06 Statistical Functionality in Python

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Information: Some commonly used statistical methods which are useful tools for analyzing data in machine learning

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1 Simple Random Variables

In Python, a common method to define a random variable and sample from it is to use the statistical functionality of scipy.stats.

Note in scipy.stats, drawing random numbers relies on generators from numpy.random package. So we usually only needs to explicit seed a global variable by calling numpy.random.seed() method to ensure reproducibility. Assigning random state is not discussed here since I can not see any necessity of it in the context of this notebook.

For all the attributes in it, read the document of scipy.stats

```
[1]: import scipy.stats as st
import numpy as np
np.random.seed(123)
```

1.1 Discrete Random Variables

1.1.1 The Bernoulli Distribution and basic operations for discrete random variables

Define a Bernoulli distribution by specifying the probability of 1

Call the bernoulli(theta) method

```
[2]: theta = 0.8
X = st.bernoulli(theta)
```

support() method gives which values it takes

```
[3]: print('X takes values in', X.support())
```

X takes values in (0, 1)

pmf() method gives the probability mass function at each point of the support

```
[4]: for x in X.support():
    print('p(X=%d) = %f' % (x, X.pmf(x)))

p(X=0) = 0.200000
p(X=1) = 0.800000
```

expectation() method gives the expectation of the random variable

```
[5]: print('E[X] = %f' % (X.expect()))
```

E[X] = 0.800000

var() method gives the variance of the random variable

```
[6]: print('V[X] = %f' % (X.var()))
```

V[X] = 0.160000

Use rvs(size) method to sample from the random variable with desired shape

```
[7]: xs = X.rvs(size=(4, 3))
print(xs)
```

[[1 1 1]

 $[1 \ 1 \ 1]$

[0 1 1]

[1 1 1]]

Use entropy() method to get the information entropy

```
[8]: print('H[X] = %f'%(X.entropy()))
```

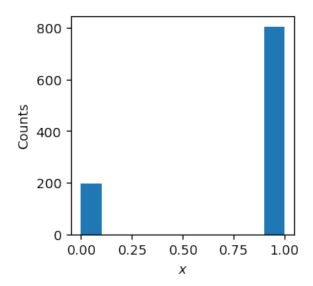
H[X] = 0.500402

Histogram of the samples counts the number of samples taking each value and is one common way to visualize the sampling result.

Use hist() method in matplotlib.pyplot

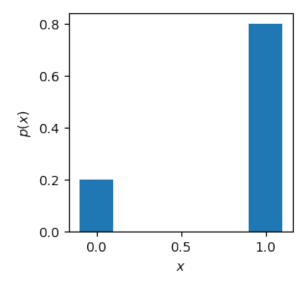
```
[9]: import matplotlib.pyplot as plt

xs = X.rvs(size=1000)
fig, ax = plt.subplots(figsize=(3, 3), dpi=100)
ax.hist(xs)
ax.set_xlabel('$x$')
ax.set_ylabel('Counts')
plt.show()
```



Use bar() method in matplotlib.pyplot to visualize the probability mass function

```
[10]: fig, ax = plt.subplots(figsize=(3, 3), dpi=100)
    ax.bar(x=X.support(), height=X.pmf(X.support()), width=0.2)
    ax.set_xlabel('$x$')
    ax.set_ylabel('$p(x)$')
    plt.show()
```

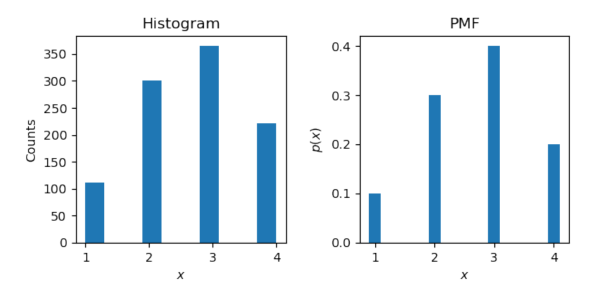


1.1.2 The Categorical Distribution

Specify all the possible values and corresponding probabilities

Call the rv_discrete(name, values=(value, probability)) method

```
[11]: # Define all the possible values
      x_{values} = np.array([1, 2, 3, 4])
      # The probabilities need to sum to 1
      px = np.array([0.1, 0.3, 0.4, 0.2])
      # Define the random variable
      X = st.rv_discrete(name='Custom Categorical', values=(x_values, px))
      # Show the basic properties of it
      print('X takes values in', X.support())
      print('All possible values are', x_values)
      for x in x_values:
          print('p(X=%d) = %f' % (x, X.pmf(x)))
      print('E[X] = %f' % (X.expect()))
      print('V[X] = %f' % (X.var()))
      print('H[X] = %f'%(X.entropy()))
      xs = X.rvs(size=(1000))
      fig, axes = plt.subplots(1, 2, figsize=(7, 3), dpi=100)
      axes[0].hist(xs)
      axes[0].set_xlabel('$x$')
      axes[0].set_ylabel('Counts')
      axes[0].set_title('Histogram')
      axes[1].bar(x=x_values, height=X.pmf(x_values), width=0.2)
      axes[1].set xlabel('$x$')
      axes[1].set_ylabel('$p(x)$')
      axes[1].set title('PMF')
      plt.subplots_adjust(wspace=0.35)
      plt.show()
     X takes values in (1, 4)
     All possible values are [1 2 3 4]
     p(X=1) = 0.100000
     p(X=2) = 0.300000
     p(X=3) = 0.400000
     p(X=4) = 0.200000
     E[X] = 2.700000
     V[X] = 0.810000
     H[X] = 1.279854
```



1.1.3 The Binomial Distribution

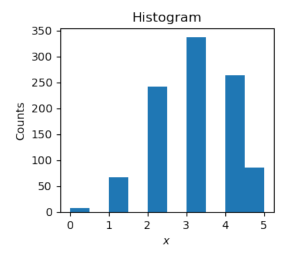
Specify the times of experiments and probability of success of each time

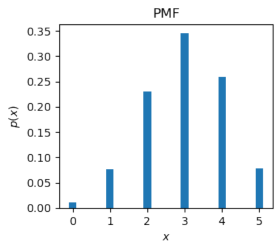
Call the binom() method

```
[12]: # Spcify the times of experiments
      # Specify the probability of success of each time
      theta = 0.6
      # Define the random variable
      X = st.binom(n, theta)
      # Show the basic properties of it
      print('X takes values in', X.support())
      x_range = np.arange(n + 1)
      print('All possible values are', x_range)
      for x in x_range:
          print('p(X=%d) = %f' % (x, X.pmf(x)))
      print('E[X] = %f' % (X.expect()))
      print('V[X] = %f' % (X.var()))
      print('H[X] = %f'%(X.entropy()))
      xs = X.rvs(size=(1000))
      fig, axes = plt.subplots(1, 2, figsize=(8, 3), dpi=100)
      axes[0].hist(xs)
      axes[0].set_xlabel('$x$')
      axes[0].set_ylabel('Counts')
      axes[0].set_title('Histogram')
      axes[1].bar(x=x_range, height=X.pmf(x_range), width=0.2)
      axes[1].set_xlabel('$x$')
```

```
axes[1].set_ylabel('$p(x)$')
axes[1].set_title('PMF')
plt.subplots_adjust(wspace=0.35)
plt.show()
```

```
X takes values in (0, 5)
All possible values are [0 1 2 3 4 5]
p(X=0) = 0.010240
p(X=1) = 0.076800
p(X=2) = 0.230400
p(X=3) = 0.345600
p(X=4) = 0.259200
p(X=5) = 0.077760
E[X] = 3.000000
V[X] = 1.200000
H[X] = 1.497998
```





