# ECSE 543 Assignment 1

Sean Stappas 260639512

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# Introduction

The programs for this assignment were created in Python 2.7. The source code is provided as listings in Appendix A. To perform the required tasks in this assignment, a custom matrix package was created, with useful methods such as add, multiply, transpose, etc. This package can be seen in Listing 1. In addition, logs of the output of the programs are provided in Appendix B.

# 1 Choleski Decomposition

The source code for the Question 1 main program can be seen in Listing 4.

# 1.a Choleski Program

The code relating specifically to Choleski decomposition can be seen in Listing 2.

### 1.b Constructing Test Matrices

The matrices were constructed with the knowledge that, if A is positive-definite, then  $A = LL^T$  where L is a lower triangular non-singular matrix. The task of choosing valid A matrices then boils down to finding non-singular lower triangular L matrices. To ensure that L is non-singular, one must simply choose nonzero values for the main diagonal.

# 1.c Test Runs

The matrices were tested by inventing x matrices, and checking that the program solves for that x correctly. The output of the program, comparing expected and obtained values of x, can be seen in Listing 8.

#### 1.d Linear Networks

can be seen in Listing 3. the csv\_to\_network\_branch\_matrices method the linear\_networks.py script reads from a CSV file where row k contains  $J_k$ ,  $R_k$  and  $E_k$ . It then converts the resistances to a diagonal admittance matrix Y and produces the J and E column The incidence matrix A is also read directly from file, as seen in Listing 4.

First, the program was tested on the circuits provided on MyCourses. These circuits are labeled 1 to 5 and have corresponding incidence matrix and network branch CSV files, located in the network\_data directory. The program obtains the expected voltages, as seen in the output in Listing 8.

Then, some additional simple test circuits were created. Circuit 6 can be seen in Figure 1 and the SPICE analysis output in Table 1. These voltages match the ones calculated by the program, as seen in Listing 8.

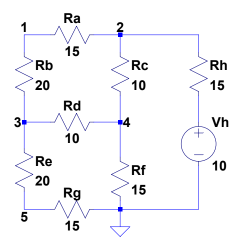


Figure 1: Test circuit 6 with nodes labeled 1 to 4.

Table 1: Output of SPICE operating point analysis of circuit 6.

| Node | Voltage (V) |
|------|-------------|
| 1    | 4.443       |
| 2    | 5.498       |
| 3    | 3.036       |
| 4    | 3.200       |
| 5    | 1.301       |

# 2 Finite Difference Mesh

The source code for the Question 2 main program can be seen in Listing 5.

### 2.a Equivalent Resistance

The code for creating all the network matrices and for finding the equivalent resistance of an N by 2N mesh can be seen in Listing 3. The resistances found by the program for values of N from 2 to 10 can be seen in Table 2.

The resistance values returned by the program for small meshes were validated using simple SPICE circuits. The voltage found at the  $V_{test}$  node for the 2x4 mesh is 1.875 V and the equivalent resistance is therefore 1875  $\Omega$ . Similarly, for the 3x6 mesh,  $V_{test}=2.379\,55\,\mathrm{V}$  and the equivalent resistance is 2379.55  $\Omega$ . These match the results found by the program, as seen in Table 2.

Table 2: Mesh equivalent resistance R versus mesh size N.

| N  | R (Omega) |
|----|-----------|
| 2  | 1875.000  |
| 3  | 2379.545  |
| 4  | 2741.025  |
| 5  | 3022.819  |
| 6  | 3253.676  |
| 7  | 3449.166  |
| 8  | 3618.675  |
| 9  | 3768.291  |
| 10 | 3902.189  |

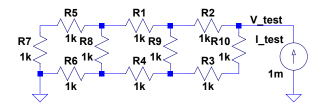


Figure 2: SPICE circuit used to test the 2x4 mesh.

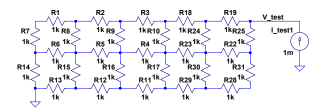


Figure 3: SPICE circuit used to test the 3x6 mesh.

# 2.b Time Complexity

The runtime data for the mesh resistance solver is tabulated in Table 3 and plotted in Figure 4. Theoretically, the time complexity of the program should be  $O(N^6)$ , and this matches the obtained data.

Table 3: Runtime of mesh resistance solver program versus mesh size N.

| N  | Runtime (s) |
|----|-------------|
| 2  | 0.001       |
| 3  | 0.017       |
| 4  | 0.100       |
| 5  | 0.482       |
| 6  | 1.461       |
| 7  | 3.266       |
| 8  | 7.534       |
| 9  | 15.002      |
| 10 | 28.363      |

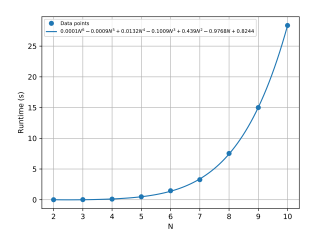


Figure 4: Runtime of mesh resistance solver program versus mesh size N.

# 2.c Sparsity Modification

The runtime data for the banded mesh resistance solver is tabulated in Table 4 and plotted in Figure 5. By inspection of the constructed network matrices, a half-bandwidth of 2N+1 was chosen. Theoretically, the banded version should have a time complexity of  $O(N^4)$ .

Table 4: Runtime of banded mesh resistance solver program versus mesh size N.

| N  | Runtime (s) |
|----|-------------|
| 2  | 0.001       |
| 3  | 0.017       |
| 4  | 0.095       |
| 5  | 0.378       |
| 6  | 1.192       |
| 7  | 3.052       |
| 8  | 6.943       |
| 9  | 14.219      |
| 10 | 26.764      |

The runtime of the banded and non-banded versions of the program are plotted in Figure 6, showing the benefits of banded elimination.

#### 2.d Resistance vs. Mesh Size

The equivalent mesh resistance R is plotted versus the mesh size N in Figure 7. The function R(N) appears logarithmic, and a log function does indeed fit the data well.

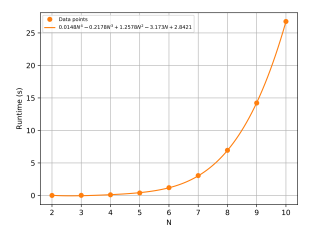


Figure 5: Runtime of banded mesh resistance solver program versus mesh size N.

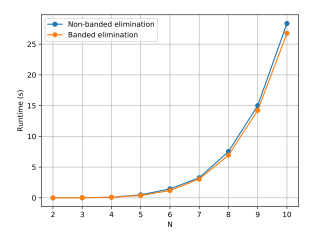


Figure 6: Comparison of runtime of banded and non-banded resistance solver programs versus mesh size N.

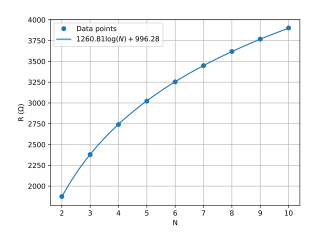


Figure 7: Resistance of mesh versus mesh size N.

# 3.a SOR Program

The source code for the finite difference methods can be seen in Listing 6. Horizontal and vertical symmetries were exploited by only solving for a quarter of the coaxial cable, and reproducing the results where necessary.

# 3.b Varying $\omega$

The number of iterations to achieve convergence for 10 values of  $\omega$  between 1 and 2 are tabulated in Table 5 and plotted in Figure 8. Based on these results, the value of  $\omega$  yielding the minimum number of iterations is 1.3.

Table 5: Number of iterations of SOR versus  $\omega$ .

| Omega | Iterations |
|-------|------------|
| 1.0   | 32         |
| 1.1   | 26         |
| 1.2   | 20         |
| 1.3   | 14         |
| 1.4   | 16         |
| 1.5   | 20         |
| 1.6   | 27         |
| 1.7   | 39         |
| 1.8   | 60         |
| 1.9   | 127        |

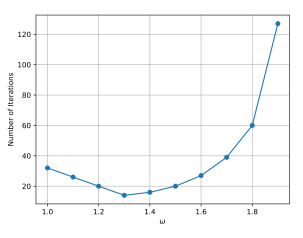


Figure 8: Number of iterations of SOR versus  $\omega$ .

The potential values found at (0.06, 0.04) versus  $\omega$  are tabulated in Table 6. It can be seen that all the potential values are identical to 3 decimal places.

# 3 Coaxial Cable

The source code for the Question 2 main program can be seen in Listing 7.

Table 6: Potential at (0.06, 0.04) versus  $\omega$  when using SOR.

| Omega | Potential (V) |
|-------|---------------|
| 1.0   | 5.526         |
| 1.1   | 5.526         |
| 1.2   | 5.526         |
| 1.3   | 5.526         |
| 1.4   | 5.526         |
| 1.5   | 5.526         |
| 1.6   | 5.526         |
| 1.7   | 5.526         |
| 1.8   | 5.526         |
| 1.9   | 5.526         |

# 3.c Varying h

With  $\omega=1.3$ , the number of iterations of SOR versus 1/h is tabulated in Table 7 and plotted in Figure 9. It can be seen that the smaller the node spacing is, the more iterations the program will take to run. Theoretically, the time complexity of the program should be  $O(N^3)$ , where the finite difference mesh is N by N, and this matches the measured data.

