Elements Of Data Science - F2020

Week 4: Hypothesis Testing

10/5/2020

Tools So Far (a common first cell)

```
In [1]: import numpy as np
    import pandas as pd
    import matplotlib.pyplot as plt
    import seaborn as sns

sns.set_style('darkgrid')
%matplotlib inline
```

TODOs

- Readings
 - PDSH Chap 5 up to but not including Hyperparameters and Model Evaluation
 - PDSH Chap 5 In Depth: Linear Regression
 - PDSH Chap 5 In Depth: Support Vector Machines
 - PDSH Chap 5 In Depth: Decision Trees and Random Forests
 - Recommended PML Chap 3
 - Optional PML Chap 2
- Quiz 4
- HW1, out next few days, due 2 weeks from release via gradescope
- Midterm: Last week of October, take-home, open-book

Additional Resources for Hypothesis Testing

- Statistical Rules of Thumb, Gerald van Belle Chapter 2 online
- "The 2019 ASA Guide to P-values and Statistical Significance: Don't Say What You Don't Mean" (Some Recommendations)(ii)

Today

Hypothesis Testing and MAB

Questions?

Hypothesis Testing

- Random Sampling
- Confidence Intervals
- A/B Tests
- Hypothesis Testing
- Permutation Tests
- p-values
- Calculating Power and Number of Observations Needed
- Multi-Armed Bandit

Questions and More Questions

- Have web conversions gone up?
- Have stock prices changed?
- Which ad generates more sales?
- Which headline generates more clicks?
- Did the number of "likes" change?

Scipy

• Routines for numerical integration, interpolation, optimization, linear algebra, and statistics.



In [2]: import scipy as sp

Mini Probability Review

- Random Variable
 - takes values from an associated probability distribution
- Distribution
 - describes probability of values of a Random Variable
- P(x): Probability
 - probability of seeing x, takes value in [0,1]
 - Ex: Probability of getting heads on a coin toss
- $P(x \mid y)$: Conditional Probability
 - probability of seeing x, given that some y is true
 - Ex: Probability of getting heads on a coin toss given that coin is fair

Population Distributions and Sampling

- The World :: Ground Truth
 - Ex: The length of taxi rides
- Our Data :: An Experiment
 - Ex: The length of taxi rides we saw in Jan 2017

Population Dists. and Sampling

- Population Distribution: The actual distribution out in world
 - Ex: Actual distribution of taxi trip length
- Random Sample: Our observations of the true population distrution
 - We hope this does not differ systematically from the true distribution
 - Ex: The taxi trip lengths recorded in Jan 2017
- Sample Size (n): The number of observations, the larger the better
 - Ex: We saw 1,000 trips

Population Dists and Sampling

- Population Mean vs. Sample Mean: μ vs. \bar{x}
 - Ex: The true mean trip length vs the one we observed
- Population Std. Dev. vs Sample Std. Dev.: σ vs. s
 - Ex: The true spread of trip length vs the one we observed
- Sample Statistic: eg. mean, median, standard deviation
 - Ex: We're interested in mean trip length
- Sampling Distribution: Distribution of the sample statistic
 - Ex: How is mean trip length distributed if we were to repeat our experiment many times?

Things To Know First

- sample size
- shape (skewed?, multimodal?)
- location (central tendencies)
- spread

What can we say about the length of an average taxi trip in Jan 2017?

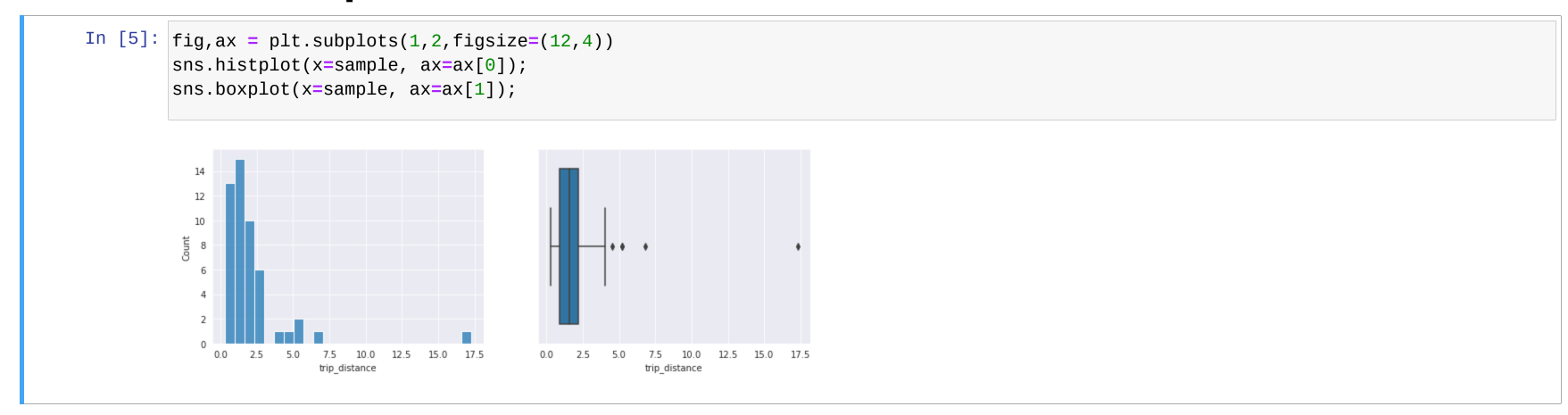
```
In [3]: df = pd.read_csv('../data/yellowcab_demo.csv',parse_dates=['pickup_datetime','dropoff_datetime'])
print(df.shape)

(1000, 6)
```

