```
Inferential statistics II - Bootstrapping
           Introduction
           In the previous frequentist mini-projects, you did frequentist calculations to perform inference from a sample of data. Such
           inference relies on theory largely developed from the 19th-Century onwards that is subject to certain assumptions or
           theoretical limits. These are fine if those assumptions hold for the particular case you're working on, and what you want to do
           has a known theoretical distribution (for example the mean of a sampling distribution that we looked at in the previous mini-
           project.)
           In this mini-project, you'll use the same medical charge data you used in the frequentist inference mini-project, but this time
           you'll make inferences about the population using bootstrapping (ie. simulating repeated re-runs of an experiment.) If
           frequentism is about using assumptions and theoretical results to calculate what we expect to happen were an experiment to
           be run again and again and again, then bootstrapping is about using computing power to essentially re-run the sample draw
           again and again and again to see what actually happens.
           Prerequisites
           While these exercises do not strictly depend on these concepts, we encourage you to complete the previous mini-projects
           before starting this one so that you can approach this assignment with a good understanding of frequentist concepts like:
            • the z-statistic
            • the t-statistic

    the difference and relationship between the two

    the Central Limit Theorem, its assumptions and consequences

    how to estimate the population mean and standard deviation from a sample

    the concept of a sampling distribution of a test statistic, particularly for the mean

    how to combine these concepts to calculate confidence intervals and p-values

    how those confidence intervals and p-values allow you to perform hypothesis (or A/B) tests

           To complete mini-project, it's important that you first complete the bootstrap resources listed in this subunit, as they contain
           valuable information about how to calculate bootstrap replicates of summary statistics. Having an basic understanding of what
           confidence intervals and p-values are will also be helpful (we touch on them in this mini-project, but please speak to your
           mentor or conduct individual research if you'd like to learn more.)
 In [1]: import pandas as pd
           import numpy as np
           from numpy.random import seed
           from scipy import stats
           import matplotlib.pyplot as plt
           Medical charge data set
 In [2]: med charges = pd.read csv('data/insurance2.csv')
 In [3]: med charges.head()
 Out[3]:
              age sex
                          bmi children smoker region
                                                          charges insuranceclaim
                     0 27.900
                                                    3 16884.92400
           1 18
                   1 33.770
                                                    2 1725.55230
           2 28 1 33.000
                                                    2 4449.46200
           3 33 1 22.705
                                                   1 21984.47061
           4 32 1 28.880
                                                    1 3866.85520
           In the previous assignment, you used the frequentist approach to estimate the lower limit for the 95% confidence interval on
           the mean hospital charge. This approach relies on statistical theory that has been developed over the years and is also limited
           to statistics for which theoretical results on the sampling distribution exist. These results are remarkably useful and applicable
           much of the time and under a surprisingly wide range of conditions.
           Having calculated the 95% lower confidence interval using frequentist theory in the previous exercise, you'll now use bootstrap
           inference to verify your calculations and check that you get consistent results without making the assumptions required before.
           After all, the distribution of charges really was very non-normal.
           Q: Use bootstrap sampling to estimate the same 95% confidence interval lower limit as before.
           A: Use conf_int = np.percentile(bs_replicates, [5]) # 95% lower limit
 In [4]: # Define a function we can use again
           def draw_bs_reps(data, func, size=1):
               11 11 11
               Draw bootstrap replicates.
                data = one-dimensional array of values
                 func = the function to run on each replicate, e.g. mean
                 size = the number of replicates to draw (generate)
               # Initialize array of replicates: bs_replicates
               bs_replicates = np.empty(size)
               # Generate replicates
               for i in range(size):
                    bs_replicates[i] = func(
                                           np.random.choice(
                                               data, size=len(data)
               return bs replicates
           #end_def
 In [5]: # Draw 10000 replicates of medical charges and
           # calculate the 95% lower confidence interval
           np.random.seed(47)
           N_{rep} = 10000
           bs_replicates = draw_bs_reps(med_charges.charges, np.mean, N_rep)
           conf_int = np.percentile(bs_replicates, [5]) # 95% lower limit
           conf_int
 Out[5]: array([12724.4679217])
           If you performed 10000 replicates immediately after setting the random seed to 47, you should get the value 12724 here,
           which compares very well with the value 12725 obtained using the t-distribution confidence interval previously. It is a most
           pleasant result to see the predictions of classical frequentist theory match with results that are now possible through the
           number-crunching ability of computers.