Table 7: Number of iterations of SOR versus 1/h. Note that  $\omega = 1.3$ .

| 1/h    | Iterations |
|--------|------------|
| 50.0   | 14         |
| 100.0  | 59         |
| 200.0  | 189        |
| 400.0  | 552        |
| 800.0  | 1540       |
| 1600.0 | 4507       |

The potential values found at  $(0.06,\ 0.04)$  versus 1/h are tabulated in Table 8 and plotted in Figure 10. By examining these values, the potential at  $(0.06,\ 0.04)$  to three significant figures is approximately  $5.25\,\mathrm{V}$ . It can be seen that the smaller the node spacing is, the more accurate the calculated potential is. However, by inspecting Figure 10 it is apparent that the potential converges relatively quickly to around  $5.25\,\mathrm{V}$  There are therefore diminishing returns to decreasing the node spacing too much, since this will also increase the runtime of the program.

# 3.d Jacobi Method

The number of iterations of the Jacobi method versus 1/h is tabulated in Table 9 and plotted in

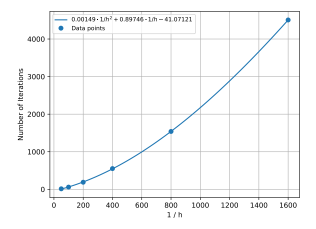


Figure 9: Number of iterations of SOR versus 1/h. Note that  $\omega = 1.3$ .

Table 8: Potential at (0.06, 0.04) versus 1/h when using SOR.

| 1/h    | Potential (V) |
|--------|---------------|
| 50.0   | 5.526         |
| 100.0  | 5.351         |
| 200.0  | 5.289         |
| 400.0  | 5.265         |
| 800.0  | 5.254         |
| 1600.0 | 5.247         |

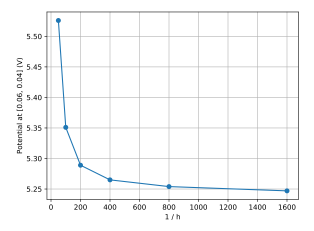


Figure 10: Potential at (0.06, 0.04) found by SOR versus 1/h. Note that  $\omega = 1.3$ .

Figure 11. Similarly to SOR, the smaller the node spacing is, the more iterations the program will take to run. We can see however that the Jacobi method takes a much larger number of iterations to converge. Theoretically, the Jacobi method should have a time complexity of  $O(N^4)$ , and this matches the data.

The potential values found at (0.06, 0.04) ver-

Table 9: Number of iterations versus  $\omega$  when using the Jacobi method.

| 1/h    | Iterations |
|--------|------------|
| 50.0   | 51         |
| 100.0  | 180        |
| 200.0  | 604        |
| 400.0  | 1935       |
| 800.0  | 5836       |
| 1600.0 | 16864      |

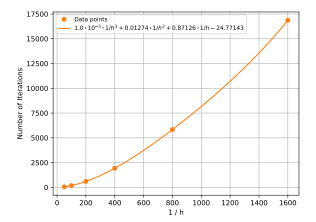


Figure 11: Number of iterations of the Jacobi method versus 1/h.

sus 1/h with the Jacobi method are tabulated in Table 10 and plotted in Figure 12. These potential values are almost identical to the SOR ones. Similarly to SOR, the smaller the node spacing is, the more accurate the calculated potential is.

Table 10: Potential at (0.06, 0.04) versus 1/h when using the Jacobi method.

| 1/h    | Potential (V) |
|--------|---------------|
| 50.0   | 5.526         |
| 100.0  | 5.351         |
| 200.0  | 5.289         |
| 400.0  | 5.265         |
| 800.0  | 5.254         |
| 1600.0 | 5.246         |
|        |               |

The number of iterations of both SOR and the Jacobi method can be seen in Figure 13, which shows the clear benefits of SOR.

# 3.e Non-uniform Node Spacing

First, we adjust the equation derived in class to set  $a_1 = \Delta_x \alpha_1$ ,  $a_2 = \Delta_x \alpha_2$ ,  $b_1 = \Delta_y \beta_1$  and

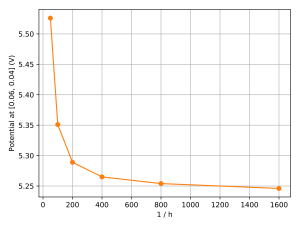


Figure 12: Potential at (0.06, 0.04) versus 1/h when using the Jacobi method.

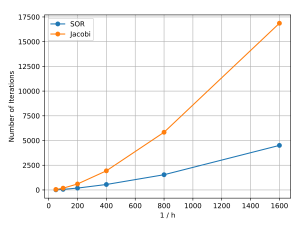


Figure 13: Comparison of number of iterations when using SOR and Jacobi methods versus 1/h. Note that  $\omega = 1.3$  for the SOR program.

 $b_2 = \Delta_y \beta_2$ . These values correspond to the distances between adjacent nodes <sup>1</sup>, and can be easily calculated by the program. Then, the five-point difference formula for non-uniform spacing can be seen in Equation 1.

$$\phi_{i,j}^{k+1} = \frac{1}{a_1 + a_2} \left( \frac{\phi_{i-1,j}^k}{a_1} + \frac{\phi_{i+1,j}^k}{a_2} \right) + \frac{1}{b_1 + b_2} \left( \frac{\phi_{i,j-1}^k}{b_1} + \frac{\phi_{i,j+1}^k}{b_2} \right)$$
(1)

This was implemented in the finite difference program, as seen in Listing 6. As can be seen in this code, many different mesh arrangements were

<sup>&</sup>lt;sup>1</sup>Note that, in the program, index i is associated to position x and index j is associated to position y. This is purely for easier printing of the matrices.

tested. The arrangement that was chosen can be seen in Figure 14. The potential at  $(0.06,\ 0.04)$  obtained from this arrangement is  $5.243\,\mathrm{V}$ , which seems like an accurate potential value. Indeed, as can be seen in Figures 10 and 12, the potential value for small node spacings tends towards  $5.24\,\mathrm{V}$  for both the Jacobi and SOR methods.

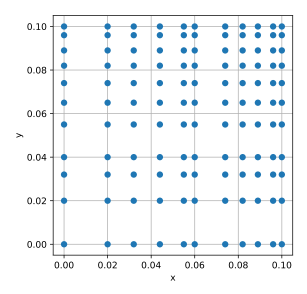


Figure 14: Final mesh arrangement used for nonuniform node spacing. Each point corresponds to a mesh point. Points are positioned closer to the inner conductor, since this is a more difficult area.