Sampling From the Population

```
In [4]: sample = df.trip_distance.sample(n=50,
                                                           # our sample size
                                         random_state=123, # needed for reproducability
                                                           # sample without replacement
                                         replace=False
        sample.describe()
Out[4]: count
                 50.000000
                  2.141800
        mean
                  2.563404
        std
                  0.300000
        min
                  0.910000
        25%
        50%
                  1.600000
        75%
                  2.187500
                 17.300000
        max
        Name: trip_distance, dtype: float64
```

Plot our Sample



Define the Sample Statistic

```
In [6]: xbar = sample.mean()
print(f'sample mean: {xbar:0.2f}')
sample mean: 2.14
```

- How good of an approximation is our sample statistic?
- Let's take more samples!

Generating Samples

```
In [7]: sample_means = []
        for i in range(1000):
            sample_mean = df.trip_distance.sample(n=50, random_state=i).mean()
            sample_means.append(sample_mean)
In [8]: # sampling distribution with original statistic
        ax = sns.histplot(x=sample_means)
        ax.set_xlabel('sample_means');
        ax.set_ylabel('frequency');
        ax.axvline(xbar,color='red');
                2.0 2.5
                        sample means
```

Central Limit Theorem

If all samples are randomly drawn from the same sample population:

For reasonably large samples (usually $n \ge 30$), the distribution of sample mean \bar{x} is normal regardless of the distribution of X.

The sampling distribution of \bar{x} becomes approximately normal as the the sample size n gets large.

Ex: X = trip_distance, \bar{x} = mean trip_distance, n = 50

What is Normal?

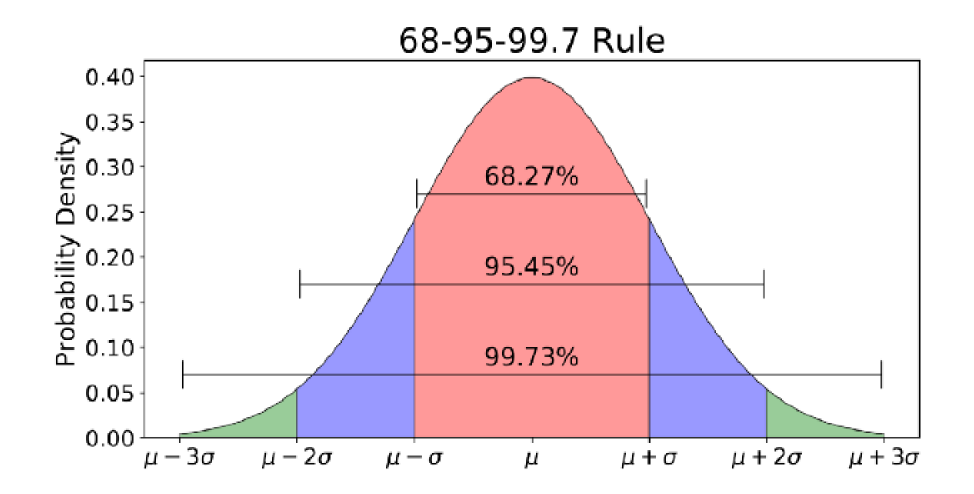
distribution defined by mean (μ) and standard deviation (σ)

$$N(x; \mu, \sigma) = \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{1}{2\sigma}(x-\mu)^2}$$

PDF (Probability Density Function):

• function of a continuous random variable that provides a relative likelihood of seeing a particular sample of a random variable.

Properties of a Normal Distribution



https://towardsdatascience.com/understanding-the-68-95-99-7-rule-for-a-normal-distribution-b7b7cbf760c2

Plotting a Standard Normal Distribution

- Standard Normal: μ =0, σ =1
- ullet Often referred to as Z

```
In [9]: x = np.random.normal(0, 1, size=100_000)
                                                                       # generate many random samples
        fig, ax = plt.subplots(1, 1, figsize=(12, 6))
        ax = sns.histplot(x=x, stat='density', kde=True);
                                                                      # using density to normalize bin counts
        ax.set_xlabel('$x$');ax.set_ylabel('$N(x;\mu=0,\sigma=1)$'); # using latex in labels
        ax.vlines([-1,1],0,sp.stats.norm.pdf(1), colors='k'); # 1 standard deviation
        ax.vlines([-2,2],0,sp.stats.norm.pdf(2), colors='r');
                                                                      # 2 standard deviations
          0.40
          0.35
           0.30
          0.15
          0.10
          0.05
           0.00
```

Confidence Intervals

Typically we only have one sample from the population (set of experimental results, set of survey results, etc.)

```
In [10]: # treat all observations as our sample
    n = len(df.trip_distance)
    n
Out[10]: 1000

In [11]: x_bar = df.trip_distance.mean()
    print(f'sample mean: {x_bar:0.4f}')
    sample mean: 2.8800
```

- What is the spread of our sample statistic?
- What other values would it be reasonable to observe?

