Chapter 31-5: Elliptic PDEs Using the Monte Carlo Method.

Monte Carlo Methods are a class of computational solutions that can be applied to various problems which rely on repeated random sampling to provide generally approximate solutions. Monte Carlo is a stochastic approach, in which a series of simulations (trials), representing the analyzed problem, with randomly selected input values, are performed. Among these trials, a specified number of properly defined successes is achieved. The ratio between the number of success trials to the number of all trials, scaled by dimensional quantity (e.g., area or function value) allows for the estimation of the unknown solution, providing the number of trials is large enough.

A capacitor is constructed out of two infinite metal plates spaced 10 cm apart from one another. One plate is placed at 5V potential with respect to the other and the space between the plates is free of charges. The number of random walks is taken to be 400 and the number of lattice points is taken to be 30.

(The description of the problem and the code solution was taken from the repository of s-ankur.)

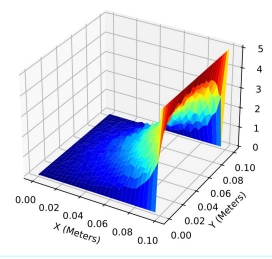
```
In [1]:
           2
             All calculations in SI units
           3
             import random
             from mpl_toolkits.mplot3d import axes3d
              import matplotlib.pyplot as plt
              from matplotlib import cm
           ጸ
             import numpy as np
           9
              %config InlineBackend.figure_formats = ['svg']
          10
          11 h = 10e-2 # Distance between plates = 10cm
             lattice_points = 30 # Number of Points in lattice
          12
          13 d = h / lattice_points # Lattice size
          14 | boundary_voltage_high = 5.0 # 5 Volts at Positive Plate
          boundary_voltage_low = 0.0 # 0 Volts at Negative Plate epsilon_naught = 8.854e-12 # Permittivity of Vaccum
              charge_density = 6e-16 # Coulomb per meter cube
          18
             N = 400 # Number of Random Walks
          19
          20
          21
              def f(x):
          22
23
                  # The Function \nabla ^2(phi) = f
                  # For Laplace f = 0
          24
                  return 0
          25
          26
          27
              def q(x):
          28
                  # Two Dimensional Boundary Conditions: two parallel metal plates at x=0,x=h
          29
                  # the plate at x=h is at high potential and x=0 is low potential
          30
                  # Assume that there are metal plates along y=0 and y=h (uncharged)
          31
                  # this is because I dont know how to simulate open boundry conditions
          32
                  if x[0] <= 0:
          33
                       return boundary_voltage_low
          34
                  if x[0] >= h:
          35
                       return boundary_voltage_high
          36
                  if x[1] \leftarrow 0 or x[1] >= h:
          37
                       return boundary_voltage_low
          38
              0.00
          39
             def f_2(x):
          40
                  # Alternative charge distribution: A charged Sphere in the centre of metal box if (h / 2 - \times[0]) ** 2 + (h / 2 - \times[1]) ** 2 <= (h / 5) ** 2:
          41
          42
                       return -charge_density * 5 / epsilon_naught
          43
          44
                  else:
          45
                       return 0
          46
          47
          48
              def g_2(x):
          49
                  # Two Dimentional Alternative Boundary Conditions: uncharged metal box
          50
                  return 0
          51
          52
          53
              def f_3(x):
```