# A Code Listings

```
Listing 1: Custom matrix package (matrices.py).
          from __future__ import division
  2
          import copy
 3
  4
          import csv
          from ast import literal_eval
          import math
          class Matrix:
10
11
                   def __init__(self, data):
12
13
                             self.data = data
14
15
                   def __str__(self):
16
                             string = ''
                             for row in self.data:
17
18
                                       string += '\n'
                                       for val in row:
19
                                               string += '{:6.2f} '.format(val)
20
21
                             return string
22
23
                    def __add__(self, other):
                             if len(self) != len(other) or len(self[0]) != len(other[0]):
                                      raise ValueError('Incompatible matrix sizes for addition. Matrix A is {}x{}, but matrix B is
25
                                        \hookrightarrow {}x{}.'
                                                                               .format(len(self), len(self[0]), len(other), len(other[0])))
26
                             rows = len(self)
27
                             cols = len(self[0])
28
29
                             return Matrix([[self[row][col] + other[row][col] for col in range(cols)] for row in range(rows)])
30
31
                   def __sub__(self, other):
32
                             if len(self) != len(other) or len(self[0]) != len(other[0]):
33
                                      raise ValueError('Incompatible matrix sizes for subtraction. Matrix A is {}x{}, but matrix B
34
                                        \hookrightarrow is \{\}x\{\}.
35
                                                                               .format(len(self), len(self[0]), len(other), len(other[0])))
                             rows = len(self)
36
                             cols = len(self[0])
37
                             return Matrix([[self[row][col] - other[row][col] for col in range(cols)] for row in range(rows)])
39
40
41
                    def __mul__(self, other):
                             m = len(self[0])
42
                             n = len(self)
43
                             p = len(other[0])
44
45
                             if m != len(other):
                                       \textbf{raise ValueError('Incompatible matrix sizes for multiplication. Matrix A is $\{\}x\{\}$, but matrix A is $\{\}x\{\}, but matrix A
                                        \hookrightarrow B is \{\}x\{\}.
47
                                                                              .format(n, m, len(other), p))
48
                             # Inspired from https://en.wikipedia.org/wiki/Matrix_multiplication
49
50
                             product = Matrix.empty(n, p)
                             for i in range(n):
51
                                      for j in range(p):
52
                                                row_sum = 0
                                                for k in range(m):
54
                                                         row_sum += self[i][k] * other[k][j]
55
                                               product[i][j] = row_sum
56
                             return product
57
58
                    def __deepcopy__(self, memo):
59
                             return Matrix(copy.deepcopy(self.data))
60
                   def __getitem__(self, item):
62
```

```
63
             return self.data[item]
64
         def __len__(self):
65
             return len(self.data)
66
67
         def is_positive_definite(self):
68
69
70
             :return: True if the matrix if positive-definite, False otherwise.
71
72
             A = copy.deepcopy(self.data)
             n = len(A)
73
             for j in range(n):
74
                 if A[j][j] <= 0:
                      return False
76
                 A[j][j] = math.sqrt(A[j][j])
77
                  for i in range(j + 1, n):
78
                      A[i][j] = A[i][j] / A[j][j]
79
80
                      for k in range(j + 1, i + 1):
                          A[i][k] = A[i][k] - A[i][j] * A[k][j]
81
             return True
82
83
         def transpose(self):
84
85
             :return: the transpose of the current matrix
86
87
88
             rows = len(self)
             cols = len(self[0])
89
             return Matrix([[self.data[row][col] for row in range(rows)] for col in range(cols)])
90
91
         def mirror_horizontal(self):
92
93
             :return: the horizontal mirror of the current matrix
94
95
96
             rows = len(self)
             cols = len(self[0])
97
             return Matrix([[self.data[rows - row - 1][col] for col in range(cols)] for row in range(rows)])
98
99
         def empty_copy(self):
100
101
102
              :return: an empty matrix of the same size as the current matrix.
103
104
             return Matrix.empty(len(self), len(self[0]))
105
         Ostaticmethod
106
         def multiply(*matrices):
107
108
             Computes the product of the given matrices.
109
110
             :param matrices: the matrix objects
111
112
             :return: the product of the given matrices
113
             n = len(matrices[0])
114
115
             product = Matrix.identity(n)
             for matrix in matrices:
116
                 product = product * matrix
117
             return product
118
119
120
         Ostaticmethod
         def empty(num_rows, num_cols):
121
122
             Returns an empty matrix (filled with zeroes) with the specified number of columns and rows.
123
124
             :param num_rows: number of rows
125
             :param num_cols: number of columns
             :return: the empty matrix
127
128
129
             return Matrix([[0 for _ in range(num_cols)] for _ in range(num_rows)])
130
131
         @staticmethod
         def identity(n):
132
```

```
.....
133
             Returns the identity matrix of the given size.
134
135
             :param n: the size of the identity matrix (number of rows or columns)
136
              :return: the identity matrix of size n
137
138
             return Matrix.diagonal_single_value(1, n)
139
140
141
         @staticmethod
142
         def diagonal(values):
143
             Returns a diagonal matrix with the given values along the main diagonal.
144
             :param values: the values along the main diagonal
146
147
             :return: a diagonal matrix with the given values along the main diagonal
148
             n = len(values)
149
             return Matrix([[values[row] if row == col else 0 for col in range(n)] for row in range(n)])
150
151
         @staticmethod
152
153
         def diagonal_single_value(value, n):
154
155
             Returns a diagonal matrix of the given size with the given value along the diagonal.
156
             :param value: the value of each element on the main diagonal
157
             :param n: the size of the matrix
158
             :return: a diagonal matrix of the given size with the given value along the diagonal.
159
160
             return Matrix([[value if row == col else 0 for col in range(n)] for row in range(n)])
161
162
         @staticmethod
163
         def column_vector(values):
164
165
166
             Transforms a row vector into a column vector.
167
             :param values: the values, one for each row of the column vector
168
             :return: the column vector
169
170
             return Matrix([[value] for value in values])
171
172
         @staticmethod
173
174
         def csv_to_matrix(filename):
175
             Reads a CSV file to a matrix.
176
177
             :param filename: the name of the CSV file
178
             :return: a matrix containing the values in the CSV file
179
180
             with open(filename, 'r') as csv_file:
181
182
                 reader = csv.reader(csv_file)
                 data = []
183
                 for row_number, row in enumerate(reader):
184
185
                     data.append([literal_eval(val) for val in row])
                 return Matrix(data)
186
                                  Listing 2: Choleski decomposition (choleski.py).
     from __future__ import division
 3
     import math
    from matrices import Matrix
 5
     def choleski_solve(A, b, half_bandwidth=None):
 8
 9
 10
         Solves an Ax = b matrix equation by Choleski decomposition.
 11
         :param A: the A matrix
```

```
13
        :param b: the b matrix
14
         : param\ half\_bandwidth:\ the\ half\_bandwidth\ of\ the\ A\ matrix
         :return: the solved x vector
15
16
        n = len(A[0])
17
        if half_bandwidth is None:
18
19
             elimination(A, b)
20
21
            elimination_banded(A, b, half_bandwidth)
22
        x = Matrix.empty(n, 1)
        back_substitution(A, x, b)
23
        return x
24
26
    def elimination(A, b):
27
28
         Performs the elimination step of Choleski decomposition.
29
30
         :param A: the A matrix
31
         :param b: the b matrix
32
33
        n = len(A)
34
35
        for j in range(n):
36
             if A[j][j] <= 0:
                 raise ValueError('Matrix A is not positive definite.')
37
38
             A[j][j] = math.sqrt(A[j][j])
             b[j][0] = b[j][0] / A[j][j]
39
             for i in range(j + 1, n):
40
                 A[i][j] = A[i][j] / A[j][j]
41
                 b[i][0] = b[i][0] - A[i][j] * b[j][0]
42
                 for k in range(j + 1, i + 1):
43
                     A[i][k] = A[i][k] - A[i][j] * A[k][j]
44
45
46
    def elimination_banded(A, b, half_bandwidth): # TODO: Keep limited band in memory, improve time
47
     48
        Performs the banded elimination step of Choleski decomposition.
49
50
51
         :param A: the A matrix
         :param b: the b matrix
52
         : param\ half\_bandwidth:\ the\ half\_bandwidth\ to\ be\ used\ for\ the\ banded\ elimination
53
54
        n = len(A)
55
        for j in range(n):
56
             if A[j][j] <= 0:
57
                 raise ValueError('Matrix A is not positive definite.')
58
             A[j][j] = math.sqrt(A[j][j])
59
             b[j][0] = b[j][0] / A[j][j]
60
             for i in range(j + 1, min(j + half_bandwidth, n)):
61
                 A[i][j] = A[i][j] / A[j][j]
62
                 b[i][0] = b[i][0] - A[i][j] * b[j][0]
63
64
                 for k in range(j + 1, i + 1):
                     A[i][k] = A[i][k] - A[i][j] * A[k][j]
65
66
67
    def back_substitution(L, x, y):
68
69
        Performs the back-substitution step of Choleski decomposition.
70
71
72
         : param \ L \colon \ the \ L \ matrix
         :param x: the x matrix
73
74
         :param y: the y matrix
        n = len(L)
76
        for i in range(n - 1, -1, -1):
77
            prev_sum = 0
78
             for j in range(i + 1, n):
79
                 prev_sum += L[j][i] * x[j][0]
80
             x[i][0] = (y[i][0] - prev_sum) / L[i][i]
81
```

```
Listing 3: Linear resistive networks (linear_networks.py).
    from __future__ import division
1
2
3
    import csv
    from matrices import Matrix
4
    from choleski import choleski_solve
5
    def solve_linear_network(A, Y, J, E, half_bandwidth=None):
8
9
         Solve the linear resistive network described by the given matrices.
10
11
         :param A: the incidence matrix
12
         :param Y: the admittance matrix
13
         :param J: the current source matrix
         :param E: the voltage source matrix
15
16
         :param half_bandwidth:
         :return: the solved voltage matrix
17
18
        A_new = A * Y * A.transpose()
19
        b = A * (J - Y * E)
20
        return choleski_solve(A_new, b, half_bandwidth=half_bandwidth)
21
22
23
    def csv_to_network_branch_matrices(filename):
24
25
         Converts a CSV file to Y, J, E network matrices.
26
27
28
         :param filename: the name of the CSV file
         : return: \ the \ Y, \ J, \ E \ network \ matrices
29
30
        with open(filename, 'r') as csv_file:
31
            reader = csv.reader(csv file)
32
             J = []
33
            Y = []
34
            E = []
35
            for row in reader:
36
                J k = float(row[0])
37
38
                R_k = float(row[1])
                E_k = float(row[2])
39
                 J.append(J_k)
40
41
                 Y.append(1 / R_k)
                 E.append(E_k)
42
            Y = Matrix.diagonal(Y)
43
44
             J = Matrix.column_vector(J)
            E = Matrix.column_vector(E)
45
46
            return Y, J, E
47
48
    def create_network_matrices_mesh(rows, cols, branch_resistance, test_current):
49
        num_horizontal_branches = (cols - 1) * rows
50
51
        num_vertical_branches = (rows - 1) * cols
        num_branches = num_horizontal_branches + num_vertical_branches + 1
52
        num_nodes = rows * cols - 1
53
54
         A = create_incidence_matrix_mesh(cols, num_branches, num_horizontal_branches, num_nodes,
55

→ num vertical branches)

         Y, J, E = create_network_branch_matrices_mesh(num_branches, branch_resistance, test_current)
56
57
58
        return A, Y, J, E
59
60
61
    def create_incidence_matrix_mesh(cols, num_branches, num_horizontal_branches, num_nodes,
     → num_vertical_branches):
        A = Matrix.empty(num_nodes, num_branches)
62
63
        node\_offset = -1
        for branch in range(num_horizontal_branches):
64
             if branch == num_horizontal_branches - cols + 1:
65
```

```
A[branch + node_offset + 1][branch] = 1
66
67
             else:
                 if branch % (cols - 1) == 0:
68
                     node_offset += 1
69
                 node_number = branch + node_offset
70
                  A[node_number][branch] = -1
71
                 A[node_number + 1][branch] = 1
72
73
         branch_offset = num_horizontal_branches
         node_offset = cols
74
         for branch in range(num_vertical_branches):
75
             if branch == num_vertical_branches - cols:
76
                 node offset -= 1
77
                 A[branch][branch + branch_offset] = 1
             else:
79
                 A[branch][branch + branch_offset] = 1
80
                 A[branch + node_offset][branch + branch_offset] = -1
81
         if num branches == 2:
82
83
             A[0][1] = -1
84
             A[cols - 1][num\_branches - 1] = -1
85
86
         return A
87
88
     def create_network_branch_matrices_mesh(num_branches, resistance, test_current):
89
         Y = Matrix.diagonal([1 / resistance if branch < num_branches - 1 else 0 for branch in
90

    range(num_branches)])
         # Negative test current here because we assume current is coming OUT of the test current node.
91
         J = Matrix.column_vector([0 if branch < num_branches - 1 else -test_current for branch in
92
          \hookrightarrow range(num_branches)])
         E = Matrix.column_vector([0 for branch in range(num_branches)])
93
94
         return Y, J, E
95
96
97
     def find_mesh_resistance(n, branch_resistance, half_bandwidth=None):
         test_current = 0.01
98
         A, Y, J, E = create_network_matrices_mesh(n, 2 * n, branch_resistance, test_current)
99
         x = solve_linear_network(A, Y, J, E, half_bandwidth=half_bandwidth)
100
         test_voltage = x[2 * n - 1 \text{ if } n > 1 \text{ else } 0][0]
101
         equivalent_resistance = test_voltage / test_current
102
         return equivalent_resistance
                                            Listing 4: Question 1 (q1.py).
 1
     from __future__ import division
     from linear_networks import solve_linear_network, csv_to_network_branch_matrices
 3
