set figure size
#plt.rc("figure", figsize=(25, 10))
##Plot the MAE of all combinations
#fig, ax = plt.subplots()
#plt.plot(mae)
##Set the tick names to names of
combinations
#ax.set_xticks(range(len(comb)))
#ax.set_xticklabels(comb,rotation='vertic
al')
##Plot the accuracy for all combinations
#plt.show()
#High computation time
#Best estimated performance is 1741
for depth=5
```

Support Vector Machine (SVM, SVR)

```
Example 1
       # https://medium.com/
      @pushkarmandot/what-is-the-
      significance-of-c-value-in-support-
      vector-machine-28224e852c5a
      # Getting the number of rows and
      columns
       r, c = <myDataFrameName1>.shape
      # Creating an array which has indexes
      of columns
      i_cols = []
      for i in range(0, c-1):
         i_cols.append(i)
      # Y is the target column, X has the rest
      X = \exp DataFrameName1>[:, 0:(c-1)]
      y = <myDataFrameName1>[:, (c-1)]
      # Validation chunk size
      val\_size = 0.1
      # Using a common seed in all
      experiments so that same chunk is used
      for validation
      seed = 0
      from sklearn.model selection import
      train_test_split
      X_train, X_test, y_train, y_test =
      train_test_split(
         X, y, test_size=val_size,
      random_state=seed)
      del X
      del y
      # All features
      X_{all} = []
      # List of combinations
      comb = []
```

```
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all) # A list of all the columns
along with dummy vars
# SVM (Non-linear Algo)
# https://medium.com/
@pushkarmandot/what-is-the-
significance-of-c-value-in-support-
vector-machine-28224e852c5a
#Import the library
from sklearn.svm import SVR
#Add the C value to the below list if you
want to run the algo
c_{list} = np.array([])
for C in c list:
  #Set the base model
  model = SVR(C=C)
  algo = "SVM"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
```

```
list])))
                  mae.append(result)
                  print(name + " %s" % result)
               comb.append(algo + " %s" % C)
             ##Set figure size
             #plt.rc("figure", figsize=(25, 10))
             ##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set_xticks(range(len(comb)))
             #ax.set xticklabels(comb,rotation='vertic
             al')
             ##Plot the accuracy for all combinations
             #plt.show()
             #very very high computation time, not
             running
Bagged Decision Trees (Bagging)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
             # Creating an array which has indexes
             of columns
             i_cols = []
             for i in range(0, c-1):
                i_cols.append(i)
             # Y is the target column, X has the rest
```

np.expm1(model.predict(X_test[:,i_cols_

```
X = \exp DataFrameName1>[:, 0:(c-1)]
y = <myDataFrameName1>[:, (c-1)]
# Validation chunk size
val size = 0.1
# Using a common seed in all
experiments so that same chunk is used
for validation
seed = 0
from sklearn.model_selection import
train_test_split
X_train, X_test, y_train, y_test =
train test split(
   X, y, test_size=val_size,
random state=seed)
del X
del y
# All features
X_all = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all) # A list of all the columns
along with dummy vars
```

```
# Bagged Decision Trees (Bagging)
#Evaluation of various combinations of
Bagged Decision Trees
#Import the library
from sklearn.ensemble import
BaggingRegressor
#from sklearn.tree import
DecisionTreeRegressor
#Add the n_estimators value to the
below list if you want to run the algo
n_{int} = np.array([])
for n_estimators in n_list:
  #Setting the base model
  model =
BaggingRegressor(n_jobs=-1,n_estimat
ors=n_estimators)
  algo = "Bag"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X_test[:,i_cols_
list])))
     mae.append(result)
     print(name + " %s" % result)
  comb.append(algo + " %s" %
n_estimators)
##Set figure size
#plt.rc("figure", figsize=(25, 10))
```

```
##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set_xticks(range(len(comb)))
             #ax.set_xticklabels(comb,rotation='vertic
             ##Plot the accuracy for all combinations
             #plt.show()
             #very high computation time, not
             running
Random Forest (Bagging) # note, we use bagging to
find the sweet spot between a simple and complex
model (see bias vs variance tradeoff)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
             # Creating an array which has indexes
             of columns
             i_cols = []
             for i in range(0, c-1):
               i cols.append(i)
             # Y is the target column, X has the rest
             X = \exp DataFrameName1>[:, 0:(c-1)]
             y = <myDataFrameName1>[:, (c-1)]
```

_

- # Validation chunk size

val_size = 0.1

_

 # Using a common seed in all experiments so that same chunk is used for validation

- seed = 0

-

from sklearn.model_selection import