1.1.4 The Poisson Distribution

Specify the rate with each the events occur

Call possion() method

```
[13]: # The rate with each the events occur
lam = 1.
# Define the random variable
X = st.poisson(lam)
# Show the basic properties of it
x_range = np.arange(6)
for x in x_range:
    print('p(X=%d) = %f' % (x, X.pmf(x)))
```

```
print('E[X] = %f' % (X.expect()))
print('V[X] = %f' % (X.var()))
print('H[X] = %f'%(X.entropy()))
xs = X.rvs(size=(1000))
fig, axes = plt.subplots(1, 2, figsize=(8, 3), dpi=100)
axes[0].hist(xs)
axes[0].set_xlabel('$x$')
axes[0].set_ylabel('Counts')
axes[0].set_title('Histogram')
axes[1].bar(x=x_range, height=X.pmf(x_range), width=0.2)
axes[1].set_xlabel('$x$')
axes[1].set_ylabel('$p(x)$')
axes[1].set_title('PMF')
plt.subplots_adjust(wspace=0.35)
plt.show()
```

```
p(X=0) = 0.367879

p(X=1) = 0.367879

p(X=2) = 0.183940

p(X=3) = 0.061313

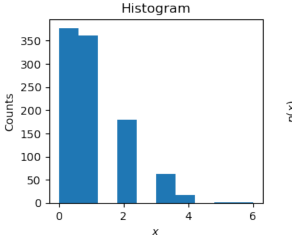
p(X=4) = 0.015328

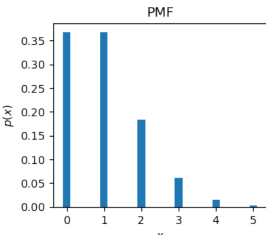
p(X=5) = 0.003066

E[X] = 1.000000

V[X] = 1.000000

H[X] = 1.304842
```





1.2 Continuous Random Variables

1.2.1 The Uniform Distribution and basic operations for continuous random variables

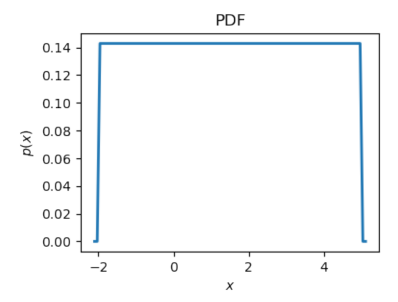
Specify the interval [a, b] where the random variable is distributed uniformly

Call the uniform(loc, scale) method

```
[14]: # Left bound
a = -2.0
# Right bound
b = 5.0
# Define the random variable
X = st.uniform(loc=a, scale=(b - a))
```

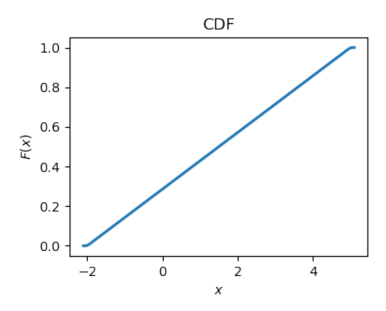
Use pdf() method to get the probability density function

```
[15]: fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
    x = np.linspace(a - 0.1, b + 0.1, 100)
    ax.plot(x, X.pdf(x), lw=2)
    ax.set_xlabel('$x$')
    ax.set_ylabel('$p(x)$')
    ax.set_title('PDF')
    plt.show()
```

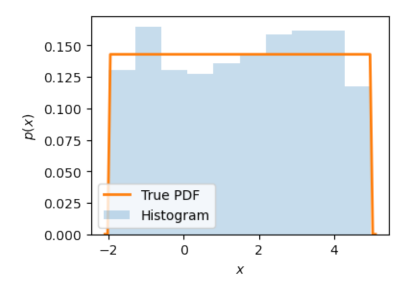


Use cdf() method to get the cumulative distribution function

```
[16]: fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
x = np.linspace(a - 0.1, b + 0.1, 100)
ax.plot(x, X.cdf(x), lw=2)
ax.set_xlabel('$x$')
ax.set_ylabel('$F(x)$')
ax.set_title('CDF')
plt.show()
```



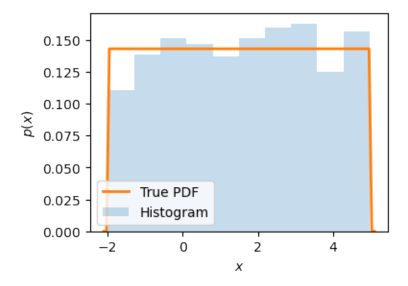
Other properties can be got use similar methods as for discrete random variables



We can also obtain $X \sim U([a,b])$ from $Z \sim U[(0,1)]$ by

$$X = a + (b - a)Z$$

```
[18]: x = np.linspace(a - 0.1, b + 0.1, 100)
# np.random.rand gives the uniform distribution over [0,1]
xs = a + (b - a) * np.random.rand(1000)
fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
ax.hist(xs, density=True, alpha=0.25, label='Histogram')
ax.plot(x, X.pdf(x), lw=2, label='True PDF')
ax.set_xlabel('$x$')
ax.set_ylabel('$p(x)$')
ax.legend(loc='best')
plt.show()
```



1.2.2 The Normal Distribution

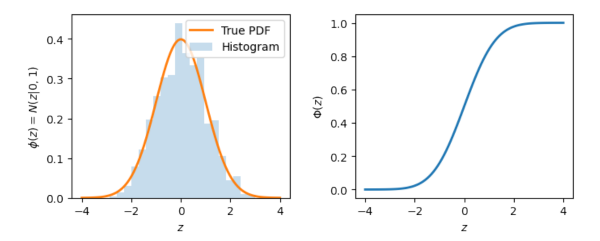
Simple call norm() defines the standard normal distribution

```
[19]: # Define the random variable
      Z = st.norm()
      # Show some basic properties
      print('E[X] = %f' % (Z.expect()))
      print('V[X] = %f' % (Z.var()))
      print('H[X] = %f'%(X.entropy()))
      z = np.linspace(-4, 4, 100)
      zs = Z.rvs(size=(1000))
      fig, axes = plt.subplots(1, 2, figsize=(8, 3), dpi=100)
      axes[0].hist(zs, bins=20, density=True, alpha=0.25, label='Histogram')
      axes[0].plot(z, Z.pdf(z), lw=2, label='True PDF')
      axes[0].set_xlabel('$z$')
      axes[0].set_ylabel('\phi(z)=N(z|0,1)$')
      axes[0].legend(loc='best')
      axes[1].plot(z, Z.cdf(z), lw=2)
      axes[1].set_xlabel('$z$')
      axes[1].set_ylabel('$\Phi(z)$')
      plt.subplots_adjust(wspace=0.3)
      plt.show()
```

E[X] = 0.000000

V[X] = 1.000000

H[X] = 1.945910



Use ppf(q/100) to get the q% quantile z_q

```
[20]: z_025 = Z.ppf(2.5 / 100.0)
z_500 = Z.ppf(50.0 / 100.0)
z_975 = Z.ppf(97.5 / 100.0)
print('2.5% quantile of Z = %1.2f' % (z_025))
print('50% quantile of Z = %1.2f' % z_500)
print('97.5% quantile of Z = %1.2f' % z_975)
```