     from choleski import choleski_solve
 5
     from matrices import Matrix
     NETWORK_DIRECTORY = 'network_data'
     L_2 = Matrix([
 9
         [5, 0],
10
         [1, 3]
11
     ])
12
     L_3 = Matrix([
13
         [3, 0, 0],
14
15
         [1, 2, 0],
         [8, 5, 1]
16
     ])
17
     L_4 = Matrix([
18
         [1, 0, 0, 0],
19
20
         [2, 8, 0, 0],
         [5, 5, 4, 0],
21
         [7, 2, 8, 7]
22
     1)
23
     matrix_2 = L_2 * L_2.transpose()
24
    matrix_3 = L_3 * L_3.transpose()
25
    matrix_4 = L_4 * L_4.transpose()
```

```
positive_definite_matrices = [matrix_2, matrix_3, matrix_4]
27
    x_2 = Matrix.column_vector([8, 3])
29
    x_3 = Matrix.column_vector([9, 4, 3])
30
    x_4 = Matrix.column_vector([5, 4, 1, 9])
31
    xs = [x_2, x_3, x_4]
32
33
34
35
    def q1b():
        print('=== Question 1(b) ===')
36
        for count, A in enumerate(positive_definite_matrices):
37
            n = count + 2
38
            print('n={} matrix is positive-definite: {}'.format(n, A.is_positive_definite()))
39
40
41
    def q1c():
42
        print('=== Question 1(c) ===')
43
44
        n = 2
        for x, A in zip(xs, positive_definite_matrices):
45
            b = A * x
46
47
            # print('A: {}'.format(A))
            # print('b: {}'.format(b))
48
49
            x_choleski = choleski_solve(A, b)
50
            print('Matrix with n={}:'.format(n))
51
52
            print('Expected x: {}'.format(x))
            print('Actual x: {}'.format(x_choleski))
53
            n += 1
54
56
    def q1d():
57
        print('=== Question 1(d) ===')
58
        for i in range(1, 7):
59
            A = Matrix.csv_to_matrix('{}/incidence_matrix_{}.csv'.format(NETWORK_DIRECTORY, i))
60
            Y, J, E = csv_to_network_branch_matrices('{}\network_branches_{}\.csv'.format(NETWORK_DIRECTORY,
61

→ i))

            # print('Y: {}'.format(Y))
62
            # print('J: {}'.format(J))
63
            # print('E: {}'.format(E))
64
65
            x = solve_linear_network(A, Y, J, E)
            print('Solved for x in network {}:'.format(i)) # TODO: Create my own test circuits here
66
67
            for j in range(len(x)):
                print('V{} = {:.3f} V'.format(j + 1, x[j][0]))
68
69
70
    def q1():
71
72
        q1b()
        q1c()
73
        q1d()
74
75
76
    if __name__ == '__main__':
77
        q1()
                                           Listing 5: Question 2 (q2.py).
    import csv
2
    import time
    import matplotlib.pyplot as plt
    import numpy as np
    import numpy.polynomial.polynomial as poly
    import sympy as sp
    from matplotlib.ticker import MaxNLocator
10
    from linear_networks import find_mesh_resistance
11
12
    def find_mesh_resistances(banded):
```

```
14
        branch_resistance = 1000
        points = {}
15
        runtimes = {}
16
        for n in range(2, 11):
17
            start_time = time.time()
18
            half_bandwidth = 2 * n + 1 if banded else None
19
            equivalent_resistance = find_mesh_resistance(n, branch_resistance, half_bandwidth=half_bandwidth)
20
            print('Equivalent resistance for \{\}x\{\} mesh: \{:.2f\} Ohms.'.format(n, 2 * n,
21
             points[n] = '{:.3f}'.format(equivalent_resistance)
22
            runtime = time.time() - start_time
23
            runtimes[n] = '{:.3f}'.format(runtime)
24
            print('Runtime: {} s.'.format(runtime))
        plot_runtime(runtimes, banded)
26
27
        return points, runtimes
28
29
    def q2ab():
30
        print('=== Question 2(a)(b) ===')
31
        _, runtimes = find_mesh_resistances(banded=False)
32
33
        save_rows_to_csv('report/csv/q2b.csv', zip(runtimes.keys(), runtimes.values()), header=('N', 'Runtime
         34
        return runtimes
35
36
    def q2c():
37
        print('=== Question 2(c) ===')
38
        pts, runtimes = find_mesh_resistances(banded=True)
39
        save_rows_to_csv('report/csv/q2c.csv', zip(runtimes.keys(), runtimes.values()), header=('N', 'Runtime
40
         41
        return pts, runtimes
42
43
44
    def plot_runtime(points, banded=False):
45
        N^6: non-banded
46
        N^4: banded
47
48
49
        :param points:
50
        :param banded:
51
52
        f = plt.figure()
        ax = f.gca()
53
        ax.xaxis.set_major_locator(MaxNLocator(integer=True))
54
        x_range = [float(x) for x in points.keys()]
        y_range = [float(y) for y in points.values()]
56
        plt.plot(x_range, y_range, '{}o'.format('C1' if banded else 'C0'), label='Data points')
57
58
        x_new = np.linspace(x_range[0], x_range[-1], num=len(x_range) * 10)
59
60
        degree = 4 if banded else 6
        polynomial_coeffs = poly.polyfit(x_range, y_range, degree)
61
        polynomial_fit = poly.polyval(x_new, polynomial_coeffs)
62
63
        N = sp.symbols("N")
        poly_label = sum(sp.S("{:.4f}".format(v)) * N ** i for i, v in enumerate(polynomial_coeffs))
64
        equation = '${}$'.format(sp.printing.latex(poly_label))
65
        plt.plot(x_new, polynomial_fit, '{}-'.format('C1' if banded else 'C0'), label=equation)
66
67
68
        plt.xlabel('N')
69
        plt.ylabel('Runtime (s)')
        plt.grid(True)
70
        plt.legend(fontsize='x-small')
71
        f.savefig('report/plots/q2{}.pdf'.format('c' if banded else 'b'), bbox_inches='tight')
72
73
74
    def plot_runtimes(points1, points2):
75
76
        f = plt.figure()
        ax = f.gca()
77
        ax.xaxis.set_major_locator(MaxNLocator(integer=True))
78
        x_range = points1.keys()
79
        y_range = points1.values()
80
```

```
81
         y_banded_range = points2.values()
82
         plt.plot(x_range, y_range, 'o-', label='Non-banded elimination')
        plt.plot(x_range, y_banded_range, 'o-', label='Banded elimination')
83
         plt.xlabel('N')
84
         plt.ylabel('Runtime (s)')
85
        plt.grid(True)
86
87
         plt.legend()
88
         f.savefig('report/plots/q2bc.pdf', bbox_inches='tight')
89
90
     def q2d(points):
91
         print('=== Question 2(d) ===')
92
         f = plt.figure()
93
         ax = f.gca()
94
         ax.xaxis.set_major_locator(MaxNLocator(integer=True))
95
        x_range = [float(x) for x in points.keys()]
96
         y_range = [float(y) for y in points.values()]
97
98
         plt.plot(x_range, y_range, 'o', label='Data points')
99
        x_new = np.linspace(x_range[0], x_range[-1], num=len(x_range) * 10)
100
101
         coeffs = poly.polyfit(np.log(x_range), y_range, deg=1)
        {\tt polynomial\_fit = poly.polyval(np.log(x\_new), coeffs)}
102
         103
          \hookrightarrow coeffs[0]))
104
105
        plt.xlabel('N')
        plt.ylabel('R ($\Omega$)')
106
        plt.grid(True)
107
        plt.legend()
108
         f.savefig('report/plots/q2d.pdf', bbox_inches='tight')
109
         save_rows_to_csv('report/csv/q2a.csv', zip(points.keys(), points.values()), header=('N', 'R (Omega)'))
110
111
112
113
     def q2():
        runtimes1 = q2ab()
114
         pts, runtimes2 = q2c()
115
         plot_runtimes(runtimes1, runtimes2)
116
         q2d(pts)
117
118
119
     def save_rows_to_csv(filename, rows, header=None):
120
121
         with open(filename, "wb") as f:
            writer = csv.writer(f)
122
            if header is not None:
123
                writer.writerow(header)
            for row in rows:
125
126
                writer.writerow(row)
127
128
     if __name__ == '__main__':
129
         q2()
130
                              Listing 6: Finite difference method (finite_diff.py).
     from __future__ import division
 3
     import math
 4
     import random
    from abc import ABCMeta, abstractmethod
    from matrices import Matrix
    MESH_SIZE = 0.2
 10
 11
 12
     class Relaxer:
13
         Performs the relaxing stage of the finite difference method.
 14
```

```
16
        __metaclass__ = ABCMeta
17
        @abstractmethod
18
        def relax(self, phi, i, j):
19
20
            Perform a relaxation iteration on a given (i, j) point of the given phi matrix.
21
22
23
            :param phi: the phi matrix
            :param i: the row index
24
            :param j: the column index
25
26
            raise NotImplementedError
27
        def reset(self):
29
30
            Optional method to reset the relaxer.
31
32
33
            pass
34
        def residual(self, phi, i, j):
35
36
            Calculate the residual at the given (i, j) point of the given phi matrix.
37
38
39
            :param phi: the phi matrix
            :param i: the row index
40
41
            :param j: the column index
            :return:
42
43
            return abs(phi[i + 1][j] + phi[i - 1][j] + phi[i][j + 1] + phi[i][j - 1] - 4 * phi[i][j])
45
46
    class GaussSeidelRelaxer(Relaxer):
47
        def relax(self, phi, i, j):
48
            return (phi[i + 1][j] + phi[i - 1][j] + phi[i][j + 1] + phi[i][j - 1]) / 4
49
50
51
    class JacobiRelaxer(Relaxer):
52
        def __init__(self, num_cols):
53
54
            self.num_cols = num_cols
55
            self.prev_row = [0] * (num_cols - 1) # Don't need to copy entire phi, just previous row
56
57
        def relax(self, phi, i, j):
            left_val = self.prev_row[j - 2] if j > 1 else 0
58
            top_val = self.prev_row[j - 1]
59
            self.prev_row[j - 1] = phi[i][j]
            return (phi[i + 1][j] + top_val + phi[i][j + 1] + left_val) / 4
61
62
        def reset(self):
63
            self.prev_row = [0] * (self.num_cols - 1)
64
65
66
    class NonUniformRelaxer(Relaxer):
67
68
        def __init__(self, mesh):
            self.mesh = mesh
69
70
        def get_distances(self, i, j):
71
            a1 = self.mesh.get_y(i) - self.mesh.get_y(i - 1)
72
            a2 = self.mesh.get_y(i + 1) - self.mesh.get_y(i)
73
            b1 = self.mesh.get_x(j) - self.mesh.get_x(j - 1)
74
            b2 = self.mesh.get_x(j + 1) - self.mesh.get_x(j)
75
            return a1, a2, b1, b2
76
77
        def relax(self, phi, i, j):
78
            a1, a2, b1, b2 = self.get_distances(i, j)
80
            return ((phi[i - 1][j] / a1 + phi[i + 1][j] / a2) / (a1 + a2)
81
                     + (phi[i][j - 1] / b1 + phi[i][j + 1] / b2) / (b1 + b2)) / (1 / (a1 * a2) + 1 / (b1 * b2))
82
83
84
        def residual(self, phi, i, j):
            a1, a2, b1, b2 = self.get_distances(i, j)
85
```

```
86
87
             return abs(((phi[i - 1][j] / a1 + phi[i + 1][j] / a2) / (a1 + a2)
                          + (phi[i][j - 1] / b1 + phi[i][j + 1] / b2) / (b1 + b2))
88
                         - phi[i][j] * (1 / (a1 * a2) + 1 / (b1 * b2)))
89
90
91
     class SuccessiveOverRelaxer(Relaxer):
92
93
         def __init__(self, omega):
             self.gauss_seidel = GaussSeidelRelaxer()
94
95
             self.omega = omega
96
         def relax(self, phi, i, j, last_row=None, a1=None, a2=None, b1=None, b2=None):
97
             return (1 - self.omega) * phi[i][j] + self.omega * self.gauss_seidel.relax(phi, i, j)
98
99
100
     class Boundary:
101
102
103
         Constant-potential boundary in the finite difference mesh, representing a conductor.
104
         __metaclass__ = ABCMeta
105
106
         @abstractmethod
107
108
         def potential(self):
109
             Return the potential on the boundary.
110
111
             raise NotImplementedError
112
113
         @abstractmethod
114
         def contains_point(self, x, y):
115
116
             Returns true if the boundary contains the given (x, y) point.
117
118
             :param x: the x coordinate of the point
119
             :param y: the y coordinate of the point
120
121
122
             raise NotImplementedError
123
124
125
     class OuterConductorBoundary(Boundary):
         def potential(self):
126
127
             return 0
128
         def contains_point(self, x, y):
129
             return x == 0 or y == 0 or x == 0.2 or y == 0.2
130
131
132
     class QuarterInnerConductorBoundary(Boundary):
133
         def potential(self):
134
135
             return 15
136
         def contains_point(self, x, y):
137
138
             return 0.06 \le x \le 0.14 and 0.08 \le y \le 0.12
139
140
     class PotentialGuesser:
141
142
         Guesses the initial potential in the finite-difference mesh.
143
144
         __metaclass__ = ABCMeta
145
146
         def __init__(self, min_potential, max_potential):
147
             self.min_potential = min_potential
148
             self.max_potential = max_potential
149
150
151
         @abstractmethod
         def guess(self, x, y):
152
153
154
             Guess the potential at the given (x, y) point, and return it.
```

```
:param x: the x coordinate of the point
156
157
              :param y: the y coordinate of the point
158
             raise NotImplementedError
159
160
161
     class RandomPotentialGuesser(PotentialGuesser):
162
163
         def guess(self, x, y):
             return random.randint(self.min_potential, self.max_potential)
164
165
166
     class LinearPotentialGuesser(PotentialGuesser):
167
         def guess(self, x, y):
168
             return 150 * x if x < 0.06 else 150 * y
169
170
171
     class RadialPotentialGuesser(PotentialGuesser):
172
173
         def guess(self, x, y):
              def radial(k, x, y, x_source, y_source):
174
                  return k / (math.sqrt((x_source - x) ** 2 + (y_source - y) ** 2))
175
176
             return 0.0225 * (radial(20, x, y, 0.1, 0.1) - radial(1, x, y, 0, y) - radial(1, x, y, x, 0))
177
178
179
     class PhiConstructor:
180
181
         Constructs the phi potential matrix with an outer conductor, inner conductor, mesh points and an inital
182
         potential
         guess.
183
184
185
         def __init__(self, mesh):
186
              outer_boundary = OuterConductorBoundary()
187
188
              inner_boundary = QuarterInnerConductorBoundary()
              self.boundaries = (inner_boundary, outer_boundary)
189
              self.guesser = RadialPotentialGuesser(0, 15)
190
              self.mesh = mesh
191
192
         def construct_phi(self):
193
194
              phi = Matrix.empty(self.mesh.num_rows, self.mesh.num_cols)
              for i in range(self.mesh.num_rows):
195
196
                  y = self.mesh.get_y(i)
                  for j in range(self.mesh.num_cols):
197
                      x = self.mesh.get_x(j)
198
                      boundary_pt = False
199
                      for boundary in self.boundaries:
200
201
                          if boundary.contains_point(x, y):
                              boundary_pt = True
202
                              phi[i][j] = boundary.potential()
203
204
                      if not boundary_pt:
                          phi[i][j] = self.guesser.guess(x, y)
205
             return phi
206
207
208
209
     class SquareMeshConstructor:
210
         {\it Constructs~a~square~mesh.}
211
212
213
         def __init__(self, size):
214
215
              self.size = size
216
         def construct_uniform_mesh(self, h):
217
218
             Constructs a uniform mesh with the given node spacing.
219
220
221
              :param h: the node spacing
              :return: the constructed mesh
222
223
             num_rows = num_cols = int(self.size / h) + 1
224
```

```
225
             return SimpleMesh(h, num_rows, num_cols)
226
         def construct_symmetric_uniform_mesh(self, h):
227
228
              Construct a symmetric uniform mesh with the given node spacing.
230
231
              :param h: the node spacing
232
              :return: the constructed mesh
233
234
             half_size = self.size / 2
             num_rows = num_cols = int(half_size / h) + 2 # Only need to store up to middle
235
             return SimpleMesh(h, num_rows, num_cols)
236