           Remember, in the previous mini-projects, we saw that there are two ways of performing a t-test from a sample, depending on
           whether we can assume the groups have equal variance or not. We can actually easily test this using the bootstrap approach!
           Q: Calculate the 95% confidence interval for the difference between the standard deviations of insurance and non-insurance
           claim charges (insured - non-insured). Calculate the differences over 10000 replicates. Plot the histogram of values and mark
           the locations of the percentiles. State the null and alternative hypothesis and comment on whether you would retain or reject
           the null hypothesis in this case and why.
           A: The null hypothesis: there is no difference between the standard deviations of insurance and non-insurance claim charges.
 In [6]: # not insured
           ins0 = med_charges.loc[med_charges.insuranceclaim == 0]
           ins1 = med charges.loc[med charges.insuranceclaim == 1]
 In [7]: np.random.seed(47)
           N \text{ rep} = 10000
           nonins_bs_replicates = draw_bs_reps(ins0.charges, np.std, N_rep)
           ins_bs_replicates = draw_bs_reps(ins1.charges, np.std, N_rep)
           diffs = ins bs replicates - nonins bs replicates
           print("insured\n", ins_bs_replicates)
           print("not insured\n", nonins bs replicates)
           print("diffs\n", diffs)
           insured
            [14165.0062503 14176.49758651 14109.56522567 ... 14112.40608661
           14420.49156918 14163.2373204 ]
           not insured
            [6159.05411195 6115.51989319 6642.76007193 ... 6510.627812 6371.59338214
            6269.25813899]
           diffs
            [8005.95213835 8060.97769333 7466.80515375 ... 7601.77827461 8048.89818704
            7893.97918141]
 In [8]: axvline kwargs = {"linestyle" : "dashed", "linewidth" : "1"}
           diffs s = pd.Series(diffs)
           diffs s.hist(bins=20, grid=False,
                       figsize=(14,8), xlabelsize=10, edgecolor='k')
           plt.xlabel("Dollars", fontsize=15)
           plt.ylabel("Frequency", fontsize=15)
           plt.axvline(np.percentile(diffs s, [97.5]), **axvline kwargs, label="2.5%")
           plt.axvline(diffs s.mean(), **axvline kwargs, color="red", label="mean")
           plt.axvline(np.percentile(diffs_s, [2.5]), **axvline_kwargs, label="97.5%")
           plt.legend(loc='upper right')
           plt.show()
                                                                                                                 ---- 2.5%
              1600
                                                                                                                 ---- 97.5%
              1400
              1200
              1000
            Frequency
               800
               600
               400
               200
                                                                   Dollars
 In [9]: conf int = np.percentile(diffs, [2.5, 97.5]) # 95%
           lower, upper = conf int
           print("The 95% confidence interval is between", round(lower,2), "and", round(upper,2))
           The 95% confidence interval is between 6685.46 and 8510.07
           A: There is definitely a difference between the standard deviations of insurance and non-insurance claim charges. A difference
           of anything close to 0 isn't even on the plot.
           Confidence interval, \alpha, and p-value
           The confidence interval is often a useful quantity to estimate. We wish to limit our expected probability of making a Type I
           error<sup>[1]</sup> to alpha (\alpha)<sup>[2]</sup>. The associated confidence interval is our estimate of the interval within which we expect the true
           population value to be found (1 - \alpha) \times 100 \% of the time we do this test.
           In the exercise above, we performed bootstrap replicates to estimate the interval and reject the null hypothesis if this interval
           did not contain zero. You will sometimes see such an interval reported in the output of statistical functions.