Plotting Confidence Intervals with Seaborn

```
In [12]: fig, ax = plt.subplots(1,1,figsize=(12,6))
         sns.barplot(x=df.trip_distance,
                      estimator=np.mean, # default
                      ci=95,
                                            # 95% CI
                      color='c',
                     );
                                               2.0
                                                         2.5
                                       trip distance
```

- How are these confidence intervals generated?
- What does a 95% confidence interval mean?

Generate Confidence Intervals

Bootstrapping: sampling with replacement

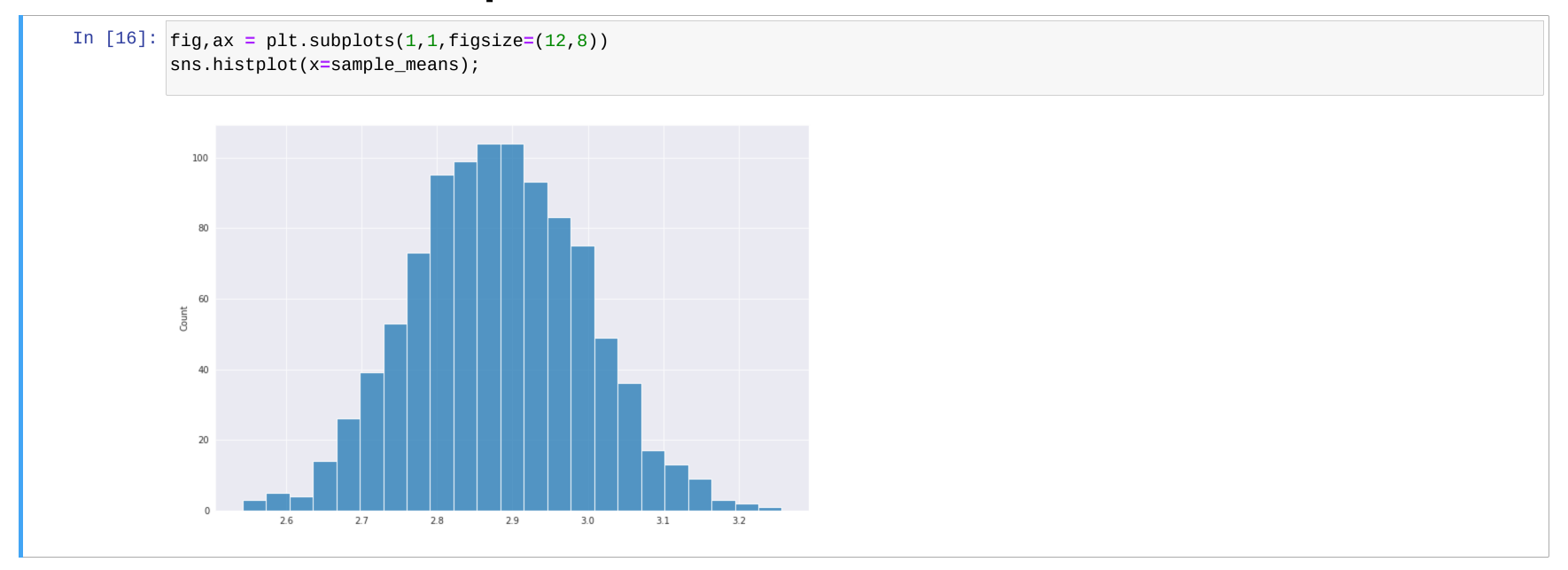
Bootstrap Confidence Interval: create confidence interval using bootstrap samples

- 1. draw a random sample of size n from the data
- 2. record the sample statistic from this random sample
- 3. repeat 1 and 2 many times
- 4. for an x% CI, trim off $\frac{1}{2}\left(1-\frac{x}{100}\right)$ of the data from both ends
- 5. those trim points are the endpoints of the the x% bootstrap CI

1. & 2. Draw a Random Sample and Record Statistic

3. Repeat Many Times

Distribution of Sample Means?



• Between what two values do 95% of these samples fall?