         def construct_symmetric_non_uniform_mesh(self, x_values, y_values):
238
239
              Construct a symmetric non-uniform mesh with the given adjacent x coordinates and y coordinates.
240
241
242
              :param x\_values: the values of successive x coordinates
              :param y_values: the values of successive y coordinates
243
              :return: the constructed mesh
244
             return NonUniformMesh(x_values, y_values)
246
247
^{248}
     class Mesh:
249
250
         Finite-difference mesh.
251
252
         __metaclass__ = ABCMeta
253
254
         @abstractmethod
255
         def get_x(self, j):
256
257
              Get the x value at the specified index.
258
259
260
              :param j: the column index.
261
             raise NotImplementedError
262
263
264
         @abstractmethod
         def get_y(self, i):
265
266
              Get the y value at the specified index.
267
268
              : param \ i: \ the \ row \ index.
270
             raise NotImplementedError
271
272
         @abstractmethod
273
274
         def get_i(self, y):
275
             Get the row index of the specified y coordinate.
276
277
              :param y: the y coordinate
278
279
              raise NotImplementedError
280
281
282
         @abstractmethod
         def get_j(self, x):
283
284
285
              Get the column index of the specified x coordinate.
286
              :param x: the x coordinate
287
288
             raise NotImplementedError
289
290
291
         def point_to_indices(self, x, y):
292
              Converts the given (x, y) point to (i, j) matrix indices.
293
```

```
295
             :param x: the x coordinate
296
              :param\ y:\ the\ y\ coordinate
              :return: the (i, j) matrix indices
297
298
             return self.get_i(y), self.get_j(x)
299
300
         def indices_to_points(self, i, j):
301
302
             Converts the given (i, j) matrix indices to an (x, y) point.
303
304
             :param i: the row index
305
             :param j: the column index
306
             :return: the (x, y) point
308
             return self.get_x(j), self.get_y(i)
309
310
311
312
     class SimpleMesh(Mesh):
         def __init__(self, h, num_rows, num_cols):
313
             self.h = h
314
315
             self.num_rows = num_rows
             self.num_cols = num_cols
316
317
         def get_i(self, y):
318
             return int(y / self.h)
319
320
         def get_j(self, x):
321
             return int(x / self.h)
322
         def get_x(self, j):
324
             return j * self.h
325
326
         def get_y(self, i):
327
328
             return i * self.h
329
330
331
     class NonUniformMesh(Mesh):
         def __init__(self, x_values, y_values):
332
333
             self.x\_values = x\_values
334
             self.y_values = y_values
             self.num_rows = len(y_values)
335
336
             self.num_cols = len(x_values)
337
         def get_i(self, y):
338
             return self.y_values.index(y)
339
340
         def get_j(self, x):
341
             return self.x_values.index(x)
342
343
         def get_x(self, j):
344
             return self.x_values[j]
345
346
347
         def get_y(self, i):
             return self.y_values[i]
348
349
     class IterativeRelaxer:
351
352
         Performs finite-difference iterative relaxation on a phi potential matrix associated with a mesh.
353
354
355
         def __init__(self, relaxer, epsilon, phi, mesh):
356
             self.relaxer = relaxer
357
             self.epsilon = epsilon
             self.phi = phi
359
             self.boundary = QuarterInnerConductorBoundary()
360
             self.num_iterations = 0
361
             self.rows = len(phi)
362
             self.cols = len(phi[0])
             self.mesh = mesh
364
```

```
self.mid_i = mesh.get_i(MESH_SIZE / 2)
365
366
              self.mid_j = mesh.get_j(MESH_SIZE / 2)
367
         def relaxation(self):
368
369
              Performs iterative relaxation until convergence is met.
370
371
372
              :return: the current iterative relaxer object
373
              while not self.convergence():
374
                  self.num_iterations += 1
375
                  self relaxation iteration()
376
                  self.relaxer.reset()
              return self
378
379
         def relaxation_iteration(self):
380
381
382
              Performs one iteration of relaxation.
383
             for i in range(1, self.rows - 1):
384
385
                  y = self.mesh.get_y(i)
                  for j in range(1, self.cols - 1):
386
387
                      x = self.mesh.get_x(j)
                      if not self.boundary.contains_point(x, y):
388
                          relaxed_value = self.relaxer.relax(self.phi, i, j)
389
                          self.phi[i][j] = relaxed_value
390
                          if i == self.mid_i - 1:
391
                              self.phi[i + 2][j] = relaxed_value
392
                          elif j == self.mid_j - 1:
393
                              self.phi[i][j + 2] = relaxed_value
394
395
         def convergence(self):
396
397
398
              Checks if the phi matrix has reached convergence.
399
              :return: True if the phi matrix has reached convergence, False otherwise
400
401
             max_i, max_j = self.mesh.point_to_indices(0.1, 0.1) # Only need to compute for 1/4 of grid
402
403
             for i in range(1, max_i + 1):
404
                 y = self.mesh.get_y(i)
                  for j in range(1, max_j + 1):
405
406
                      x = self.mesh.get_x(j)
                      if not self.boundary.contains_point(x, y) and self.relaxer.residual(self.phi, i, j) >=
407

    self.epsilon:

                          return False
408
              return True
409
410
         def get_potential(self, x, y):
411
412
413
              Get the potential at the given (x, y) point.
414
              :param x: the x coordinate
415
416
              :param\ y:\ the\ y\ coordinate
              :return: the potential at the given (x, y) point
417
418
              i, j = self.mesh.point_to_indices(x, y)
419
             return self.phi[i][j]
420
421
422
     def non_uniform_jacobi(epsilon, x_values, y_values):
423
424
         Perform Jacobi relaxation on a non-uniform finite-difference mesh.
425
426
          :param epsilon: the maximum error to achieve convergence
427
          :param x_values: the values of successive x coordinates
428
429
          :param y\_values: the values of successive y coordinates
          :return: the relaxer object
430
431
         mesh = SquareMeshConstructor(MESH_SIZE).construct_symmetric_non_uniform_mesh(x_values, y_values)
432
         relaxer = NonUniformRelaxer(mesh)
433
```

```
phi = PhiConstructor(mesh).construct_phi()
434
435
         return IterativeRelaxer(relaxer, epsilon, phi, mesh).relaxation()
436
437
     def successive_over_relaxation(omega, epsilon, h):
438
439
         {\it Perform~SOR~on~a~uniform~symmetric~finite-difference~mesh.}
440
441
         : param\ omega:\ the\ omega\ value\ for\ SOR
442
          :param epsilon: the maximum error to achieve convergence
443
         :param h: the node spacing
444
         :return: the relaxer object
445
446
         mesh = SquareMeshConstructor(MESH_SIZE).construct_symmetric_uniform_mesh(h)
447
448
         relaxer = SuccessiveOverRelaxer(omega)
         phi = PhiConstructor(mesh).construct_phi()
449
         return IterativeRelaxer(relaxer, epsilon, phi, mesh).relaxation()
450
451
452
     def jacobi_relaxation(epsilon, h):
453
454
         Perform Jacobi relaxation on a uniform symmetric finite-difference mesh.
455
456
          :param epsilon: the maximum error to achieve convergence
457
          :param h: the node spacing
458
459
          :return: the relaxer object
460
         mesh = SquareMeshConstructor(MESH_SIZE).construct_symmetric_uniform_mesh(h)
461
         relaxer = GaussSeidelRelaxer()
462
         phi = PhiConstructor(mesh).construct_phi()
463
         return IterativeRelaxer(relaxer, epsilon, phi, mesh).relaxation()
464
                                            Listing 7: Question 3 (q3.py).
     from __future__ import division
 1
 2
     import csv
     import time
 4
     import matplotlib.pyplot as plt
     import numpy as np
     import numpy.polynomial.polynomial as poly
     import sympy as sp
 9
 10
     from finite_diff import successive_over_relaxation, jacobi_relaxation, \
 11
         non_uniform_jacobi
12
 13
     EPSILON = 0.00001
 14
     X_QUERY = 0.06
 15
     Y QUERY = 0.04
 16
     NUM_H_ITERATIONS = 6
17
 18
 19
     def q3b():
20
21
         print('=== Question 3(b) ===')
         h = 0.02
22
         min_num_iterations = float('inf')
23
24
         best_omega = float('inf')
25
26
         omegas = []
         num_iterations = []
27
         potentials = []
28
29
         for omega_diff in range(10):
30
             omega = 1 + omega_diff / 10
31
             print('Omega: {}'.format(omega))
32
             iter_relaxer = successive_over_relaxation(omega, EPSILON, h)
33
             print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
34
             print('Num iterations: {}'.format(iter_relaxer.num_iterations))
```

```
potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
36
             print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
37
             if iter_relaxer.num_iterations < min_num_iterations:</pre>
38
                 best_omega = omega
39
             min_num_iterations = min(min_num_iterations, iter_relaxer.num_iterations)
40
41
42
             omegas.append(omega)
43
             num_iterations.append(iter_relaxer.num_iterations)
             potentials.append('{:.3f}'.format(potential))
44
45
         print('Best number of iterations: {}'.format(min_num_iterations))
46
         print('Best omega: {}'.format(best_omega))
47
48
         f = plt.figure()
49
50
         x_range = omegas
         y_range = num_iterations
51
         plt.plot(x_range, y_range, 'o-', label='Number of iterations')
52
         plt.xlabel('$\omega$')
53
         plt.ylabel('Number of Iterations')
54
         plt.grid(True)
55
56
         f.savefig('report/plots/q3b.pdf', bbox_inches='tight')
57
58
         save_rows_to_csv('report/csv/q3b_potential.csv', zip(omegas, potentials), header=('Omega', 'Potential
         save_rows_to_csv('report/csv/q3b_iterations.csv', zip(omegas, num_iterations), header=('Omega',
59
          60
         return best_omega
61
62
63
     def q3c(omega):
64
         print('=== Question 3(c): SOR ===')
65
         h = 0.04
66
         h_values = []
67
         potential_values = []
68
         iterations_values = []
69
         for i in range(NUM_H_ITERATIONS):
70
            h = h / 2
71
             print('h: {}'.format(h))
72
73
             print('1/h: {}'.format(1 / h))
             iter_relaxer = successive_over_relaxation(omega, EPSILON, h)
74
             # print(phi.mirror_horizontal())
75
             potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
76
             num_iterations = iter_relaxer.num_iterations
77
             print('Num iterations: {}'.format(num_iterations))
79
             print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
80
81
82
             h_values.append(1 / h)
             potential_values.append('{:.3f}'.format(potential))
83
             iterations_values.append(num_iterations)
84
85
86
         f = plt.figure()
         x_range = h_values
87
         y_range = potential_values
88
         plt.plot(x_range, y_range, 'o-', label='Data points')
89
90
91
         plt.xlabel('1 / h')
         plt.ylabel('Potential at [0.06, 0.04] (V)')
92
         plt.grid(True)
93
         f.savefig('report/plots/q3c_potential.pdf', bbox_inches='tight')
94
95
96
         f = plt.figure()
         x_range = h_values
97
         y_range = iterations_values
98
99
         x_new = np.linspace(x_range[0], x_range[-1], num=len(x_range) * 10)
100
         polynomial_coeffs = poly.polyfit(x_range, y_range, deg=3)
101
         polynomial_fit = poly.polyval(x_new, polynomial_coeffs)
102
         N = sp.symbols("1/h")
103
```

```
poly_label = sum(sp.S("{:.5f}".format(v)) * N ** i for i, v in enumerate(polynomial_coeffs))
104
                 equation = '${}$'.format(sp.printing.latex(poly_label))
105
                plt.plot(x_new, polynomial_fit, '{}-'.format('CO'), label=equation)
106
107
                plt.plot(x_range, y_range, 'o', label='Data points')
108
                plt.xlabel('1 / h')
109
110
                plt.ylabel('Number of Iterations')
                plt.grid(True)
111
                plt.legend(fontsize='small')
112
113
                f.savefig('report/plots/q3c_iterations.pdf', bbox_inches='tight')
114
115
                 save_rows_to_csv('report/csv/q3c_potential.csv', zip(h_values, potential_values), header=('1/h',
116
                  → 'Potential (V)'))
                 save\_rows\_to\_csv('report/csv/q3c\_iterations.csv', \ zip(h\_values, iterations\_values), \ header=('1/h', report/csv/q3c\_iterations.csv', report/csv', report/csv'
117
                         'Iterations'))
118
                return h_values, potential_values, iterations_values
119
120
121
122
         def q3d():
                print('=== Question 3(d): Jacobi ===')
123
124
                h = 0.04
125
                h_values = []
                potential_values = []
126
                 iterations_values = []
127
                for i in range(NUM_H_ITERATIONS):
128
                        h = h / 2
129
                        print('h: {}'.format(h))
130
                        iter_relaxer = jacobi_relaxation(EPSILON, h)
131
                        potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
132
133
                        num_iterations = iter_relaxer.num_iterations
134
135
                        print('Num iterations: {}'.format(num_iterations))
                        print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
136
137
                        h_values.append(1 / h)
138
                        potential_values.append('{:.3f}'.format(potential))
139
140
                        iterations_values.append(num_iterations)
141
                f = plt.figure()
142
                x_range = h_values
143
                y_range = potential_values
144
                plt.plot(x_range, y_range, 'C1o-', label='Data points')
145
                plt.xlabel('1 / h')
146
                plt.ylabel('Potential at [0.06, 0.04] (V)')
147
148
                 plt.grid(True)
                 f.savefig('report/plots/q3d_potential.pdf', bbox_inches='tight')
149
150
151
                f = plt.figure()
                x_range = h_values
152
                y_range = iterations_values
153
154
                plt.plot(x_range, y_range, 'C1o', label='Data points')
                plt.xlabel('1 / h')
155
156
                plt.ylabel('Number of Iterations')
157
                 x_new = np.linspace(x_range[0], x_range[-1], num=len(x_range) * 10)
158
159
                 polynomial_coeffs = poly.polyfit(x_range, y_range, deg=4)
160
                 polynomial_fit = poly.polyval(x_new, polynomial_coeffs)
                 N = sp.symbols("1/h")
161
                 poly_label = sum(sp.S("{:.5f})".format(v if i < 3 else -v)) * N ** i for i, v in
162
                        enumerate(polynomial_coeffs))
                 equation = '${}$'.format(sp.printing.latex(poly_label))
163
                plt.plot(x_new, polynomial_fit, '{}-'.format('C1'), label=equation)
164
165
166
                 plt.grid(True)
                plt.legend(fontsize='small')
167
168
                 f.savefig('report/plots/q3d_iterations.pdf', bbox_inches='tight')
169
```