           The partner of the confidence interval is the p-value. The p-value and the confidence interval are linked through our choice of
           α. The p-value tells us how likely it is, under the null hypothesis, to get an outcome at least as extreme as what was observed.
           If this fails to reach the level of our pre-specified \alpha, we decide the null hypothesis is sufficiently unlikely to be true and thus
           reject it. To calculate this p-value via the bootstrap, we have to put ourselves in a position where we are simulating the null
           hypothesis being true and then calculate the fraction of times we observe a result at least as extreme as that actually
           observed.
           Remember how, previously, you used the t-test to calculate the p-value for the observed difference between the means of
           insured and non-insured medical cases. We're now going to repeat this, this time using the bootstrap approach.
          [1] Type 1 error: where we wrongly reject the null hypothesis, and it is, instead, true)
           ^{[2]} alpha (a) is the desired significance level, typically 0.05 or lower.
           See Also: What is the difference between an alpha level and a p-value?
           Q: Perform a bootstrapped hypothesis test at the 5% significance level (\alpha = 0.05) to calculate the p-value of the observed
           difference between insurance and non-insurance charges, state your null and alternative hypotheses and whether you retain
           or reject the null hypothesis for the given significance level.
           A: The null hypothesis: there is no significant difference between insurance and non-insurance charges. (The difference seen
           in the dataset is due to chance.)
           The alernative hypothesis: there is a significant difference between insurance and non-insurance charges.
                  To do the two-sample bootstrap test, we shift both arrays to have the same mean, since we are simulating the
                  hypothesis that their means are, in fact, equal. We then draw bootstrap samples out of the shifted arrays and
                  compute the difference in means. This constitutes a bootstrap replicate, and we generate many of them. The
                  p-value is the fraction of replicates with a difference in means greater than or equal to what was observed.
           reference: DataCamp Statistical thinking in Python, part 2
           Null hypothesis: charges are identically distributed between insured and uninsured patients (there are no differences due to
           insurance; shuffling samples shouldn't matter.)
           Test statistic: if the populations are "identical", they should have the same mean charge. The difference in mean charge should
           be 0.
            1. Use the function, diff_of_means to compute the empirical difference of means that was observed between charges for
               insured vs uninsured patients in the original dataset.
            2. Shift both arrays to have the same mean
            3. Draw 10,000 replicates out of the shifted arrays.
            4. Use the function, diff_of_means to compute the difference of means for he N permuted samples.
            5. Compare the differences from the original data with the randomly computed differences.
            6. Compute the p-value as the fraction of replicates with a difference in means greater than or equal to what was observed
In [10]: # Define the functions we will use
           def diff of means(data 1, data 2):
               Difference in means of two arrays.
               data1 - data2
               # The difference of means of data_1, data_2
               diff = np.mean(data_1) - np.mean(data_2)
               return diff
           #end def
In [11]: # not insured
           # ins0.charges is the Series/array of interest
           ins0 = med charges.loc[med charges.insuranceclaim == 0]
           ins0_charges = ins0.charges.values # numpy array
           # not insured
           # insl.charges is the Series/array of interest
           ins1 = med_charges.loc[med_charges.insuranceclaim == 1]
           ins1_charges = ins1.charges.values # numpy array
In [12]: # Compute empirical (observed in the data we have) difference in means
           # Use absolute value bcause we don't care (or pretend not to know)
           # which way the difference goes.
           empirical_diff_means = diff_of_means(ins1.charges, ins0.charges)
           # Mean of all charges (if insurance doesn't matter, this is relevant)
           # This is equivalent to the mean of the concatenated sub-populations
           mean_charges = np.mean(med_charges.charges)
           # Generate shifted arrays
           ins0_shifted = ins0_charges - np.mean(ins0_charges) + mean_charges
           ins1_shifted = ins1_charges - np.mean(ins1_charges) + mean_charges
In [13]: # Compute 10,000 bootstrap replicates from shifted arrays
           nonins_bs_replicates = draw_bs_reps(ins0_shifted, np.mean, size=10000)
           ins_bs_replicates = draw_bs_reps(ins1_shifted, np.mean, size=10000)
           # Get replicates of difference of means
           # subtracing in the same order as for empirical_diff_means
           bs replicates diff = ins bs replicates - nonins bs replicates
           # Compute and print p-value
           p = np.sum(bs replicates diff >= empirical diff means) / len(bs replicates diff)
           print('p-value =', p)
           p-value = 0.0
           We reject the null hypothesis for the given significance level.