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```
# Alternative charge distribution: TWO charged Sphere in the centre of metal box if (h / 3 - x[0]) ** 2 + (h / 2 - x[1]) ** 2 <= (h / 5) ** 2:
55
 56
         return -charge_density * 5 / epsilon_naught if (2 * h / 3 - x[0]) ** 2 + (h / 2 - x[1]) ** 2 <= (h / 5) ** 2:
 57
58
 59
             return charge_density * 5 / epsilon_naught
 60
         else:
61
             return 0
62
63
    @np.vectorize
 64
65 def poisson_approximation_fixed_step(*A):
66
         # Returns the Value of Potential Feild at a given point A with N random walks
67
         result = 0
         F = 0
68
 69
         for i in range(N):
 70
             x = list(A)
71
             while True:
 72
                  if x[0] \leftarrow 0 or x[0] \rightarrow h or x[1] \leftarrow 0 or x[1] \rightarrow h:
 73
                      break
 74
                  random\_number = random.randint(0, 3)
 75
                  if random_number == 0:
 76
                      x[0] += d
 77
                  elif random_number == 1:
 78
                      x[0] -= d
 79
                  elif random_number == 2:
 80
                      x[1] += d
 81
                  elif random_number == 3:
82
                      x[1] -= d
                  F += \bar{f}(\bar{x}) * h ** 2
 83
 84
             result += g(x) / N
 85
         result = result - F
 86
         return result
87
88
   def plot(x, y, z):
89
90
         # Function for plotting the potential
91
         fig = plt.figure()
92
         ax = fig.add_subplot(111, projection="3d")
93
94
         ax.plot_surface(x, y, np.array(z), cmap=cm.jet, linewidth=0.1)
         plt.xlabel("X (Meters)")
plt.ylabel("Y (Meters)")
95
96
97
         ax.set_zlabel("Potential (Volts)")
98
         plt.show()
99
100
101 if __name__ == "__main__":
102
         # Experiment E: 2D Capacitor
103
         print(
104
             f"Calculating Monte Carlo with {lattice_points}x{lattice_points} lattice points and {N} random wa
105
106
         lattice_x, lattice_y = np.mgrid[
107
             0 : h : lattice_points * 1j, 0 : h : lattice_points * 1j
108
109
         z = poisson_approximation_fixed_step(lattice_x.ravel(), lattice_y.ravel()).reshape(
110
              lattice_x.shape
111
112
         plot(lattice_x, lattice_y, z)
113
114
         # Experiment F: Metal box with positively charged metal ball inside
115
          f = f2
      #
116
      #
          g = g2
         print(
117
118
             f"Calculating Monte Carlo with {lattice_points}x{lattice_points} lattice points and {N} random wa
119
         lattice_x, lattice_y = np.mgrid[
120
121
             0 : h : lattice_points * 1j, 0 : h : lattice_points * 1j
122
123
         z = poisson\_approximation\_fixed\_step(lattice\_x.ravel(), lattice\_y.ravel()).reshape(
124
              lattice_x.shape
125
126
         plot(lattice_x, lattice_y, z)
127
128
         # Experiment G: Metal Box with two spheres (positive and negative)
129
130
         f = f_3
131
         g = g_2
         print(
132
133
             f"Calculating Monte Carlo with {lattice_points}x{lattice_points} lattice points and {N} random wa
134
```

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```
135
        lattice_x, lattice_y = np.mgrid[
             0 : h : lattice_points * 1j, 0 : h : lattice_points * 1j
136
137
        z = poisson_approximation_fixed_step(lattice_x.ravel(), lattice_y.ravel()).reshape(
138
139
             lattice_x.shape
140
        )
141
        plot(lattice_x, lattice_y, z)
^{142} Calculating Monte Carlo with 30x30 lattice points and 400 random walks
```



```
Traceback (most recent call last)
 ~\AppData\Local\Temp/ipykernel_11260/807967070.py in <module>
                         116 # g = g2
                          117
                                                                            print(
                                                                                                           f"Calculating Monte Carlo with \{lattice\_points\}x\{lattice\_points\} \ lattice\_points \ and \ \{N\} \ rather than the points \ and \ rather than the points \ lattice\_points \ latt
 --> 118
ndom walks for {'Laplace' if laplace else 'Poisson'}"
                        119
                                                                              lattice_x, lattice_y = np.mgrid[
                        120
NameError: name 'laplace' is not defined
```

```
In []:
```

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