170

```
171
         save_rows_to_csv('report/csv/q3d_potential.csv', zip(h_values, potential_values), header=('1/h',
              'Potential (V)'))
         save_rows_to_csv('report/csv/q3d_iterations.csv', zip(h_values, iterations_values), header=('1/h',
172
              'Iterations'))
173
         return h_values, potential_values, iterations_values
174
175
176
     def q3e():
177
         print('=== Question 3(e): Non-Uniform Node Spacing ===')
178
179
         print('Jacobi (for reference)')
180
         iter_relaxer = jacobi_relaxation(EPSILON, 0.01)
181
         print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
182
183
         print('Num iterations: {}'.format(iter_relaxer.num_iterations))
184
         jacobi_potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
         print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, jacobi_potential))
185
186
         print('Uniform Mesh (same as Jacobi)')
187
         x_values = [0.00, 0.01, 0.02, 0.03, 0.04, 0.05, 0.06, 0.07, 0.08, 0.09, 0.1, 0.11]
188
189
         y_{values} = [0.00, 0.01, 0.02, 0.03, 0.04, 0.05, 0.06, 0.07, 0.08, 0.09, 0.1, 0.11]
         iter_relaxer = non_uniform_jacobi(EPSILON, x_values, y_values)
190
191
         print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
         print('Num iterations: {}'.format(iter_relaxer.num_iterations))
192
         uniform_potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
193
         print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, uniform_potential))
194
         print('Jacobi potential: {} V, same as uniform potential: {} V'.format(jacobi_potential,
195

    uniform_potential))

196
         print('Non-Uniform (clustered around (0.06, 0.04))')
197
         x_{values} = [0.00, 0.01, 0.02, 0.03, 0.05, 0.055, 0.06, 0.065, 0.07, 0.09, 0.1, 0.11]
198
         y_values = [0.00, 0.01, 0.03, 0.035, 0.04, 0.045, 0.05, 0.07, 0.08, 0.09, 0.1, 0.11]
199
         iter_relaxer = non_uniform_jacobi(EPSILON, x_values, y_values)
200
201
         print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
         print('Num iterations: {}'.format(iter_relaxer.num_iterations))
202
         potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
203
         print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
204
205
         print('Non-Uniform (more clustered around (0.06, 0.04))')
206
207
         x_values = [0.00, 0.01, 0.02, 0.03, 0.055, 0.059, 0.06, 0.061, 0.065, 0.09, 0.1, 0.11]
         v_values = [0.00, 0.01, 0.035, 0.039, 0.04, 0.041, 0.045, 0.07, 0.08, 0.09, 0.1, 0.11]
208
         iter_relaxer = non_uniform_jacobi(EPSILON, x_values, y_values)
209
         print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
210
         print('Num iterations: {}'.format(iter_relaxer.num_iterations))
211
         potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
212
         print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
213
214
         print('Non-Uniform (clustered near outer conductor)')
215
         x_values = [0.00, 0.020, 0.032, 0.044, 0.055, 0.06, 0.074, 0.082, 0.089, 0.096, 0.1, 0.14]
216
217
         y_values = [0.00, 0.020, 0.032, 0.04, 0.055, 0.065, 0.074, 0.082, 0.089, 0.096, 0.1, 0.14]
         iter_relaxer = non_uniform_jacobi(EPSILON, x_values, y_values)
218
         print('Quarter grid: {}'.format(iter_relaxer.phi.mirror_horizontal()))
219
220
         print('Num iterations: {}'.format(iter_relaxer.num_iterations))
         potential = iter_relaxer.get_potential(X_QUERY, Y_QUERY)
221
         print('Potential at ({}, {}): {:.3f} V'.format(X_QUERY, Y_QUERY, potential))
222
223
         plot_mesh(x_values, v_values)
224
225
226
     def plot_mesh(x_values, y_values):
227
         f = plt.figure()
228
         ax = f.gca()
229
         ax.set_aspect('equal', adjustable='box')
230
231
         x_range = []
         y_range = []
232
         for x in x_values[:-1]:
233
             for y in y_values[:-1]:
234
235
                 x_range.append(x)
236
                 y_range.append(y)
         plt.plot(x_range, y_range, 'o', label='Mesh points')
237
```

```
plt.xlabel('x')
238
239
         plt.ylabel('y')
         plt.grid(True)
240
         f.savefig('report/plots/q3e.pdf', bbox_inches='tight')
241
243
     def plot_sor_jacobi(h_values, potential_values, potential_values_jacobi, iterations_values,
244
         iterations_values_jacobi):
245
         f = plt.figure()
         plt.plot(h_values, potential_values, 'o-', label='SOR')
246
         plt.plot(h_values, potential_values_jacobi, 'o-', label='Jacobi')
247
         plt.xlabel('1 / h')
248
         plt.ylabel('Potential at [0.06, 0.04] (V)')
         plt.grid(True)
250
251
         plt.legend()
         f.savefig('report/plots/q3d_potential_comparison.pdf', bbox_inches='tight')
252
253
254
         f = plt.figure()
         plt.plot(h_values, iterations_values, 'o-', label='SOR')
255
         plt.plot(h_values, iterations_values_jacobi, 'o-', label='Jacobi')
256
257
         plt.xlabel('1 / h')
         plt.ylabel('Number of Iterations')
258
259
         plt.grid(True)
260
         plt.legend()
         f.savefig('report/plots/q3d_iterations_comparison.pdf', bbox_inches='tight')
261
262
263
     def save_rows_to_csv(filename, rows, header=None):
264
         with open(filename, "wb") as f:
265
             writer = csv.writer(f)
266
267
             if header is not None:
                 writer.writerow(header)
268
             for row in rows:
269
270
                 writer.writerow(row)
271
272
     def q3():
273
         o = q3b()
274
275
         h_values, potential_values, iterations_values = q3c(o)
276
         _, potential_values_jacobi, iterations_values_jacobi = q3d()
         plot_sor_jacobi(h_values, potential_values, potential_values_jacobi, iterations_values,
277
              iterations_values_jacobi)
         q3e()
278
279
     if __name__ == '__main__':
281
         t = time.time()
282
         q3()
283
         print('Total runtime: {} s'.format(time.time() - t))
284
```