2.5% quantile of Z = -1.9650% quantile of Z = 0.0097.5% quantile of Z = 1.96

Define any normal by specifying the mean μ and the standard deviation σ :

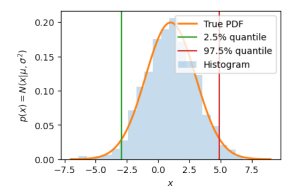
$$X \sim \mathcal{N}(\mu, \sigma^2)$$

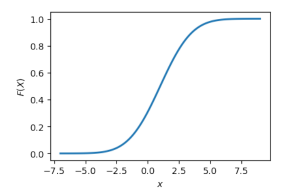
```
[21]: # Mean
      mu = 1
      # Standard deviation
      sigma = 2
      # Define the random variable
      X = st.norm(mu, sigma)
      # Basic properties
      print('E[X] = %f' % (X.expect()))
      print('V[X] = %f' % (X.var()))
      print('H[X] = %f'%(X.entropy()))
      x = np.linspace(mu - 8, mu + 8, 100)
      xs = X.rvs(size=(1000))
      fig, axes = plt.subplots(1, 2, figsize=(10, 3), dpi=100)
      axes[0].hist(xs, bins=20, density=True, alpha=0.25, label='Histogram')
      axes[0].plot(x, X.pdf(x), lw=2, label='True PDF')
      axes[0].set_xlabel('$x$')
```

```
axes[0].set_ylabel('$p(x)=N(x|\mu,\sigma^2)$')
x_025 = X.ppf(2.5 / 100)
x_975 = X.ppf(97.5 / 100)
axes[0].plot(x_025 * np.ones(2), axes[0].set_ylim(), label='2.5% quantile')
axes[0].plot(x_975 * np.ones(2), axes[0].set_ylim(), label='97.5% quantile')
axes[0].legend(loc='best')
axes[0].legend(loc='best')
axes[1].plot(x, X.cdf(x), lw=2)
axes[1].set_xlabel('$x$')
axes[1].set_ylabel('$F(X)$')
plt.subplots_adjust(wspace=0.3)
plt.show()
```

```
E[X] = 1.000000
V[X] = 4.000000
```

H[X] = 2.112086

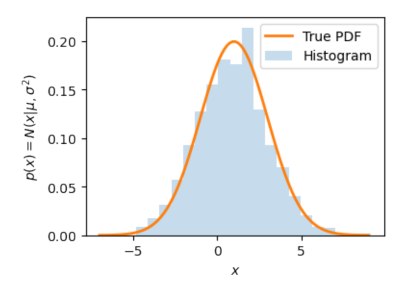




Also can get any normal distribution $X \sim \mathcal{N}(\mu, \sigma^2)$ using the standard normal distribution $Z \sim \mathcal{N}(0, 1)$:

$$X = \mu + \sigma Z$$

```
[22]: Z = st.norm()
mu, sigma = 1, 2
X = st.norm(mu, sigma)
x = np.linspace(mu - 8, mu + 8, 100)
xs = mu + sigma * Z.rvs(size=(1000))
fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
ax.hist(xs, bins=20, density=True, alpha=0.25, label='Histogram')
ax.plot(x, X.pdf(x), lw=2, label='True PDF')
ax.set_xlabel('$x$')
ax.set_ylabel('$p(x)=N(x|\mu,\sigma^2)$')
ax.legend(loc='best')
plt.subplots_adjust(wspace=0.3)
plt.show()
```



2 Collections of Random Variables

Consider the joint probability mass function of two discrete random variables

Take two discrete random variables X and Y. Say X takes m values $0, 1, \dots, m-1$ and Y takes n values $0, 1, \dots, n-1$. Then the joint probability mass function of X and Y can be viewed as the $m \times n$ matrix:

$$A_{i,j} = p(X = i, Y = j)$$

```
[23]: # Take m = 4, n = 3 as an example
A = np.random.rand(4, 3)
# Scale the matrix so that the elements sum to 1
A = A / np.sum(A)
print(A)
print('Sum of elements in A is: %1.2f' % np.sum(A))
```

```
[[0.11736006 0.09213332 0.12528823]

[0.12411247 0.06798429 0.05991044]

[0.03254855 0.09134394 0.11771173]

[0.09943759 0.05207599 0.02009339]]

Sum of elements in A is: 1.00
```

Sum over the second axis gets the marginal distribution of X:

$$p(x) = \sum_{y} p(x, y)$$

```
[24]: p_x = np.sum(A, axis=1)
print('Marginal Distribution of X is:', p_x)
print('Sum of marginal probabilities is %f' % np.sum(p_x))
```

Marginal Distribution of X is: [0.33478161 0.25200721 0.24160421 0.17160698] Sum of marginal probabilities is 1.000000

Find the expectation of X:

$$\mathbb{E}[X] = \sum_{x} x p(x)$$

```
[25]: E_X = np.sum(np.arange(4) * p_x)
print('E[X] = %1.2f' % E_X)
```

E[X] = 1.25

Find the variance of X:

$$\mathbb{V}[X] = \mathbb{E}[(X - \mathbb{E}[X])^2] = \mathbb{E}[X^2] - (\mathbb{E}[X])^2$$

V[X] = 1.20

Do the same for Y

```
[27]: p_y = np.sum(A, axis=0)
    print('Marginal Distribution of Y is:', p_y)
    print('Sum of marginal probabilities is %f' % np.sum(p_y))
    E_Y = np.sum(np.arange(3) * p_y)
    print('E[Y] = %1.2f' % E_Y)
    E_Y2 = np.sum(np.arange(3)**2 * p_y)
    V_Y = E_Y2 - E_Y**2
    print('V[Y] = %1.2f' % V_Y)
```

Marginal Distribution of Y is: $[0.37345868\ 0.30353753\ 0.32300379]$ Sum of marginal probabilities is 1.000000 E[Y] = 0.95V[Y] = 0.69

Find the covariance of the two random variables:

$$\mathbb{C}[X,Y] = \sum_{x,y} (x - \mathbb{E}[X])(y - \mathbb{E}[Y])p(x,y)$$

```
[28]: C_XY = 0.0
for x in range(4):
    for y in range(3):
        C_XY += (x - E_X) * (y - E_Y) * A[x, y]
print('C[X,Y] = %1.2f' % C_XY)
```

$$C[X,Y] = -0.07$$

To sample from (X,Y), we need to sample a set of index (i,j) with probability $A_{i,j}$. It's like sampling from a categorical distribution with $m \times n = mn$ different labels $c_0 = (0,0), c_1 = (0,1), \dots, c_{mn-1} = (m-1,n-1)$ with each a probability $A_{0,0}, A_{0,1}, \dots, A_{m-1,n-1}$

