# Chapter 14. Generating more samples; resampling methods

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# Introduction to Resampling

Resampling methods are powerful and widely used in computational statitics. By repeatedly re-sampling data some of the assumptions of classical statistical methods can be relaxed. These computationally intensive methods are products of the computer age. Resampling methods provide a natural way to find uncertainty when performing statistical inferences.

Re-sampling methods draw heavily on the central limit theorem (CLT) (Chapter XX) and the weak law of large numbers (Chapter XX). The weak law of large numbers tells us that a resampled estimate of a static converges to the correct value, when the conditions re met. The CLT tells us that the distribution of mean estiamtes is Normally distributed, as the number of resamples increases.

There are several use cases for resampling methods. Specifically re-sampling methods:

- Allow computation of statistics from limited data samples.
- Esimate a probability distribution of a statistic.
- Make minimal distributional assumptions, when compared to classical frequentist statistics.
- Are computationally intensive, but often highly parallelizable.

Commonly used re-sampling methods include:

- Randomization or Permutation methods: hypothesis tests.
- Non-parametric bootstrap resampling: compute statistics.
- Jackknife: or leave one out re-sampling.
- Cross validation: re-sample into multiple folds without replacement.

Re-sampling methods are general and powerful but, there is no magic involved! When using re-sampling methods always keep in mind these pitfalls: - If a sample is biased, the re-sampled statistic estimated from that sample will be biased. - The sample variance and Cis can be no better than the data sample allows.

#### Randomization and permutation methods

Randomization and permutation methods were pioneered by Fisher as early as 1911. Fisher fully developed the theory in his 1935 book. Scalability of these permutation methods remain limited, even with modern computers.

# **Cross-validation**

Today, cross-validation is widely used in the testing of machine learning models. Cross-validation was originally proposed by Kurtz in 1948. Mosier extended the method to double cross validation in 1951. The modern method of nested or multicross-validation were introduced by Krus and Fuller in 1982.

#### Jack knife methods

Jack knife methods are often effective when there are only limited data samples. Maurice Quenouille originally suggested this method in 1949. The jack knife was fully developed by John W. Tukey, who gave the method its name, in 1958. Tukey saw that method as a simple tool useful for many purposes like a pocket knife.

#### **Bootstrap**

The bootstrap is an extremely general and powerful re-sampling method. In principle, the bootstrap algorithm and can provide estimates of the distributions of most any statistic. The bootstrap method was first suggested by Efron and Hinkley in 1978 and further developed by Efron in 1979. A full treatment was provided in Efron's 1980 book.

As with other re-sampling methods, the bootstrap algorithm is computationally intensive. However, with increased computing power, use of bootstrap methods continues to expand. Further, the algorithm can be readily parallelized.

### Point Estimates vs. Distribution Estimates

The goal of **frequentist statistics** is to compute a **point estimate** and **confidence interval** for the point estimate. By a point estimate, we mean a single most likely value for a statistic. The confidence interval is based on the properties of some assumed probability distribution of the statistic. Up to now we have worked with hypothesis tests and summary statistics computed with frequentist methods. For example, with the t-test we found the most likely value of the t-statistic and the confidence interval assuming the difference of means follows a t-distribution.

Are there alternative to this frequentist approach? Yes, indeed there are. The most widely used alternative are found in **Bayesian statistics**. In this lesson, we will focus on re-sampling methods, particularly **bootstrap** methods. The bootstrap lies somewhere between classical frequentist statistics and Bayesian methods.

Rather than computing a point estimate directly, bootstrap methods compute a **bootstrap distribution** of the statistic. The bootstrap distribution is comprised of values of the statistic computed from **bootstrap samples** of the original data sample. Based on this distribution a mostly likely point estimate of the statistic, or **bootstrap estimate** (the mean of the distribution) is computed. The **bootstrap confidence interval** is also computed from the bootstrap distribution. This approach is in contrast to the purely frequentist approach of computing point estimates and confidence intervals using the original data sample.