```
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X \text{ all} = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{``All''}
X_all.append([n, i_cols])
print(X_all) # A list of all the columns
along with dummy vars
# Random Forest (Bagging)
# Evaluation of various combinations of
RandomForest
#Import the library
from sklearn.ensemble import
RandomForestRegressor
#Add the n estimators value to the
below list if you want to run the algo
```

```
n_{int} = np.array([])
for n estimators in n list:
  #Set the base model
  model =
RandomForestRegressor(n_jobs=-1,n_e
stimators=n estimators,random state=s
eed)
  algo = "RF"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X_testl[:,i_cols
_list])))
     mae.append(result)
     print(name + " %s" % result)
  comb.append(algo + " %s" %
n estimators)
# since we know the outcome, we can
skip the algorithm and append the result
if (len(n_list)==0):
  mae.append(1213)
  comb.append("RF" + " %s" % 50 )
##Set figure size
#plt.rc("figure", figsize=(25, 10))
##Plot the MAE of all combinations
#fig, ax = plt.subplots()
#plt.plot(mae)
##Set the tick names to names of
combinations
#ax.set_xticks(range(len(comb)))
#ax.set_xticklabels(comb,rotation='vertic
al')
##Plot the accuracy for all combinations
#plt.show()
```

#Best estimated performance is 1213
when the number of estimators is 50

 Extra Trees (Bagging) # note, we use bagging to find the sweet spot between a simple and complex model (see bias vs variance tradeoff)

Example 1

_

- # Getting the number of rows and columns

r, c = <myDataFrameName1>.shape

-

Creating an array which has indexes of columns

- i_cols = []

-

- for i in range(0, c-1):

i_cols.append(i)

-

- # Y is the target column, X has the rest

- $X = \mbox{myDataFrameName1>[:, 0:(c-1)]}$

- $y = \mbox{myDataFrameName1>}[:, (c-1)]$

-

Validation chunk size

val_size = 0.1

_

 # Using a common seed in all experiments so that same chunk is used for validation

 \cdot seed = 0

-

 from sklearn.model_selection import train test split

- X_train, X_test, y_train, y_test = train_test_split(

 X, y, test_size=val_size, random_state=seed)

-

- del X

- del y

-

All features

X_all = []

_

```
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{``All''}
X_all.append([n, i_cols])
print(X_all)
            # A list of all the columns
along with dummy vars
# Extra Trees (Bagging)
#Evaluation of various combinations of
ExtraTrees
#Import the library
from sklearn.ensemble import
ExtraTreesRegressor
#Add the n_estimators value to the
below list if you want to run the algo
n_{int} = np.array([])
for n estimators in n list:
  #Set the base model
  model =
ExtraTreesRegressor(n_jobs=-1,n_estim
ators=n_estimators,random_state=seed
)
  algo = "ET"
```

List of combinations

```
#Accuracy of the model using all
             features
               for name,i_cols_list in X_all:
             model.fit(X_train[:,i_cols_list],Y_train)
                  result =
             mean absolute error(np.expm1(y test),
             np.expm1(model.predict(X_test[:,i_cols_
             list])))
                  mae.append(result)
                  print(name + " %s" % result)
               comb.append(algo + " %s" %
             n_estimators)
             # since we know the outcome, we can
             skip the algorithm and append the result
             if (len(n_list)==0):
               mae.append(1254)
               comb.append("ET" + " %s" % 100 )
             ##Set figure size
             #plt.rc("figure", figsize=(25, 10))
             ##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set_xticks(range(len(comb)))
             #ax.set xticklabels(comb,rotation='vertic
             ##Plot the accuracy for all combinations
             #plt.show()
             #Best estimated performance is 1254
             for 100 estimators
Adaboost # note, we use bagging to find the sweet
```

spot between a simple and complex model (see bias vs variance tradeoff)

Example 1