4 & 5 Find CI Endpoints

```
In [17]: # 4. For a 95% conf. int., trim off .5*(1-(95/100)) of the data from both ends
         # calculate where to trim
         trim = .5*(1-.95) * num_iterations
         # find the closest integer
         trim = int(np.round(trim))
         trim
Out[17]: 25
In [18]: # for 1000 iterations and a 95% CI, we want to find the 25th value and (1000-25)th value
         # 5. those trim points are the endpoints of the the x% Bootstrap CI
         ci = np.sort(sample_means)[[trim,-trim]] # sort the array!
         Сi
Out[18]: array([2.66425, 3.1133])
```

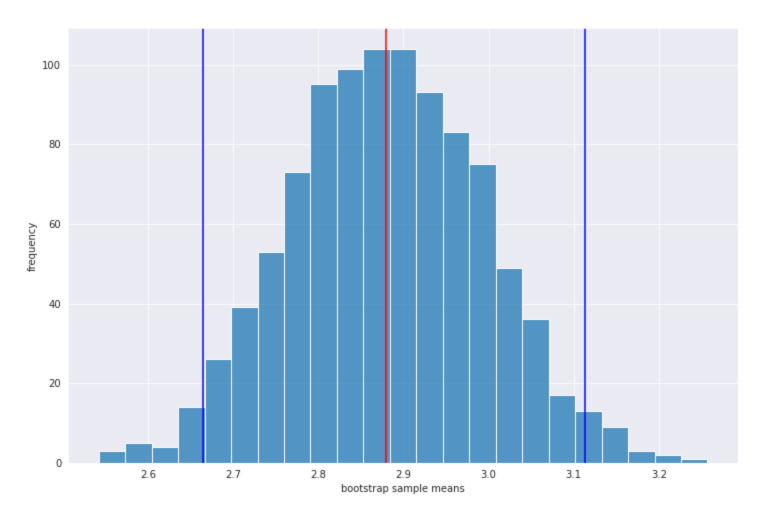
Plotting Distribution of Sample Means With Cls

```
In [19]: fig,ax = plt.subplots(1,1,figsize=(12,8))
    ax = sns.histplot(sample_means)

ylim = ax.get_ylim()
    ax.set_ylim(ylim)

ax.axvline(df.trip_distance.mean(), color='r');

ax.vlines(ci, *ylim, color='b')
    ax.set_xlabel('bootstrap sample means');
    ax.set_ylabel('frequency');
```



Interpreting Cls

- Tells us something about the variablity of this statistic.
- Tells us how confident we should be that our parameter lies in the interval.
- Does NOT tell us: "the probability that the true value lies within that interval".

If confidence intervals are constructed using a given confidence level from an infinite number of independent sample statistics, the proportion of those intervals that contain the true value of the parameter will be equal to the confidence level.

Questions re Cls?

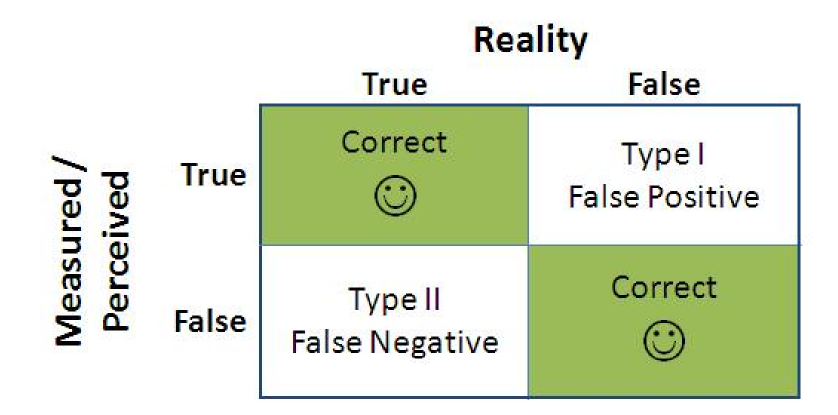
A/B Tests

- Do one of two treatments produce superior results?
 - testing two prices to determine which generates more profit
 - testing two web headlines to determine which produces more clicks
 - testing two advertisements to see which produces more conversions
- Often Used Test Statistics
 - difference in means
 - difference in counts

Hypothesis Testing

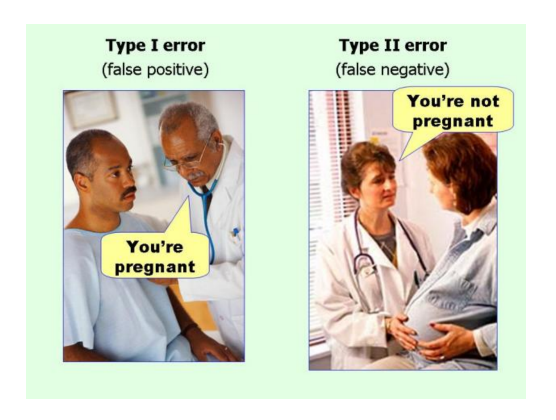
- Ex: Does one webpage lead to more sales than another?
- Null Hypothesis: H_0
 - the thing we're observing is happening due to random chance
 - there are no differences between two groups
 - Ex: A difference in sales is just random
- Alternative Hypothesis: H_1
 - the thing we're observing is happening not due to random chance
 - there is a difference between two groups
 - Ex: A difference in sales is not just random
- Experiment: given data, do we accept or reject H_0 ?
 - Ex: if we collect sales can we say that a difference between the two pages isn't random?