# B Output Logs

```
Listing 8: Output of Question 1 program (q1.txt).
```

```
=== Question 1(b) ===
  n=2 matrix is positive-definite: True
   n=3 matrix is positive-definite: True
3
   n=4 matrix is positive-definite: True
    === Question 1(c) ===
6
   Matrix with n=2:
    Expected x:
      8.00
      3.00
   Actual x:
10
     8.00
11
12
      3.00
   Matrix with n=3:
```

```
14
    Expected x:
15
      9.00
      4.00
16
      3.00
17
    Actual x:
18
      9.00
19
20
      4.00
21
      3.00
    Matrix with n=4:
22
    Expected x:
      5.00
24
      4.00
25
      1.00
      9.00
27
28
    Actual x:
     5.00
29
      4.00
30
31
      1.00
     9.00
32
    === Question 1(d) ===
33
34
    Solved for x in network 1:
    V1 = 5.000 V
35
36
    Solved for x in network 2:
    V1 = 50.000 V
37
    Solved for x in network 3:
38
    V1 = 55.000 V
    Solved for x in network 4:
40
    V1 = 20.000 V
41
    V2 = 35.000 V
    Solved for x in network 5:
43
    V1 = 5.000 V
44
    V2 = 3.750 V
45
    V3 = 3.750 V
46
47
    Solved for x in network 6:
    V1 = 4.443 V
48
    V2 = 5.498 V
49
    V3 = 3.036 V
50
    V4 = 3.200 V
51
   V5 = 1.301 V
52
```