```
# Getting the number of rows and
columns
r, c = <myDataFrameName1>.shape
# Creating an array which has indexes
of columns
i_cols = []
for i in range(0, c-1):
  i_cols.append(i)
# Y is the target column, X has the rest
X = \exp DataFrameName1>[:, 0:(c-1)]
y = \langle myDataFrameName1 \rangle [:, (c-1)]
# Validation chunk size
val size = 0.1
# Using a common seed in all
experiments so that same chunk is used
for validation
seed = 0
from sklearn.model_selection import
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X_all = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
```

- from sklearn.metrics import

#Scoring parameter

```
mean_absolute_error
#Add this version of X to the list
n = \text{``All''}
X_all.append([n, i_cols])
print(X_all)
             # A list of all the columns
along with dummy vars
#Evaluation of various combinations of
AdaBoost
#Import the library
from sklearn.ensemble import
AdaBoostRegressor
#Add the n estimators value to the
below list if you want to run the algo
n_{int} = np.array([])
for n estimators in n list:
  #Set the base model
  model =
AdaBoostRegressor(n_estimators=n_es
timators,random state=seed)
  algo = "Ada"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X_test[:,i_cols_
list])))
     mae.append(result)
    print(name + " %s" % result)
  comb.append(algo + " %s" %
n estimators)
```

-

```
# since we know the outcome, we can
             skip the algorithm and append the result
             if (len(n_list)==0):
               mae.append(1678)
               comb.append("Ada" + " %s" % 100 )
             ##Set figure size
             #plt.rc("figure", figsize=(25, 10))
             ##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set_xticks(range(len(comb)))
             #ax.set_xticklabels(comb,rotation='vertic
             al')
             ##Plot the accuracy for all combinations
             #plt.show()
             #Best estimated performance is 1678
             with n=100
Stochastic Gradient Boosting (Boosting) # note, we
use bagging to find the sweet spot between a simple
and complex model (see bias vs variance tradeoff)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
             # Creating an array which has indexes
             of columns
             i_cols = []
             for i in range(0, c-1):
               i_cols.append(i)
             # Y is the target column, X has the rest
             X = \exp DataFrameName1>[:, 0:(c-1)]
             y = \langle myDataFrameName1 \rangle [:, (c-1)]
```

Validation chunk size

```
val_size = 0.1
# Using a common seed in all
experiments so that same chunk is used
for validation
seed = 0
from sklearn.model_selection import
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X_all = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all) # A list of all the columns
along with dummy vars
#Evaluation of various combinations of
SGB
```

-

#Import the library

- from sklearn.ensemble import GradientBoostingRegressor

```
#Add the n_estimators value to the
below list if you want to run the algo
n_list = np.array([])
for n_estimators in n_list:
  #Set the base model
  model =
GradientBoostingRegressor(n_estimator
s=n_estimators,random_state=seed)
  algo = "SGB"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X testl[:,i cols
list])))
     mae.append(result)
     print(name + " %s" % result)
  comb.append(algo + " %s" %
n estimators)
# since we know the outcome, we can
skip the algorithm and append the result
if (len(n_list)==0):
  mae.append(1278)
  comb.append("SGB" + " %s" % 50 )
##Set figure size
#plt.rc("figure", figsize=(25, 10))
##Plot the MAE of all combinations
#fig, ax = plt.subplots()
#plt.plot(mae)
##Set the tick names to names of
combinations
#ax.set_xticks(range(len(comb)))
#ax.set_xticklabels(comb,rotation='vertic
al')
##Plot the accuracy for all combinations
```

```
#plt.show()
             #Best estimated performance is?
XGBoost # note, we use bagging to find the sweet
spot between a simple and complex model (see bias
vs variance tradeoff)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
             # Creating an array which has indexes
             of columns
             i cols = []
             for i in range(0, c-1):
               i_cols.append(i)
             # Y is the target column, X has the rest
             X = \exp DataFrameName1>[:, 0:(c-1)]
             y = <myDataFrameName1>[:, (c-1)]
             # Validation chunk size
             val_size = 0.1
             # Using a common seed in all
             experiments so that same chunk is used
             for validation
             seed = 0
             from sklearn.model_selection import
             train_test_split
             X_train, X_test, y_train, y_test =
             train_test_split(
                X, y, test_size=val_size,
             random state=seed)
             del X
```

- del y

-

All features

 \cdot X_all = []

-

List of combinations