Errors in Hypothesis Tests



https://www.gilliganondata.com/wp-content/uploads/2009/08/Typel_TypelI1.JPG

Errors in Hypothesis Tests



https://flowingdata.com/wp-content/uploads/2014/05/Type-I-and-II-errors1-620x465.jpg

Significance and Power

- P (reject $H_0 \mid H_0$ true) = Significance of test (Type I Error)
 - Probablity of saying things aren't by chance when they are
 - Ex: Saying Ad A leads to more sales, even though there's no difference

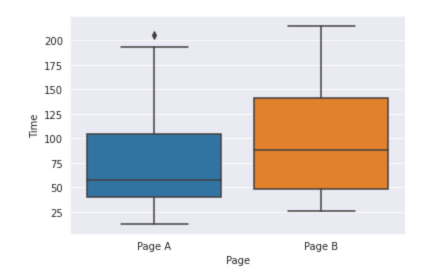
- P (reject $H_0 \mid H_1$ true) = Power of test (1-Type II Error)
 - Probability of saying things aren't by chance when they aren't
 - Ex: Saying here's no difference between ads, even though there is

Ex: Webpages and Sales

- Question: Which webpage leads to more sales?
- Potential Issue: what if sales are large but infrequent?
- Proxy Variable: stand in for true value of interest
 - Ex: Assume 'time on page' is correlated with sales

Ex: Webpages and Sales

```
In [21]: sns.boxplot(x='Page', y='Time', data=session_times);
```





Ex: Webpages and Sales, Define the Metric

- Metric: the measure we're interested in
 - Ex: We're interested in a difference of means (Page B Page A)

```
In [23]: mean_a = session_times[session_times.Page == 'Page A'].Time.mean()
    mean_b = session_times[session_times.Page == 'Page B'].Time.mean()
    observed_metric = mean_b-mean_a
    print('observed metric: {:0.2f}'.format(observed_metric))
observed metric: 21.40
```

- Is this surprising? Should we reject the null?
 - Assuming that H0 is true, is this observation surprising?

Permutation Test

Recall the Central Limit Theorem:

For reasonably large samples, the distribution of sample mean \bar{x} is normal regardless of the distribution of X.

- How do we generate additional samples of the difference in means? Resampling!
- Need to repeatedly split the data into two groups and take the differince in means
- One way to do this: combine, permute (reorder) and split

Websites and Sales, Permutation Test

```
In [24]: # 0. get group sizes
         n_a = sum(session_times.Page == 'Page A')
         n_b = sum(session_times.Page == 'Page B')
         n = n_a + n_b
         print(f'{n_a=} {n_b=} {n=}')
         n a=21 n b=15 n=36
In [25]: # 1. combine groups together (assume H0 is true)
         session_times.Time[:2]
Out[25]: 0
               12.6
              151.8
         Name: Time, dtype: float64
In [26]: # 2. permute observations
         permuted = session_times.Time.sample(frac=1, replace=False, random_state=123)
         permuted[:2]
Out[26]: 6
              126.6
               79.2
         Name: Time, dtype: float64
In [27]: # 3. create new groups
         rand_mean_a = permuted[:n_a].mean()
         rand_mean_b = permuted[n_a:].mean()
         # 4. calculate metric
         rand_mean_diff = (rand_mean_b - rand_mean_a)
         print('{:.2f}'.format(rand_mean_diff))
         -17.41
```

Websites and Sales, Permutation Test Continued

```
In [28]: # 5. repeat many times
    rand_mean_diffs = []
    iterations = 10_000

for i in range(iterations):
        permuted = session_times.Time.sample(frac=1,replace=False,random_state=i)

        rand_mean_a = permuted[:n_a].mean()
        rand_mean_b = permuted[n_a:].mean()

        rand_mean_diffs.append(rand_mean_b - rand_mean_a)

rand_mean_diffs[:5]

Out[28]: [16.257142857142857,
        16.874285714285705,
        10.2914285714285705,
        10.291428571428564,
        10.565714285714321]
```

Websites and Sales, Permutation Test Continued

```
In [29]: # 6. see where our original observation falls
         fig,ax = plt.subplots(1,1,figsize=(12,8))
         ax = sns.histplot(x=rand_mean_diffs, stat='density')
         ax.set_xlabel('random mean differences');ax.set_ylabel('frequency');
          ax.axvline(observed_metric, color='r');
            0.020
            0.015
            0.005
            0.000
                                    -20
                                        random mean differences
```

• Would be easier to interpret if this was a Standard Normal distribution...

Normalization: z-score

Convert our distribution to an approximation of standard normal

- 1. shift mean to 0
- 2. set standard deviation of 1

$$z = \frac{x - \bar{x}}{s}$$

```
In [30]: xbar = np.mean(rand_mean_diffs)
s = np.std(rand_mean_diffs)
rand_zscores = (rand_mean_diffs - xbar) / s

In [31]: observed_metric_zscore = (observed_metric - xbar) / s
```

Websites and Sales, Permutation Test Continued

```
In [32]: # 5. see where our original observation falls (normalized)
          fig,ax = plt.subplots(1,1,figsize=(12,8))
          ax = sns.histplot(rand_zscores, stat='density')
          ax.set_xlabel('random mean differences normed');ax.set_ylabel('frequency');
          ax.axvline(observed_metric_zscore,color='r');
             0.40
             0.35
             0.30
             0.25
           D 0.20
            0.15
             0.10
             0.05
             0.00
                                      random mean differences normed
```

Why Use Permutation Tests?