# Overview of the Bootstrap Algorithm

The bootstrap method follows a simple algorithm. Estimates of the statistic are accumulated by these steps:

- 1. Randomly sample (e.g. Bernoulli sample) N data with replacement from an original data sample of N values. That is, the re-sample is the same size as the original data sample.
- 2. Re-compute the statistic with the current sample sample. This is a **bootstrap sample** of the statistic.
- 3. Repeat steps 1 and 2 to accumulate the required number of bootstrap samples.
- 4. Accumulated statistic values form the **bootstrap distribution**.
- 5. The mean of the computed statistic values is the **bootstrap point estimate** of the statistic.

For example, you can compute the bootstrap mean as:

$$Meanboot = \frac{\Sigma_i mean(sample_i)}{nsample}$$

where, for example with 10 data values and example of the ith re-sample might be:

$$sample_i = X_1 + X_2 + X_3 + X_4 + X_5 + X_6 + X_7 + X_8 + X_1 + X_5$$

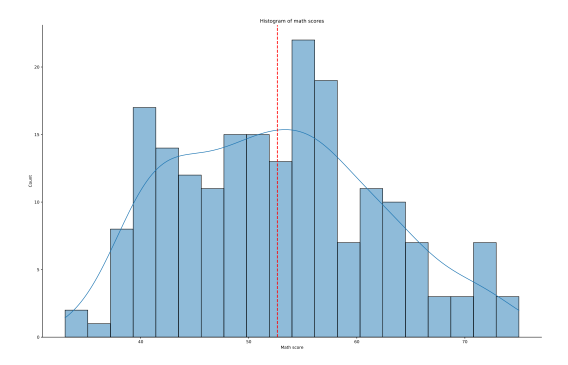
### Example; single sample bootstrap

In the previous example you computed the means and confidence intervals of the sample distributions of the male and female adult child height data. Now you will bootstrap the difference in means to determine if it is significant.

```
import pandas as pd
import numpy as np
import numpy.random as nr
import matplotlib.pyplot as plt
import seaborn as sns
import statsmodels.api as sm

test_scores = pd.read_csv('../data/hsb2.csv', index_col=0)
test_scores.head()
```

```
##
        female race ses schtyp prog read write math science socst
## id
## 70
             0
                        1
                                1
                                      1
                                                         41
                                                                  47
                                                                         57
## 121
             1
                        2
                                1
                                      3
                                           68
                                                  59
                                                        53
                                                                  63
                                                                         61
## 86
             0
                        3
                                1
                                      1
                                           44
                                                  33
                                                        54
                                                                  58
                                                                         31
## 141
             0
                   4
                        3
                                1
                                      3
                                           63
                                                  44
                                                         47
                                                                  53
                                                                         56
## 172
                                                                         61
```



The code in the cell below generates bootstrap samples from the full male and female data sub-sets and then computes the difference in the means. The result is the bootstrap distribution of the difference of means. The point estimate of the difference in means is then the mean of this distribution. And, the confidence intervals for this point estimate are also computed from this distribution.

Run the code and examine the plotted results.