```
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
              # A list of all the columns
print(X_all)
along with dummy vars
#XGBoost
#Evaluation of various combinations of
XGB
#Import the library
from xgboost import XGBRegressor
#Add the n_estimators value to the
below list if you want to run the algo
n_{int} = np.array([])
for n_estimators in n_list:
  #Set the base model
  model =
XGBRegressor(n_estimators=n_estimat
ors, seed = seed)
  algo = "XGB"
  #Accuracy of the model using all
features
  for name,i_cols_list in X_all:
model.fit(X_train[:,i_cols_list],Y_train)
     result =
```

```
mean_absolute_error(np.expm1(y_test),
             np.expm1(model.predict(X_test[:,i_cols_
             list])))
                  mae.append(result)
                  print(name + " %s" % result)
               comb.append(algo + " %s" %
             n_estimators)
             # since we know the outcome, we can
             skip the algorithm and append the result
             if (len(n_list)==0):
               mae.append(1169)
               comb.append("XGB" + " %s" %
             1000)
             ##Set figure size
             #plt.rc("figure", figsize=(25, 10))
             ##Plot the MAE of all combinations
             #fig, ax = plt.subplots()
             #plt.plot(mae)
             ##Set the tick names to names of
             combinations
             #ax.set xticks(range(len(comb)))
             #ax.set_xticklabels(comb,rotation='vertic
             al')
             ##Plot the accuracy for all combinations
             #plt.show()
             #Best estimated performance is 1169
             with n=1000
Multi-layer Perceptrons (Deep Learning)
      Example 1
             # Getting the number of rows and
             columns
             r, c = <myDataFrameName1>.shape
             # Creating an array which has indexes
             of columns
             i_cols = []
             for i in range(0, c-1):
```

```
i_cols.append(i)
# Y is the target column, X has the rest
X = \exp DataFrameName1>[:, 0:(c-1)]
y = <myDataFrameName1>[:, (c-1)]
# Validation chunk size
val_size = 0.1
# Using a common seed in all
experiments so that same chunk is used
for validation
seed = 0
from sklearn.model_selection import
train_test_split
X_train, X_test, y_train, y_test =
train_test_split(
   X, y, test_size=val_size,
random_state=seed)
del X
del y
# All features
X_all = []
# List of combinations
comb = []
# Dictionary to store the Mean Absolute
Error for all algorithms
mae = []
#Scoring parameter
from sklearn.metrics import
mean_absolute_error
#Add this version of X to the list
n = \text{"All"}
X_all.append([n, i_cols])
print(X_all)
              # A list of all the columns
along with dummy vars
```

-

```
#MLP (Deep Learning)
#Evaluation of various combinations of
multi-layer perceptrons
#Import libraries for deep learning
from keras.wrappers.scikit_learn import
KerasRegressor
from keras.models import Sequential
from keras.layers import Dense
# define baseline model
def baseline(v):
   # create model
   model = Sequential()
   model.add(Dense(v*(c-1),
input_dim=v*(c-1), init='normal',
activation='relu'))
   model.add(Dense(1, init='normal'))
   # Compile model
model.compile(loss='mean absolute err
or', optimizer='adam')
   return model
# define smaller model
def smaller(v):
   # create model
   model = Sequential()
   model.add(Dense(v*(c-1)/2,
input_dim=v*(c-1), init='normal',
activation='relu'))
   model.add(Dense(1, init='normal',
activation='relu'))
   # Compile model
model.compile(loss='mean_absolute_err
or', optimizer='adam')
   return model
# define deeper model
def deeper(v):
# create model
model = Sequential()
```