- data can be numeric or boolean (ex. temperature, conversion, etc)
- sample sizes can be different
- assumptions about normally distributed data are not needed

How sure are we?

• p-value

The probability of finding the observed, or more extreme, result when the null hypothesis (H_0) is true.

- does mean : P (data | H_0 is true)
- does NOT mean : $P(H_0 \text{ is not true} \mid \text{data})$
- Our question about significance becomes:

"How often did we see a value as or more extreme than our observed metric?"

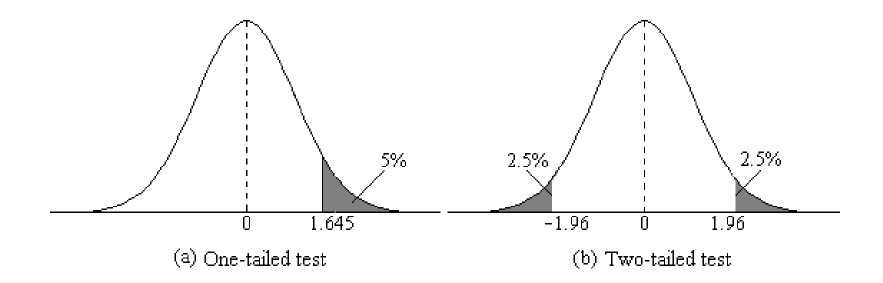
Calculating p

```
In [33]: # find absolute values greater than our observed_metric
gt = np.abs(np.array(rand_mean_diffs)) >= np.abs(observed_metric)

In [34]: # how many are greater?
num_gt = gt.sum()

# proportion of total that are as or more extreme
p = num_gt / len(rand_mean_diffs)
p
Out[34]: 0.2655
```

One-Tailed vs Two-Tailed Tests



 $\frac{https://towardsdatascience.com/one-tailed-or-two-tailed-test-that-is-the-question-1283387f631c?}{gi=9568e456cd13}$

Choosing One-Tailed vs Two-Tailed

- Do we have a strong reason for a one-tailed? One-Tailed
 - Ex: H_0 is "difference is less than or equal to 0"
 - Need a strong reason
- Otherwise? Two-tailed
 - Ex: H_0 is "there is no real difference between groups"
 - More conservative
 - Usually a better choice

One-Tailed Test Example

```
In [35]: # one-tailed test
sum(np.array(rand_mean_diffs) >= observed_metric) / len(rand_mean_diffs)
Out[35]: 0.1328
```

Note that this is less than our Two-Tailed value!

```
In [36]: # two-tailed test
sum(np.abs(np.array(rand_mean_diffs)) >= np.abs(observed_metric)) / len(rand_mean_diffs)
Out[36]: 0.2655
```

Choosing α

- alpha (α): significance level
 - What we compare our p-value to
 - Best to choose this before calculating metrics
 - Probability of rejecting the null when it is true (Type I Error)
- Common values:
 - .1 (Error 1 out of 10 times)
 - .05 (Error 1 out of 20 times)
 - .01 (Error 1 out of 100 times)
- Should depend on how bad a Type I (False Positive) Error is

Another Example: Price vs Conversion

- Does Price A lead to higher conversions than Price B?
- Conversion: Turning a visit into a sale
- H_0 : conversions for Price A \leq conversions for Price B
 - Price A does not lead to more conversions
- H_1 : conversions for Price A > conversions for Price B
 - Price A leads to more conversions

Another Example: Price vs Conversion

- Does Price A lead to higher conversions than Price B?
- Conversion: Turning a visit into a sale
- H_0 : conversions for Price A \leq conversions for Price B
 - Price A does not lead to more conversions
- H_1 : conversions for Price A > conversions for Price B
 - Price A leads to more conversions

- Metric of Interest?
 - difference in percent conversion

- First: Choose our α : 0.05
- Reminder of Permutation Test:
 - 0. get group sizes
 - 1. combine groups together
 - 2. permute observations
 - 3. create two new groups (same sizes as originals)
 - 4. calculate metric
 - 5. repeat many times
 - 6. see where our original observation falls

- What are our samples?
 - 1 = Conversion
 - 0 = No conversion
- How many samples are there?

```
In [40]: n = df.sum().sum()
n
Out[40]: 46327
```