           Q: To put the above result in perspective, plot the histogram of your bootstrapped differences along with lines marking the
           locations of the observed difference. (Why would we plot more than one line, given that we only have one observed
           A: "Why would we plot more than one line, given that we only have one observed difference?)"
           I have no idea.
          axvline kwargs = {"linestyle" : "dashed", "linewidth" : "1"}
           plt.figure(figsize=(12,8))
           plt.hist(ins bs replicates, bins=20, label="insurance means")
           plt.hist(nonins_bs_replicates, bins=20, label="non-insurance means")
           plt.hist(bs_replicates_diff, bins=20, label="difference in means")
           plt.xlabel("Dollars", fontsize=15)
           plt.ylabel("Frequency", fontsize=15)
           plt.axvline(empirical_diff_means, **axvline_kwargs, label="empirical mean difference")
           plt.axvline(mean_charges, **axvline_kwargs, color="k", label="mean of all charges")
           plt.legend(loc='best')
           plt.show()
              1600
              1400
               1200
            Frequency
               800
               600
               400
                                                            empirical mean difference
                                                         ---- mean of all charges
               200
                                                            insurance means

    non-insurance means

    difference in means

                                                          5000
                                                                                   10000
                    -2500
                                                                       7500
                                                                                               12500
                                                                Dollars
           Q: Compare your p-value above with that obtained using the t-test function in the previous assignment. Do you think you
           would want to try to perform enough bootstrap replicates to observe a random difference as large as that we did observe?
           A: In the previous assignment, I got the p-value, 4.46e-31. This time, I got 0.0.
           They're effectively equivalent.
           I see no reason to perform more bootstrap replicates.
                  The number of bootstrap samples recommended in literature has increased as available computing power has
                  increased. If the results may have substantial real-world consequences, then one should use as many samples
                  as is reasonable, given available computing power and time. Increasing the number of samples cannot
                  increase the amount of information in the original data; it can only reduce the effects of random sampling errors
                  which can arise from a bootstrap procedure itself. Moreover, there is evidence that numbers of samples
                  greater than 100 lead to negligible improvements in the estimation of standard errors. In fact, according to the
                  original developer of the bootstrapping method, even setting the number of samples at 50 is likely to lead to
                  fairly good standard error estimates. [Wikipedia]
                  (https://en.wikipedia.org/wiki/Bootstrapping_(statistics)#Recommendations)
           Why did we do a boostrap test? Why not a permuation test?
           Try a permuation test. According o Stack Exchange, "The permutation test is best for testing hypotheses and bootstrapping is
           best for estimating confidence intervals."
           "Permutation tests test a specific null hypothesis of exchangeability, i.e. that only the random sampling/randomization explains
           the difference seen."
           Permutation tests requie that we have wo sample arrays (which we do) and cannot look for a difference in distribution (which
           we don;t care about here).
In [15]: # not insured
           ins0 = med charges.loc[med charges.insuranceclaim == 0]
           # not insured
           ins1 = med charges.loc[med charges.insuranceclaim == 1]
           # we don't need to concatenate the populations;
           # they're already mixed in the original dataset
           # but for purposes of writing a moe generic function, we'll
           # separate and concatnate
           def permutation sample(data1, data2):
               """Generate a permutation sample from two data sets."""
               # Concatenate the data sets: data
               data = np.concatenate((data1, data2))
               # Permute the concatenated array: permuted data
               permuted data = np.random.permutation(data)
               # Split the permuted array into two parts
               # Note position of colons in the slices!
               perm sample 1 = permuted data[:len(data1)]
               perm sample 2= permuted data[len(data2):]
               return perm sample 1, perm sample 2
           #end def
           def draw perm reps(data 1, data 2, func, size=1):
               """Generate multiple permutation replicates."""