```
model.add(Dense(v*(c-1),
input_dim=v*(c-1), init='normal',
activation='relu'))
model.add(Dense(v*(c-1)/2,
init='normal', activation='relu'))
model.add(Dense(1, init='normal',
activation='relu'))
# Compile model
model.compile(loss='mean_absolute_err
or', optimizer='adam')
return model
# Optimize using dropout and decay
from keras.optimizers import SGD
from keras.layers import Dropout
from keras.constraints import maxnorm
def dropout(v):
  #create model
  model = Sequential()
  model.add(Dense(v*(c-1),
input_dim=v*(c-1), init='normal',
activation='relu',W_constraint=maxnorm
(3)))
  model.add(Dropout(0.2))
  model.add(Dense(v*(c-1)/2,
init='normal', activation='relu',
W constraint=maxnorm(3)))
  model.add(Dropout(0.2))
  model.add(Dense(1, init='normal',
activation='relu'))
  # Compile model
  sqd =
SGD(lr=0.1,momentum=0.9,decay=0.0,
nesterov=False)
model.compile(loss='mean_absolute_err
or', optimizer=sqd)
  return model
# define decay model
def decay(v):
  # create model
  model = Sequential()
  model.add(Dense(v*(c-1),
input_dim=v*(c-1), init='normal',
```

```
activation='relu'))
  model.add(Dense(1, init='normal',
activation='relu'))
  # Compile model
  sad =
SGD(lr=0.1,momentum=0.8,decay=0.01
,nesterov=False)
model.compile(loss='mean_absolute_err
or', optimizer=sgd)
  return model
est_list = []
#uncomment the below if you want to
run the algo
#est_list = [('MLP',baseline),
('smaller', smaller), ('deeper', deeper),
('dropout',dropout),('decay',decay)]
for name, est in est list:
  algo = name
  #Accuracy of the model using all
features
  for m,i_cols_list in X_all:
     model =
KerasRegressor(build fn=est, v=1,
nb_epoch=10, verbose=0)
model.fit(X_train[:,i_cols_list],Y_train)
     result =
mean_absolute_error(np.expm1(y_test),
np.expm1(model.predict(X test[:,i cols
list])))
     mae.append(result)
    print(name + " %s" % result)
  comb.append(algo)
# since we know the outcome, we can
skip the algorithm and append the result
if (len(est_list)==0):
  mae.append(1168)
  comb.append("MLP" + " baseline" )
```

print("mae--> %s" % mae) print("comb--> %s" % comb) ##Set figure size plt.rc("figure", figsize=(25, 10)) #Plot the MAE of all combinations fig, ax = plt.subplots()

plt.plot(mae)

#Set the tick names to names of combinations

ax.set_xticks(range(len(comb)))

ax.set_xticklabels(comb,rotation='vertica

#Plot the accuracy for all combinations

plt.show()

#Best estimated performance is MLP=1168

- Classification
- Model Selection
 - Example 1

Since XGBRegressor is showing the best performance, we can select it as our best model. Therefore, we now need to finalize the model with all of the available data.

- # note, X_train and X_test are both coming from the training set CSV. axis=0 is stacking rows on top of one another.
- $X = np.concatenate((X_train, X_test), axis=0)$
- del X_train
- del X test
- Y = np.concatenate((y_train,y_test),axis=0)
- del y_train
- del y_test