• Turning counts into samples

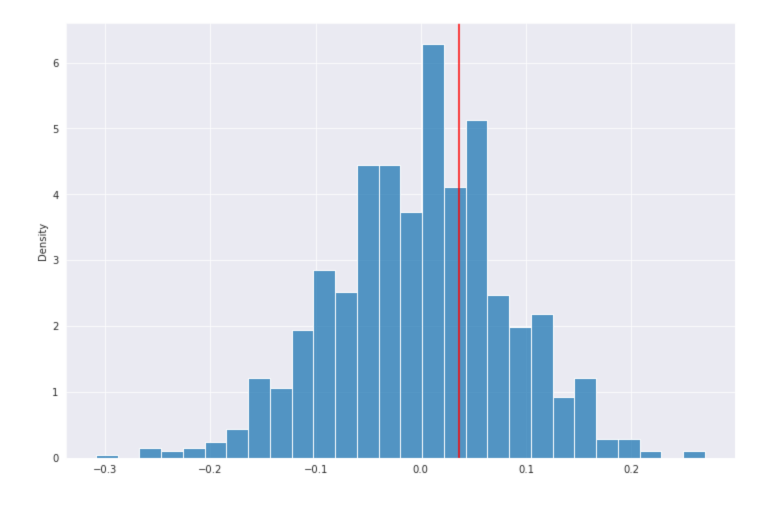
```
In [41]: n_conversion = df.loc['Conversion'].sum()
n_conversion

Out[41]: 382

In [42]: samples = np.zeros(n)
samples[:n_conversion] = 1
    assert sum(samples) == n_conversion
```

```
In [43]: n_a, n_b = df.sum(axis=0)
         print(f'{n_a=} {n_b=} {n=}')
         assert n_a + n_b = n
         n a=23739 n b=22588 n=46327
In [44]: # tqdm gives us a progress bar
         # conda install -n eods-f20 ipywidgets
         from tqdm.notebook import tqdm
In [45]: np.random.seed(123)
         rand_conv_diffs = []
         for i in tqdm(range(1000)):
             permutation = np.random.permutation(samples)
             rand_conv_a = sum(permutation[:n_a]) / n_a
             rand_conv_b = sum(permutation[n_a:]) / n_b
             rand_conv_diffs.append(100 * (rand_conv_a - rand_conv_b))
          100%
                                    1000/1000 [00:15<00:00, 63.82it/s]
```

```
In [46]: fig,ax = plt.subplots(1,1,figsize=(12,8))
ax = sns.histplot(x=rand_conv_diffs, stat='density')
ax.axvline(diff_pct_conv,color='r');
```



```
In [47]: # calculate a two-tailed p-value
sum(np.array(rand_conv_diffs) >= diff_pct_conv) / len(rand_conv_diffs)
Out[47]: 0.35
```

Equation Based Proportion Test

Equation Based Hypothesis Test: t-Test

- based on the Student-t distribution
- more involved to describe
- works for numeric data (can't use it for the last example)

• very close to the 0.27 value we found via permutation test

How Many Observations Do We Need?

Common question: How many observations do we need?

These 4 things are related:

- effect size: Min size of effect you want to detect
 - Ex: "10% increase in clicks"
- power: Prob. of detecting given effect size with given sample size
 - $P(\text{reject } H_0 \mid H_1 \text{ true}) \text{ or } 1 P(\text{accept } H_0 \mid H_1 \text{ true}) \text{ aka 1-Type II Error}$
- alpha: significance level at which to reject the null
 - $P(\text{reject } H_0 \mid H_0 \text{ true})$ aka Type I Error
- number of observations: usually what we're interested in finding

Knowing 3 gets you the 4th!

How Many Observations Do We Need? Continued

How Many Observations Do We Need? Continued

What happens to our power if we can only get 1000 observations?

Issues with Multiple Testing

- p-hacking: keep trying comparisons till you find something that works
- multiple tests: the more tests you run, the more likely a Type 1 Error
 - Bonferonni correction: $\frac{\alpha}{m}$

Statistically Significant?

The ASA Statement on p-Values: Context, Process, and Purpose Wasserstein & Lazar, 09 Jun 2016]

- Don't base your conclusions solely on whether an association or effect was found to be "statistically significant" (i.e., the p-value passed some arbitrary threshold such as p < 0.05).
- Don't believe an association/effect exists just because it was statistically significant.
- Don't believe an association/effect is absent just because it was not stat. significant.
- Don't believe that your p-value:
 - 1. gives the **probability that chance alone** produced the observed association/effect or
 - 2. the probability that your **test hypothesis is true**.]
- Don't conclude anything about **scientific or practical importance** based on statistical significance (or lack thereof).

Statistically Significant?

Moving to a World Beyond "p < 0.05" Wasserstein, Schirm & Lazar, 20 Mar 2019

- Try to avoid "Statistically Significant"
- "Accept uncertainty. Be thoughtful, open, and modest." Remember "ATOM."

Statistically Significant?

Moving to a World Beyond "p < 0.05" Wasserstein, Schirm & Lazar, 20 Mar 2019

- Try to avoid "Statistically Significant"
- "Accept uncertainty. Be thoughtful, open, and modest." Remember "ATOM."
- ATOM
 - A: Seek better measures, more sensitive designs, larger samples
 - **T:** Begin with clearly expressed objectives
 - **T**: Ask "What are the practical implications?"
 - O:: Be open/transparent in analysis and communication
 - M: Accept limititaions, assumptions, reproduction, recognizing differences in stakes

Comparing More Than 2 Groups

- ANOVA
 - need more stats than we have time for
- Multi-Armed Bandit (MAB)
 - can compare many distributions
 - don't need to make assumptions about underlying distributions
 - can also be used for early stopping of experiment

Multi-Armed Bandit



Question: Which arm should we pull?

Greedy MAB

greedy: do something simple that heads towards the goal

1. pull arm with highest payout

But what if there's a better choice, we just haven't seen it yet?