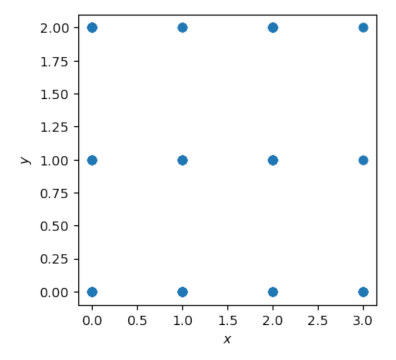
In practice, sample a label c_k and then turn in label to i and j by

$$i = k//n$$
$$j = k \mod n$$

```
[29]: # Define the random variable
XY = st.rv_discrete('Joint XY', values=(np.arange(12), A.reshape(12)))

# The function to sample c_k and transform label k to (i,j)
def sample_XY():
    """
        Samples X and Y once
    """
        k = XY.rvs()
        i = k // 3
        j = k % 3
```

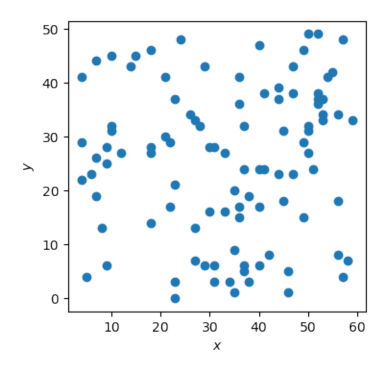
```
# Sample from XY
num_samples = 100
x_samples = np.zeros((num_samples), dtype=int)
y_samples = np.zeros((num_samples), dtype=int)
for ii in range(num_samples):
    x_samples[ii], y_samples[ii] = sample_XY()
# Visualize the sampling result
fig, ax = plt.subplots(figsize=(4, 4), dpi=100)
ax.scatter(x_samples, y_samples)
ax.set_xlabel('$x$')
ax.set_ylabel('$y$')
plt.show()
```



Try for a larger distribution where m = 60, n = 50

```
[30]: m, n = 60, 50
A = np.random.rand(m, n)
A = A / np.sum(A)
p_x = np.sum(A, axis=1)
E_X = np.sum(np.arange(m) * p_x)
print('E[X] = %1.2f' % E_X)
E_X2 = np.sum(np.arange(m)**2 * p_x)
```

```
V_X = E_X2 - E_X**2
      print('V[X] = %1.2f' % V_X)
      p_y = np.sum(A, axis=0)
      E_Y = np.sum(np.arange(n) * p_y)
      print('E[Y] = %1.2f' \% E_Y)
      E_Y2 = np.sum(np.arange(n)**2 * p_y)
      V_Y = E_Y2 - E_Y**2
      print('V[Y] = %1.2f' % V_Y)
      C XY = 0.0
      for x in range(m):
          for y in range(n):
              C_XY += (x - E_X) * (y - E_Y) * A[x, y]
      print('C[X,Y] = %1.2f' % C_XY)
     E[X] = 29.64
     V[X] = 303.68
     E[Y] = 24.11
     V[Y] = 209.01
     C[X,Y] = 2.90
[31]: # Define the random variable
      XY = st.rv_discrete('Joint XY', values=(np.arange(m * n), A.reshape(-1)))
      # The function to sample c k and transform label k to (i,j)
      def sample_XY():
          11 11 11
          Samples X and Y once
          k = XY.rvs()
          i = k // n
          j = k \% n
          return i, j
      # Sample from XY
      num_samples = 100
      x_samples = np.zeros((num_samples), dtype=int)
      y_samples = np.zeros((num_samples), dtype=int)
      for ii in range(num_samples):
          x_samples[ii], y_samples[ii] = sample_XY()
      # Visualize the sampling result
      fig, ax = plt.subplots(figsize=(4, 4), dpi=100)
      ax.scatter(x_samples, y_samples)
      ax.set_xlabel('$x$')
      ax.set_ylabel('$y$')
      plt.show()
```



3 The Multivariate Normal

3.1 Diagonal Covariance Case

To define the N-dimensional multivariate normal with diagonal covariance

$$\mathbf{X} \sim \mathcal{N}(\boldsymbol{\mu}, \operatorname{diag}(\sigma_1^2, \cdots, \sigma_n^2))$$

where μ is an N-dimensional vector and σ_i are positive numbers, call the multivariate_normal(mean=mu, cov=Sigma) method

```
[32]: # Define the mean vector
mu = [1.0, 2.0]
# Define the variance of each component
sigma2 = np.array([1.5, 0.8])**2
# Build the covariance matrix
Sigma = np.diag(sigma2)
# Define the random variable
X = st.multivariate_normal(mean=mu, cov=Sigma)
```

Take a sample from it

```
[33]: X.rvs()
```

[33]: array([-0.79612333, 1.1444333])

Evaluate the PDF at one point

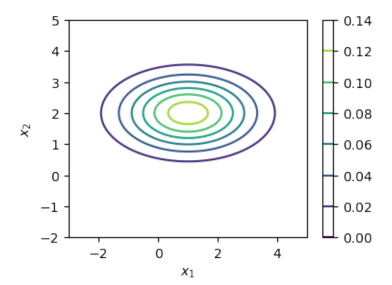
```
[34]: X.pdf([0.5, -1.0])
```

[34]: 0.0001108864139020332

Plot a contour to visualize the PDF

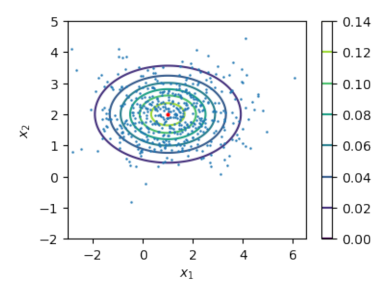
```
[35]: # Points along x1 direction
      x1 = np.linspace(mu[0] - 4, mu[0] + 4, 100)
      # Points alone x2 direction
      x2 = np.linspace(mu[1] - 4, mu[0] + 4, 100)
      # Create the grid
      X1, X2 = np.meshgrid(x1, x2)
      # Reshape these two grids into location lists
      X_{loc} = np.hstack([X1.reshape(-1, 1), X2.reshape(-1, 1)])
      # Evalute the PDF at all there locations
      p_x = X.pdf(X_{loc})
      # Reshape p_x to grid-shape
      P_X = p_x.reshape(100, 100)
      # Plot the contours
      fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
      c = ax.contour(X1, X2, P_X)
      ax.set_xlabel('$x_1$')
      ax.set_ylabel('$x_2$')
```

```
fig.colorbar(mappable=c, ax=ax)
plt.show()
```



Take some samples from X and add them in the contour figure

```
[36]: num_samples = 500
    x_samples = X.rvs(size=num_samples)
    fig, ax = plt.subplots(figsize=(4, 3), dpi=100)
    ax.plot(x_samples[:, 0], x_samples[:, 1], '.', markersize=1.5)
# Mark the mean in red
    ax.plot(X.mean[0], X.mean[1], 'ro', markersize=2)
    c = ax.contour(X1, X2, P_X)
    ax.set_xlabel('$x_1$')
    ax.set_ylabel('$x_2$')
    fig.colorbar(mappable=c, ax=ax)
    plt.show()
```