```
print("I am here 0 - debug")
n_estimators = 1000
#Best model definition
best model =
XGBRegressor(n_estimators=n_estimators,seed=see
print("I am here 0.0 - debug")
best model.fit(X,Y)
print("I am here 0.1 - debug")
del X
del Y
#Read test dataset
dataset_test = pd.read_csv("test.csv")
print("I am here 0.2 - debug")
#Drop unnecessary columns
ID = dataset test['id']
dataset_test.drop('id',axis=1,inplace=True)
#One hot encode all categorical attributes
cats = []
print("I am here 1 - debug")
for i in range(0, split):
  # label encoding
  label encoder = LabelEncoder()
  label_encoder.fit(labels[i])
  feature =
label_encoder.transform(dataset_test.iloc[:,i])
  feature = feature.reshape(dataset_test.shape[0], 1)
  #One hot encoding
  onehot encoder =
OneHotEncoder(sparse=False,n_values=len(labels[i])
  feature = onehot encoder.fit transform(feature)
  cats.append(feature)
print("I am here 2 - debug")
# Making a 2D array from a list of 1D arrays
encoded_cats = np.column_stack(cats)
del cats
# Concatenating encoded attributes with continous
attributes
X test = np.concatenate((encoded cats,
dataset_test.iloc[:,split:].values), axis=1)
```

print("I am here 3 - debug") del encoded_cats del dataset_test # Making predictions using the best model now predictions = np.expm1(best_model.predict(X_test))

DATABASE

- Good Reads
 - Data Redundancy and Data Inconsistency
 - https://pediaa.com/what-is-the-difference-between-data-redundancy-and-data-inconsistency/
 - Data redundancy can lead to data inconsistency

SAS/SPSS

- Fantastic YouTube Channels
 - Research By Design

- Statistical Significance vs. Effect Size (https://www.youtube.com/playlist?list=PLVI_iGT5ZuRlCryAkbFeRiutKec8ck4Qk) # Playlist
- One Way Anova Interpretation
 - How to do a One-Way ANOVA in SPSS (12-6)
 - https://www.youtube.com/watch?v=rS3k8ONVN-o
 - Example
 - Following a series of complaints about wicked witches, the Wizard of Oz conducts a study to determine if certain regions of Oz have more problems with wicked witches than other regions. He randomly surveys five Munchkins from each of four regions and records the number of complaints he received about wicked witchiness from each one. For example, each Munchkin may say he/she received anywhere from 1-5 complaints
 - Before we start interpreting our ANOVA results, we should first check the assumptions for the test. One of which is
 - Homogeneity of Variances
 - Which is tested by performing a Levene Test (see my other video https://www.youtube.com/watch? v=4mkEZxgxMRA)
 - Since our results show the the the Significance of the Levene Test is 0.918 this means our group variances are homogenous (not related, this is great)
 - This means we will not need the Welch ANOVA test or the Games Howell either.
 - Now we can proceed to the ANOVA / F Table
 - Note, the outputted table is not in APA format. Never use raw SPSS output in place of APA formatted tables.
 - Since we used the compare means and not the general linear model option
 - To interpret and report this ANOVA we would write (see ~07:00)
 - $F(3,16) = 10.49, p < 0.001, \eta^2_p$ = 0.66
 - note η^2_p is the "effect size" (see my other video for more info https://www.youtube.com/watch?

Standard Error (note, the std error can also be thought of as the standard deviation of a bunch of sampled means from a given population (see StatQuickie: Standard Deviation vs Standard Error by Josh Starmer (https://www.youtube.com/watch? v=A82brFpdr9g&list=PLblh5JKOoLUK0FLuzwntyYI10UQFUhsY9&index=13)))

- Standard Error by Bozeman Science https://www.youtube.com/ watch?v=BwYj69LAQOI
 - stdError = stdDev / sqrt(n)
 - where StdDev and n is the number of observations/ measurements
 - Note, the more observations we have, the lower the standard error will become
- Standard Deviation by Bozeman Science https://www.youtube.com/watch?v=09kiX3p5Vek
 - stdDev = sqrt(summation((x x_mean)^2) / (n -1)) # Note this is for sample variance, not population variance
 - where x is each sample in data set, x_mean is the average of the samples, n is the number of samples in our data
 - Why we normally divide by n-1 https:// www.khanacademy.org/math/ap-statistics/summarizingquantitative-data-ap/more-standard-deviation/v/anothersimulation-giving-evidence-that-n-1-gives-us-an-unbiasedestimate-of-variance
 - Variance and Std Deviation I Why divide by n-1 by zedstatistics https://www.youtube.com/watch? v=wpY9o_OyxoQ (great video)
 - Example
 - Objective: Describe the spread of the data
 - Out intuition tells us to just subtract the highest and lowest observations to get the range
 - However, this can be susceptible to outliers. Therefore, we need a way to incorporate the rest of the observations.
 - First thought: We can solve for the mean and the look at the average deviation from it!
 - However, this will give us negative deviations for the left side of the mean while the right side will give positive.
 - Note, by definition, the sum of the deviations to the left and to the

right will be <u>zero</u>. Hence, why we square the residuals from the mean.

- Second though: Let's find the average squared deviation from the mean
 - Note, there is a topic called "moments" that addresses why it is better to square the differences instead of using the absolution value ~6:00
 - Also, we would think to get the average squared deviation we will need to divide by just n (not n - some#)
 - So why is it so common to divide by n-1?
 - See video around ~6:00 for explanation
- Std deviation measures the variance in the data
- Note, variance = (standard deviation)^2