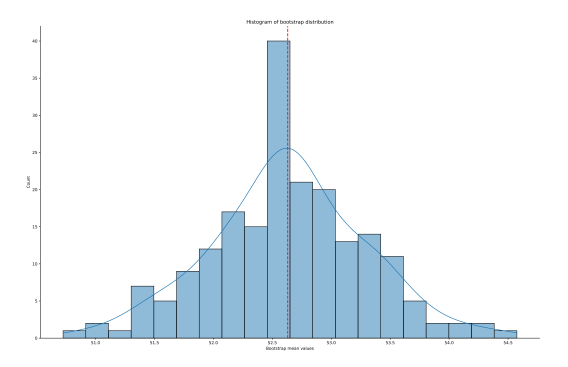
```
## Compute and plot the one sample bootstrap distribution of the mean

def bootstrap_statistic(x, b, statistic):
    n_samps = len(x)
    boot_vals = []
    for _ in range(b):
        boot_vals.append(statistic(nr.choice(x, size=n_samps, replace=True)))
    boot_estimate = np.mean(boot_vals)
    print('Bootstap point estimate = {:6.2f}'.format(boot_estimate))
    return(boot_estimate, boot_vals)

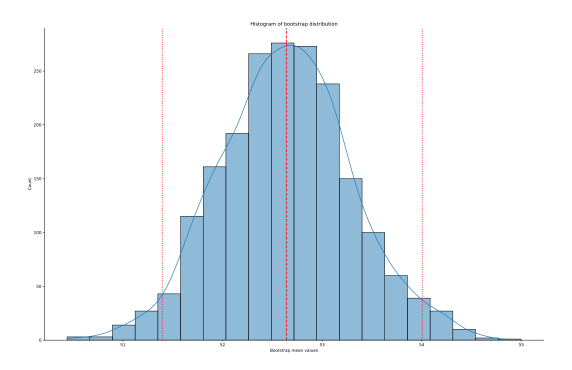
bootstrap_mean_estimate, boot_means = bootstrap_statistic(math, 200, np.mean)
```

```
## Bootstap point estimate = 52.63
```

```
plot_hist(boot_means, 'Bootstrap mean values', 'Histogram of bootstrap distribution')
plt.show()
```



```
## Compute and plot the one sample bootstrap of means with confidence intervals
def bootstrap_cis(boot_samples, alpha=0.05):
      n = len(boot_samples)
      sorted = np.sort(boot_samples)
      index_lci = int(n * alpha / 2)
      index_uci = int(n * (1 - alpha / 2))
      print('At alpha = {0:3.2f}, lower and upper bootstap confidence intervals = {1:6.2f}
                                                                                             {2:6.2f}'.
      return(sorted[index_lci], sorted[index_uci])
bootstrap_mean_estimate, boot_means = bootstrap_statistic(math, 2000, np.mean)
## Bootstap point estimate = 52.64
LCI, UCI = bootstrap_cis(boot_means)
## At alpha = 0.05, lower and upper bootstap confidence intervals = 51.40
plot_hist(boot_means, 'Bootstrap mean values', 'Histogram of bootstrap distribution')
plt.axvline(x=LCI, color='red', linestyle='dotted', linewidth=2)
plt.axvline(x=UCI, color='red', linestyle='dotted', linewidth=2)
plt.show()
```



# Example; two sample bootstrap

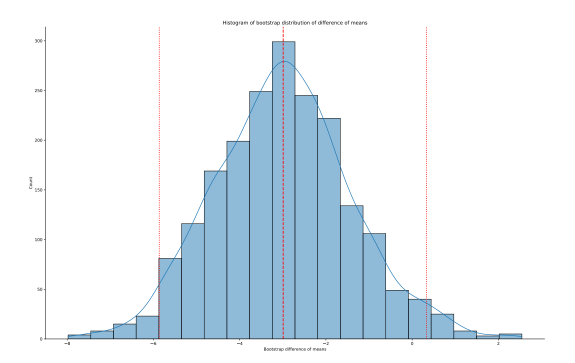
```
# Bootstrap the difference of means of low and mid SES students

def two_boot_two_stat(sample_1, sample_2, b, statistic_1, two_samp_statistic):
    two_boot_values = []
    n_samps_1 = len(sample_1)
    n_samps_2 = len(sample_2)
    for _ in range(b):
        boot_estimate_1 = statistic_1(nr.choice(sample_1, size=n_samps_1, replace=True))
        boot_estimate_2 = statistic_1(nr.choice(sample_2, size=n_samps_2, replace=True))
        two_boot_values.append(two_samp_statistic(boot_estimate_1, boot_estimate_2))
        boot_estimate = np.mean(two_boot_values)
        print('Bootstap point estimate = {:6.2f}'.format(boot_estimate))
        return(boot_estimate, two_boot_values)

math_low_ses = test_scores.loc[test_scores.loc[:,'ses']==1,'math']
math_mid_ses = test_scores.loc[test_scores.loc[:,'ses']==2,'math']
bootstrap_diff_of_mean, boot_diffs = two_boot_two_stat(math_low_ses, math_mid_ses, 2000, np.mean, lambd
```

## Bootstap point estimate = -2.99

```
plot_hist(boot_diffs, 'Bootstrap difference of means', 'Histogram of bootstrap distribution of differen
plt.axvline(x=LCI, color='red', linestyle='dotted', linewidth=2)
plt.axvline(x=UCI, color='red', linestyle='dotted', linewidth=2)
plt.show()
```



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