3.2 Full Covariance Case

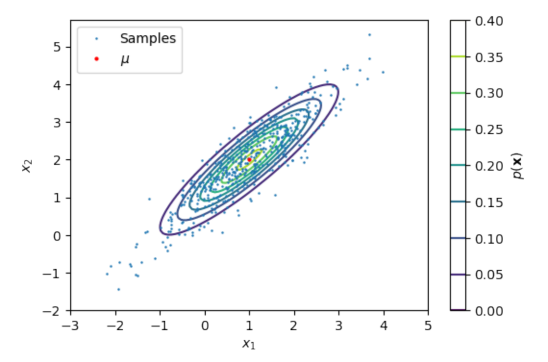
To define the N-dimensional multivariate normal

$$\mathbf{X} \sim \mathcal{N}(oldsymbol{\mu}, oldsymbol{\Sigma})$$

where μ is an N-dimensional vector and Σ is a positive-definite matrix, call the multivariate_normal(mean=mu, cov=Sigma) method

```
[37]: # Define the mean vector
      mu = [1.0, 2.0]
      # Define the covariance matrix
      Sigma = np.array([[1.0, 0.9], [0.9, 1.0]])
      # Define the random variable
      X = st.multivariate_normal(mean=mu, cov=Sigma)
      # Plot the contours
      x1 = np.linspace(mu[0] - 4, mu[0] + 4, 100)
      x2 = np.linspace(mu[1] - 4, mu[0] + 4, 100)
      X1, X2 = np.meshgrid(x1, x2)
      X_loc = np.hstack([X1.reshape(-1, 1), X2.reshape(-1, 1)])
      p_x = X.pdf(X_{loc})
      P_X = p_x.reshape(100, 100)
      fig, ax = plt.subplots(figsize=(6, 4), dpi=100)
      c = ax.contour(X1, X2, P_X)
      ax.set_xlabel('$x_1$')
      ax.set_ylabel('$x_2$')
      fig.colorbar(mappable=c, ax=ax, label='$p(\mathbf{x})$')
      # Sample from X and plot the samples and mean
      num_samples = 500
      x_samples = X.rvs(size=num_samples)
```

```
ax.plot(x_samples[:, 0], x_samples[:, 1], '.', markersize=1.5, label='Samples')
ax.plot(X.mean[0], X.mean[1], 'ro', markersize=2, label='$\mu$')
plt.legend(loc='best')
plt.show()
```



Notice the covariance matrix must be positive definite. This can be checked by calling numpy.linalg.eigh() method

```
[38]: # A positive definite matrix
Sigma_good = np.array([[1.0, 0.9], [0.9, 1.0]])
# A non-positive definite matrix
Sigma_bad = np.array([[0.4, 0.9], [0.9, 1.0]])
# Check their eigenvalues
print('Eigenvalues of Sigma_good is', np.linalg.eigh(Sigma_good)[0])
print('Eigenvalues of Sigma_bad is', np.linalg.eigh(Sigma_bad)[0])
```

Eigenvalues of Sigma_good is [0.1 1.9]
Eigenvalues of Sigma_bad is [-0.2486833 1.6486833]

If we would like to get the N-dimensional multivariate normal

$$\mathbf{X} \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$$

from an N-dimensional standard normal distribution

$$Z \sim \mathcal{N}(0, I)$$

by

$$X = \mu + AZ$$

where

$$\mathbf{\Sigma} = \mathbf{A}\mathbf{A}^T$$

Call numpy.linalg.cholesky() method to decompose the matrix Σ

```
[39]: Sigma = np.array([[1.0, 0.9], [0.9, 1.0]])
A = np.linalg.cholesky(Sigma)
Sigma == np.dot(A, A.T)
```

```
[39]: array([[ True, True], [ True, True]])
```

3.3 Marginalization

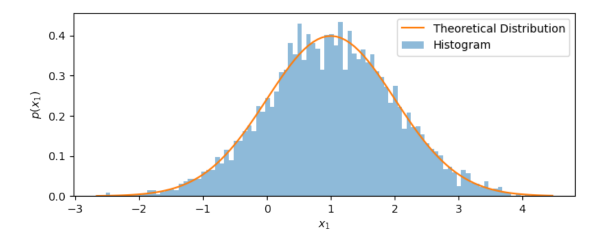
Consider the N-dimensional multivariate normal

$$\mathbf{X} \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$$

where μ is an N-dimensional vector and Σ is a positive-definite matrix. If we marginalize X_i , it will follow the normal distribution

$$\mathcal{N} \sim (\mu_i, \Sigma_{i,i})$$

```
[40]: # Define the mean vector
      mu = [1.0, 2.0]
      # Define the covariance matrix
      Sigma = np.array([[1.0, 0.9], [0.9, 1.0]])
      # Define the random variable
      X = st.multivariate_normal(mean=mu, cov=Sigma)
      # Sample from X
      num samples = 5000
      x samples = X.rvs(size=num samples)
      # Marginalize along X1
      x1_samples = x_samples[:, 0]
      # Plot the histogram
      fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
      ax.hist(x1_samples, density=True, bins=100, alpha=0.5, label='Histogram')
      # Define the theoretical distribution
      X1_theory = st.norm(loc=mu[0], scale=np.sqrt(Sigma[0, 0]))
      x1 = np.linspace(x1_samples.min(), x1_samples.max(), 100)
      ax.plot(x1, X1_theory.pdf(x1), label='Theoretical Distribution')
      ax.set_xlabel('$x_1$')
      ax.set_ylabel('$p(x_1)$')
      plt.legend(loc='best')
      plt.show()
```



3.4 Conditioning

Consider the N-dimensional multivariate normal

$$\mathbf{X} \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$$

where μ is an N-dimensional vector and Σ is a positive-definite matrix.

Assume that μ can be decomposed in two blocks of dimension N_1 and N_2 $(N_1 + N_2 = N)$:

$$oldsymbol{\mu} = egin{bmatrix} oldsymbol{\mu}_1 \ oldsymbol{\mu}_2 \end{bmatrix}$$

Similarly for Σ :

$$oldsymbol{\Sigma} = egin{bmatrix} oldsymbol{\Sigma}_{1,1} & oldsymbol{\Sigma}_{1,2} \ oldsymbol{\Sigma}_{1,2}^T & oldsymbol{\Sigma}_{1,1} \end{bmatrix}$$

The conditional PDF is

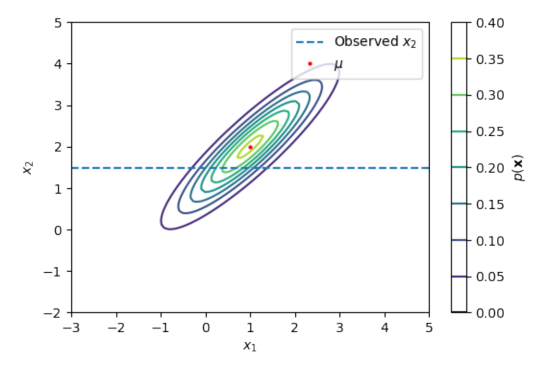
$$p(\mathbf{x}_1|\mathbf{x}_2) = \frac{p(\mathbf{x}_1, \mathbf{x}_2)}{p(\mathbf{x}_2)} \propto p(\mathbf{x}_1, \mathbf{x}_2) = \mathcal{N}(\mathbf{x}_1|\boldsymbol{\mu}_{1|\mathbf{x}_2}, \boldsymbol{\Sigma}_{1,1|\mathbf{x}_2})$$