               # Initialize array of replicates: perm replicates
               perm replicates = np.empty(size)
               for i in range(size):
                  # Generate permutation sample
                   perm_sample_1, perm_sample_2 = permutation_sample(data_1, data_2)
                    # Compute the test statistic
                    perm replicates[i] = func(perm sample 1, perm sample 2)
               return perm replicates
           #end def
           # diff of means has already been defined
           Null hypothesis: charges are identically distributed between insured and uninsured patients (there are no differences due to
           insurance.
           Test statistic: if the populations are "identical", they should have the same mean charge. The difference in mean charge should
           be 0.
            1. Use the function, diff_of_means to compute the empirical difference of means that was observed between charges for
               insured vs uninsured patients in the original dataset.
            2. Draw 10,000 permutation replicates, Using the function, diff_of_means to compute the difference of means for the
               permuted samples.
            3. Compare the differences from the original data with the randomly computed differences.
            4. Compute the p-value.
In [16]: # Compute empirical difference from dataset
           empirical diff means = diff of means(ins1.charges, ins0.charges)
           # Draw 10,000 permutation replicates
           perm replicates = draw perm reps(ins1.charges, ins0.charges,
                                                 diff of means, size=10000)
           # Compute p-value
           p = np.sum(perm_replicates >= empirical_diff_means) / len(perm_replicates)
           print("p value:", p)
           p value: 0.0
In [17]: observed_diff = np.mean(ins1.charges) - np.mean(ins0.charges)
           observed diff
Out[17]: 7602.506384231368
In [18]: axvline_kwargs = {"linestyle" : "dashed", "linewidth" : "1"}
           plt.figure(figsize=(10,8))
           plt.hist(perm_replicates, bins=20, label="difference in means")
           plt.xlabel("Dollars", fontsize=15)
           plt.ylabel("Frequency", fontsize=15)
           plt.axvline(np.percentile(perm_replicates, [97.5]), **axvline_kwargs, label="2.5%")
           plt.axvline(np.percentile(perm_replicates, [2.5]), **axvline_kwargs, label="97.5%")
           plt.axvline(observed diff, **axvline kwargs, color="red", label="observed")
           plt.legend(loc='upper right')
           plt.show()
                                                                                ---- 2.5%
              1400
                                                                                 ---- 97.5%
                                                                                    observed
                                                                                    difference in means
              1200
              1000
            Frequency
               800
               400
               200
                    -2000
                                                    2000
                                                                   4000
                                                                                   6000
                                                                                                  8000
                                                        Dollars
           Q: Consider the two variants of the t-test we performed in the previous assignment. Which one would you use now?
           A: By, "two variants", do you mean manually by formula vs. using scipy.stats?
           I would absolutely use scipy.stats. It's less prone or error cause by a mistake in parentheses, ordering of operations, or
           memory.
           Or, by "two variants", do you mean assuming variance is the same or different? I would assume different.
```

Q: If, instead of being asked whether the means of two groups were different, you were working with an ad-recommender team who wanted to release a new recommendation algorithm, and you were looking at click-through rate both for the current algorithm (call it A) and from trials of their new algorithm (call it B), would you perform a two-sided test as above? What would be your null and alternative hypotheses and what would be the real-world consequence of rejecting the null hypothesis?

The null hypohesis is, as usual, that nothing changes. There is no difference in clickthrough rate between algorithm A and

The alternative hypothesis is that there is a change. Se would like the change to be positive (higher click through rate).

The real-world consequence would be that we might replace algorithm A. If we are correct, he clickthrough rate will really increase and we might make more money. But it costs money to replace the recommendation algorithm, so we had better

Loading [MathJax]/jax/output/HTML-CSS/fonts/TeX/fontdata.js

algorithm B.

A: We should perform an A/B test.

make enugh more to offset that.