Exploration Vs Exploitation

- **Exploration:** There might be a better arm
 - keep choosing different arms randomly
- Exploitation: We want to make use of the best
 - keep pulling the best arm

ϵ -Greedy MAB

- choose a small epsilon (ϵ) between 0 and 1
- 1. generate random number between 0 and 1
- 2. if $< \epsilon$, choose arm randomly
- 3. if $\geq \epsilon$, choose best arm
- 4. GOTO 1

MAB Example

- We have two ads
- We don't know how often each will lead to a response
- We need to decide which ad to add to each page request

• We'll use an ϵ -greedy MAB to decide which ad to show

```
In [54]: # epsilon probability
epsilon = 0.40
```

- Rounds 1 and 2
 - Pull each arm once

```
In [55]: pulls_A = [ad_A.rvs()]
    pulls_B = [ad_B.rvs()]
    pulls_A,pulls_B

Out[55]: ([0], [1])
```

- Round 3
 - With probability 1ϵ , choose the best arm (B)

```
In [56]: be_greedy = np.random.rand() > epsilon
be_greedy

Out[56]: True

In [57]: pulls_B.append(ad_B.rvs())
    pulls_A,pulls_B

Out[57]: ([0], [1, 0])
```

```
In [58]: def mab(ad_A, ad_B, pulls_A, pulls_B, epsilon):
             be_greedy = np.random.rand() > epsilon
             if not be_greedy: # randomly choose
                 if np.random.rand() < 0.5:</pre>
                     pulls_A.append(ad_A.rvs())
                     choice = 'A'
                 else:
                     pulls_B.append(ad_B.rvs())
                     choice = 'B'
             else: # be greedy
                 resp_A = sum(pulls_A) / len(pulls_A)
                 resp_B = sum(pulls_B) / len(pulls_B)
                 if resp_A > resp_B:
                     pulls_A.append(ad_A.rvs())
                     choice = 'A'
                 else:
                     pulls_B.append(ad_B.rvs())
                     choice = 'B'
             return pulls_A, pulls_B, be_greedy, choice
```

• Round 4

```
In [60]: for i in range(10):
             pulls_A, pulls_B, be_greedy, choice = mab(ad_A, ad_B, pulls_A, pulls_B, epsilon)
             print(f'{str(be_greedy):5s} {choice} => '+
                   f'{np.mean(pulls_A):0.2f}:{np.mean(pulls_B):0.2f} {str(pulls_A):20s}, {str(pulls_B):20s}')
         True B => 0.00:0.50 [0]
                                                    , [1, 0, 1, 0]
         False A => 0.50:0.50 [0, 1]
                                                    , [1, 0, 1, 0]
         True B \Rightarrow 0.50:0.40[0, 1]
                                                   , [1, 0, 1, 0, 0]
         True A \Rightarrow 0.67:0.40 [0, 1, 1]
                                                   , [1, 0, 1, 0, 0]
                                                   , [1, 0, 1, 0, 0]
         False A => 0.75:0.40 [0, 1, 1, 1]
         True A => 0.80:0.40 [0, 1, 1, 1, 1]
                                                   , [1, 0, 1, 0, 0]
         False B => 0.80:0.33 [0, 1, 1, 1, 1]
                                                   , [1, 0, 1, 0, 0, 0]
                                                   , [1, 0, 1, 0, 0, 0, 1]
         False B => 0.80:0.43 [0, 1, 1, 1, 1]
         False B => 0.80:0.50 [0, 1, 1, 1, 1]
                                                   , [1, 0, 1, 0, 0, 0, 1, 1]
         False B => 0.80:0.44 [0, 1, 1, 1, 1]
                                                    , [1, 0, 1, 0, 0, 0, 1, 1, 0]
```

• Which arm seems best?

```
In [61]: print(f'conversion rates: A: {np.mean(pulls_A):0.2f} B: {np.mean(pulls_B):0.2f}')
conversion rates: A: 0.80 B: 0.44
```

• Did we pick the best one?

```
In [62]: print(f'ground truth: A: {ad_A.pmf(1):0.2f} B: {ad_B.pmf(1):0.2f}')
ground truth: A: 0.78 B: 0.24
```

MAB Variations

- Thompson's Sampling: uses Baysian approach
- UCB1: maximize expected reward using Upper Confidence Bounds
- UCBC: Upper Confidence Bound with Clusters

• ...

Questions?

Python Argument Unpacking

```
In [63]: ax = sns.histplot(np.random.normal(size=100));
         plt.tight_layout()
         # to get y-axis min and max, use ax.get_ylim()
         ymin,ymax = ax.get_ylim()
         ax.set_ylim(ymin,ymax)
         # to add a vertical line use vlines, need ymin, ymax
         ax.vlines(1, ymin, ymax, color='red');
         # ax.get_ylim() returns a tuple, so can't pass directly to vlines
         print(type(ax.get_ylim()))
         # can unpack the arguments with * (asterisk)
         ax.vlines(-1, *ax.get_ylim(), color='orange');
         # or, for vertical lines can use axvline
         ax.axvline(-2,color='black');
         <class 'tuple'>
```