Where

$$\boldsymbol{\mu}_{1|\mathbf{x}_2} = \boldsymbol{\mu}_1 + \boldsymbol{\Sigma}_{1,2} \boldsymbol{\Sigma}_{2,2}^{-1} (\mathbf{x}_2 - \boldsymbol{\mu}_2), \ \boldsymbol{\Sigma}_{1,1|\mathbf{x}_2} = \boldsymbol{\Sigma}_{1,1} - \boldsymbol{\Sigma}_{1,2} \boldsymbol{\Sigma}_{2,2}^{-1} \boldsymbol{\Sigma}_{1,2}^T$$

```
[41]: # Define the mean vector
mu = [1.0, 2.0]
# Define the covariance matrix
Sigma = np.array([[1.0, 0.9], [0.9, 1.0]])
# Define the random variable
X = st.multivariate_normal(mean=mu, cov=Sigma)
# Observe x2 component of a sample of X
x_sample = X.rvs()
x2_observed = x_sample[1]
```

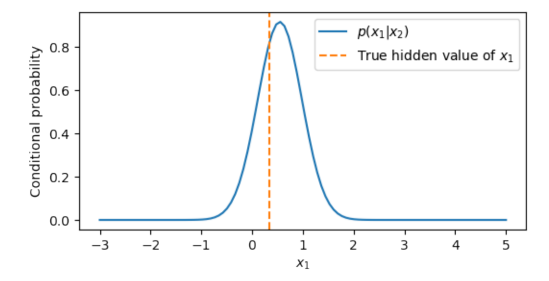
```
x1_hidden = x_sample[0]
# Plot the contours
x1 = np.linspace(mu[0] - 4, mu[0] + 4, 100)
x2 = np.linspace(mu[1] - 4, mu[0] + 4, 100)
X1, X2 = np.meshgrid(x1, x2)
X_loc = np.hstack([X1.reshape(-1, 1), X2.reshape(-1, 1)])
p_x = X.pdf(X_loc)
P_X = p_x.reshape(100, 100)
fig, ax = plt.subplots(figsize=(6, 4), dpi=100)
c = ax.contour(X1, X2, P_X)
ax.set_xlabel('$x_1$')
ax.set_ylabel('$x_2$')
fig.colorbar(mappable=c, ax=ax, label='$p(\mathbf{x})$')
# Plot the observed x2 as a line
ax.plot(ax.set_xlim(), x2_observed * np.ones(2), '--', label='Observed $x_2$')
ax.plot(X.mean[0], X.mean[1], 'ro', markersize=2, label='$\mu$')
plt.legend(loc='best')
plt.show()
```



Let's see the theoretical conditional probability distribution

```
[42]: # Devide the matices into blocks
mu1 = mu[0]
mu2 = mu[1]
```

```
Sigma11 = Sigma[0, 0]
Sigma12 = Sigma[0, 1]
Sigma22 = Sigma[1, 1]
# Conditional mean
mu1\_cond = mu1 + Sigma12 * Sigma22**(-1) * (x2\_observed - mu2)
# Conditional variance
Sigma11_cond = Sigma11 - Sigma12 * Sigma22**(-1) * Sigma12
# Plot the theoretical distribution
X1_cond = st.norm(loc=mu1_cond, scale=np.sqrt(Sigma11_cond))
fig, ax = plt.subplots(figsize=(6, 3), dpi=100)
ax.plot(x1, X1_{cond.pdf}(x1), label='p(x_1|x_2)')
# Plot the true value of hidden x1
ax.plot(x1_hidden * np.ones(2),
        ax.set_ylim(),
        1--1,
        label='True hidden value of $x_1$')
ax.set_xlabel('$x_1$')
ax.set_ylabel('Conditional probability')
plt.legend(loc='best')
plt.show()
```



4 Monte Carlo Methods

4.1 Estimating expectations

Based on the **strong law of large numbers**, we can estimate some complicated expectations. For example, we can estimate the integral

$$I = \mathbb{E}[g(X)] = \int g(x)p(x)dx$$

where $X \sim p(x)$ and g(x) is a function of x by

$$\bar{I}_N = \frac{g(X_1) + g(X_2) + \dots + g(X_N)}{N} \stackrel{N \to \infty}{\longrightarrow} I$$
, a.s.

where X_1, X_2, \cdots are independent copies of X

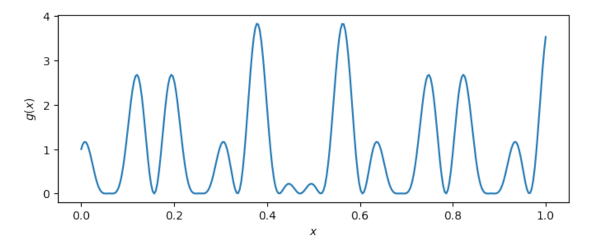
For example, assume $X \sim U([0,1])$ and pick

$$g(x) = (\cos(50x) + \sin(20x))^2$$

The correct value for the expectation can be found analytically and it is

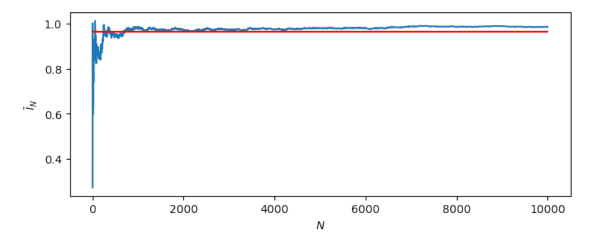
$$\mathbb{E}[g(x)] = 0.965$$

```
[43]: # Define the function
g = lambda x: (np.cos(50 * x) + np.sin(20 * x))**2
# Visualize this complicated function
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
x = np.linspace(0, 1, 300)
ax.plot(x, g(x))
ax.set_xlabel('$x$')
ax.set_ylabel('$g(x)$')
plt.show()
```



Apply the Monte Carlo method for estimating integrals. We can call the numpy.cumsum() to get the cumulative sum of the elements along a given axis.

```
[44]: # Number of samples to take
N = 10000
# Get samples from U([0,1])
x_samples = np.random.rand(N)
# Evaluate the sample average for all sample sizes
I_running = np.cumsum(g(x_samples)) / np.arange(1, N + 1)
# Plot the running estimation
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.plot(np.arange(1, N + 1), I_running)
ax.plot(np.array([1, N + 1]), 0.965 * np.ones(2), 'r')
ax.set_xlabel('$N$')
ax.set_ylabel(r'$\bar{I}_N$')
plt.show()
```



4.2 Estimating variances

Take a random variable $X \sim p(x)$ and some function g(x), to estimate the **variance**:

$$V = \mathbb{V}[g(x)] = \mathbb{E}\left[\left(g(X) - \mathbb{E}[g(X)]\right)^2\right] = \mathbb{E}\left[\left(g(X) - I\right)^2\right] = \mathbb{E}\left[g(x)^2\right] - I^2$$

Take independent identical copies of X:

$$X_1, X_2, \dots \sim p(x)$$

Estimate the mean using a sample average:

$$\bar{I}_N = \frac{1}{N} \sum_{i=1}^{N} g(X_i)$$

Estimate the variance by:

$$\bar{V}_N = \frac{1}{N} \sum_{i=1}^{N} g^2(X_i) - \bar{I}_N^2$$

Take the same example, assume $X \sim U([0,1])$ and pick

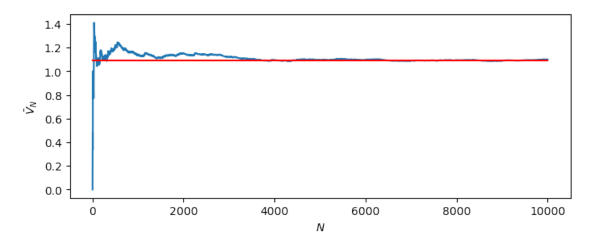
$$g(x) = (\cos(50x) + \sin(20x))^2$$

The correct value for the variance is

$$\mathbb{V}[g(x)] \approx 1.093$$

Apply the Monte Carlo method for estimating variances

```
[45]: # Number of samples to take
N = 10000
# Get samples from U([0,1])
x_samples = np.random.rand(N)
# Evaluate the sample average for all sample sizes
I_running = np.cumsum(g(x_samples)) / np.arange(1, N + 1)
I2_running = np.cumsum(g(x_samples)**2) / np.arange(1, N + 1)
V_running = I2_running - I_running**2
# Plot the running estimation
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.plot(np.arange(1, N + 1), V_running)
ax.plot(np.arange(1, N + 1]), 1.093 * np.ones(2), 'r')
ax.set_xlabel('$N$')
ax.set_ylabel(r'$\bar{V}_N$')
plt.show()
```



4.3 Estimating the CDF

Consider estimating the cumulative distribution function of Y = g(X):

$$F(y) = p(Y \le y) = p(q(X) \le y).$$

It can be written as an expectation using the indicator function:

$$F(y) = \mathbb{E}[1_{[-\infty,y]}(g(X))].$$

It means considering the random variables $1_{-[\infty,y]}(g(X_1)), 1_{-[\infty,y]}(g(X_2)), \ldots$ which are independent and identically distributed. By the strong law of large numbers:

$$\bar{F}_N(y) = \frac{1}{N} \sum_{i=1}^N 1_{[-\infty,y]}(g(X_i)) \to F(y) \text{ a.s..}$$

This estimate is called the empirical CDF.

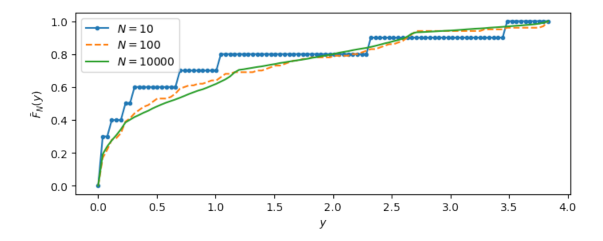
$$\bar{F}_N(y) = \frac{1}{N} \sum_{i=1}^N 1_{[-\infty, y]}(g(X_i)) = \frac{\text{number of } g(X_i) \le y}{N}.$$

Take the same example, assume $X \sim U([0,1])$ and pick

$$g(x) = (\cos(50x) + \sin(20x))^2$$

To calculate the empirical CDF, call the ECDF() method

```
[46]: from statsmodels.distributions.empirical_distribution import ECDF
      # Maximum number of samples to take
      N = 10000
      # Get samples from U([0,1])
      x_samples = np.random.rand(N)
      # Get the corresponding Y samples
      y_samples = g(x_samples)
      # Build ECDF with 10 samples
      ecdf_10 = ECDF(y_samples[:10])
      # Build ECDF with 100 samples
      ecdf 100 = ECDF(y samples[:100])
      # Build ECDF with all samples
      ecdf N = ECDF(y samples)
      # Plot the ECDFs
      fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
      y = np.linspace(y_samples.min(), y_samples.max(), 100)
      ax.plot(y, ecdf_10(y), '.-', label='$N=10$')
      ax.plot(y, ecdf_100(y), '--', label='$N=100$')
      ax.plot(y, ecdf_N(y), label='$N=%d$' % N)
      ax.set_xlabel('$y$')
      ax.set_ylabel(r'$\bar{F}_N(y)$')
      plt.legend(loc='best')
      plt.show()
```



Try to use the empirical CDF to find the probability of that Y takes specific values. For example, find

$$p(1 \le Y \le 3) = F(3) - F(1) \approx \bar{F}_N - \bar{F}_N(1)$$

```
[47]: # Estimate of the probability with 10 samples:

p_Y_10 = ecdf_10(3.0) - ecdf_10(1.0)

print('N = %d:\t\tp(1 <= Y <= 3) ~= %1.2f' % (10, p_Y_10))

# Estimate of the probability with 100 samples:

p_Y_100 = ecdf_100(3.0) - ecdf_100(1.0)

print('N = %d:\tp(1 <= Y <= 3) ~= %1.2f' % (100, p_Y_100))

# Estimate of the probability with all samples:

p_Y_N = ecdf_N(3.0) - ecdf_N(1.0)

print('N = %d:\tp(1 <= Y <= 3) ~= %1.2f' % (N, p_Y_N))
```

N = 10: $p(1 \le Y \le 3) \sim 0.20$ N = 100: $p(1 \le Y \le 3) \sim 0.30$ N = 10000: $p(1 \le Y \le 3) \sim 0.33$

4.4 Estimating the PDF via Histograms

Take a random variable $X \sim p(x)$ and some function g(x), consider the derived random variable Y = g(X), we would like to estimate the **probability density function** p(y) of Y

Take M small bins $[b_0, b_1], \dots, [b_{M-1}, b_M]$ in the y space and approximate p(y) with a constant inside each bin:

$$\bar{p}_M(y) = \sum_{j=1}^{M} c_j 1_{[b_{j-1}, b_j]}(y)$$

The constants c_i are:

$$c_i = P(b_{i-1} \le Y \le b_i) = F(b_i) - F(b_i - 1)$$

So, we can approximate the constants c_i with the empirical CDF:

$$\bar{c}_{j,N} = \bar{F}_N(b_j) - \bar{F}_N(b_{j-1}) = \frac{\text{number of samples that fall in bin } [b_{j-1}, b_j]}{N} \to c_j \text{ a.s.}$$

Putting everything together the approximation becomes:

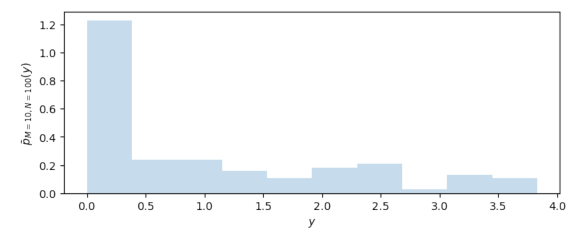
$$\bar{p}_{M,N}(y) = \sum_{j=1}^{M} \bar{c}_{j,N} 1_{[b_{j-1},b_j]}(y)$$

Take the same example, assume $X \sim U([0,1])$ and pick

$$g(x) = (\cos(50x) + \sin(20x))^2$$

In practice, the rule of thumb is to make N about ten times M. First try with M = 10, N = 100

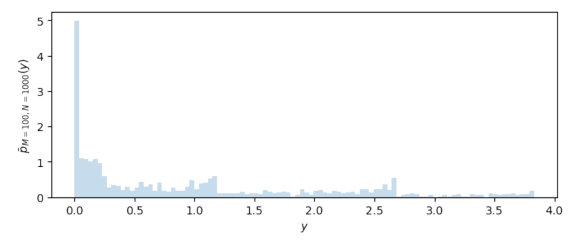
```
[48]: # Maximum number of samples to take
N = 100
# Number of bins
M = 10
# Get samples from U([0,1])
x_samples = np.random.rand(N)
# Get corresponding samples of Y
y_samples = g(x_samples)
# Plot the estimation
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.hist(y_samples, label='$N=%d$' % N, bins=M, density=True, alpha=0.25)
ax.set_xlabel('$y$')
ax.set_ylabel(r'$\bar{p}_{M=%d,N=%d}(y)$' % (M, N))
plt.show()
```



Increase to M = 100, N = 1000

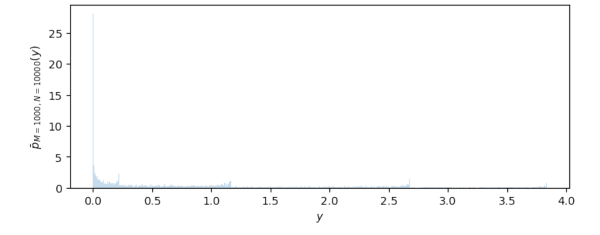
```
[49]: M, N = 100, 1000
x_samples = np.random.rand(N)
y_samples = g(x_samples)
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.hist(y_samples, label='$N=%d$' % N, bins=M, density=True, alpha=0.25)
```

```
ax.set_xlabel('$y$')
ax.set_ylabel(r'$\bar{p}_{M=%d,N=%d}(y)$' % (M, N))
plt.show()
```



Increase to M = 1000, N = 10000

```
[50]: M, N = 1000, 10000
x_samples = np.random.rand(N)
y_samples = g(x_samples)
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.hist(y_samples, label='$N=%d$' % N, bins=M, density=True, alpha=0.25)
ax.set_xlabel('$y$')
ax.set_ylabel(r'$\bar{p}_{M=%d,N=%d}(y)$' % (M, N))
plt.show()
```



4.5 Estimating predictive quantiles

To find the q-predictive quantile of Y:

$$F_Y(\mu_q) = \frac{q}{100}$$

Turn it into a root finding problem:

$$F(\mu_q) - \frac{q}{100} = 0$$

Take X_1, X_2, \cdots independent identical copies of X, fins the sampling estimate of the CDF of Y = g(X) denoted by

$$\bar{F}_N(y)$$

Then numerically solve the root finding problem and obtain the quantile $\bar{\mu}_{q,N}$

Take the same example, assume $X \sim U([0,1])$ and pick

$$g(x) = (\cos(50x) + \sin(20x))^2$$

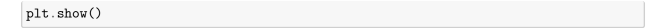
Use **np.quantile()** to solve the root finding problem. Notice we do not call the **ECDF()** method to actually estimate the empirical CDF if we only want the quantiles.

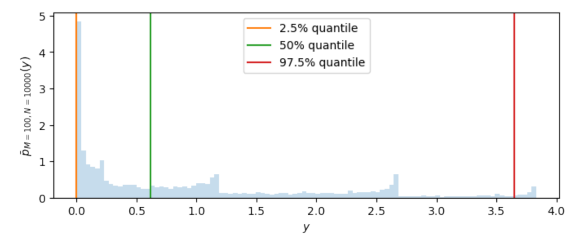
```
[51]: # Maximum number of samples to take
N = 10000
# Get samples from U([0,1])
x_samples = np.random.rand(N)
# Get corresponding samples of Y
y_samples = g(x_samples)
# Find the 2.5%, 50% and 97.5% quantiles
mu_025 = np.quantile(y_samples, 2.5 / 100)
print('mu_025 = %1.2f' % mu_025)
mu_500 = np.quantile(y_samples, 50.0 / 100)
print('mu_500 = %1.2f' % mu_500)
mu_975 = np.quantile(y_samples, 97.5 / 100)
print('mu_975 = %1.2f' % mu_975)
```

mu_025 = 0.00 mu_500 = 0.61 mu_975 = 3.65

Plot the quantiles on the histogram of samples from Y

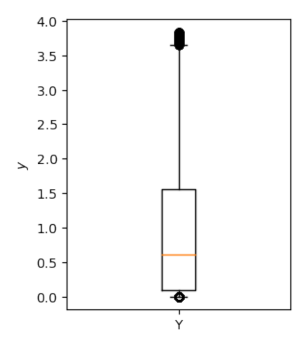
```
[52]: # Here use a relatively samll bin number for better visualization
M = 100
fig, ax = plt.subplots(figsize=(8, 3), dpi=100)
ax.hist(y_samples, bins=M, density=True, alpha=0.25)
ax.set_xlabel('$y$')
ax.set_ylabel(r'$\bar{p}_{M=%d,N=%d}(y)$' % (M, N))
ax.plot(mu_025*np.ones(2),ax.set_ylim(),label='2.5% quantile')
ax.plot(mu_500*np.ones(2),ax.set_ylim(),label='50% quantile')
ax.plot(mu_975*np.ones(2),ax.set_ylim(),label='97.5% quantile')
plt.legend(loc='best')
```





The predictive quantiles are often summarized using box plots by calling the boxplot() method

```
[53]: fig,ax=plt.subplots(figsize=(3,4),dpi=100)
ax.boxplot(y_samples,whis=[2.5,97.5],labels=['Y'])
ax.set_ylabel('$y$')
plt.show()
```



In the box plot: - The y-axis indicates possible values - The orange line indicates the median - The

box encapsulates 50% of the probability around the median - The whiskers are extreme quantiles (here selected to be 2.5% and 97.5% quantiles) - The plot also shows the samples that fall outside the extreme quantiles

4.6 Quantifying epistemic uncertainties in Monte Carlo estimates

Based on the *strong law of large numbers*, we can estimate use sampling to estimate expectations. For example, we can estimate the integral

$$I = \mathbb{E}[g(X)] = \int g(x)p(x)dx$$

where $X \sim p(x)$ and g(x) is a function of x by

$$\bar{I}_N = \frac{g(X_1) + g(X_2) + \dots + g(X_N)}{N} \stackrel{N \to \infty}{\longrightarrow} I$$
, a.s.

where X_1, X_2, \cdots are independent copies of X Also, we can estimate the variance by:

$$\bar{V}_N = \frac{1}{N} \sum_{i=1}^{N} g^2(X_i) - \bar{I}_N^2$$

Assume $Y_1 = g(X_1), Y_2 = g(X_2), \dots, Y_N = g(X_N)$ are i.i.d. random variables with

$$\mathbb{E}[Y_i] = \mathbb{E}[g(X_i)] = I, \mathbb{V}[Y_i] = \sigma^2 \le +\infty$$

According to the *central limit theorem* (details are skipped here and are provided in 02 Probability Theory):

$$I \sim \mathcal{N}(\bar{I}_N, \frac{\bar{\sigma}_N^2}{N})$$
 for large N

In practice, we usually write something like:

$$I \approx \bar{I}_N \pm \frac{1.96}{\sqrt{N}} \bar{\sigma}_N$$

with about 95% probability

```
[54]: # Number of samples to take
N = 1000
# Get samples from U([0,1])
x_samples = np.random.rand(N)
# Get corresponding samples of Y
y_samples = g(x_samples)
# Evaluate the sample average for all sample sizes
I_running = np.cumsum(y_samples) / np.arange(1, N + 1)
I2_running = np.cumsum(y_samples**2) / np.arange(1, N + 1)
V_running = I2_running - I_running**2
# Get the lower bound and upper bound of the 95% condidence interval
I_lb_running = I_running - 1.96 * np.sqrt(V_running/np.arange(1,N+1))
I_ub_running = I_running + 1.96 * np.sqrt(V_running/np.arange(1,N+1))
# Plot all the estimates
```