### Listing 9: Output of Question 2 program (q2. txt).

```
=== Question 2(a)(b) ===
2
    Equivalent resistance for 2x4 mesh: 1875.00 Ohms.
    Runtime: 0.000999927520752 s.
    Equivalent resistance for 3x6 mesh: 2379.55 Ohms.
    Runtime: 0.0169999599457 s.
    Equivalent resistance for 4x8 mesh: 2741.03 Ohms.
    Runtime: 0.100000143051 s.
    Equivalent resistance for 5x10 mesh: 3022.82 Ohms.
    Runtime: 0.481999874115 s.
9
    Equivalent resistance for 6x12 mesh: 3253.68 Ohms.
10
    Runtime: 1.46099996567 s.
    Equivalent resistance for 7x14 mesh: 3449.17 Ohms.
12
    Runtime: 3.26600003242 s.
13
    Equivalent resistance for 8x16 mesh: 3618.67 Ohms.
14
    Runtime: 7.53400015831 s.
15
16
    Equivalent resistance for 9x18 mesh: 3768.29 Ohms.
    Runtime: 15.001999855 s.
17
    Equivalent resistance for 10x20 mesh: 3902.19 Ohms.
18
    Runtime: 28.3630001545 s.
19
    === Question 2(c) ===
20
21
    Equivalent resistance for 2x4 mesh: 1875.00 Ohms.
    Runtime: 0.00100016593933 s.
22
    Equivalent resistance for 3x6 mesh: 2379.55 Ohms.
23
    Runtime: 0.0169999599457 s.
    Equivalent resistance for 4x8 mesh: 2741.03 Ohms.
25
    Runtime: 0.0950000286102 s.
26
    Equivalent resistance for 5x10 mesh: 3022.82 Ohms.
```

```
28
    Runtime: 0.378000020981 s.
    Equivalent resistance for 6x12 mesh: 3253.68 Ohms.
    Runtime: 1.19199991226 s.
30
    Equivalent resistance for 7x14 mesh: 3449.17 Ohms.
31
    Runtime: 3.05200004578 s.
32
    Equivalent resistance for 8x16 mesh: 3618.67 Ohms.
33
    Runtime: 6.9430000782 s.
34
35
    Equivalent resistance for 9x18 mesh: 3768.29 Ohms.
36
    Runtime: 14.2189998627 s.
    Equivalent resistance for 10x20 mesh: 3902.19 Ohms.
    Runtime: 26.763999939 s.
38
    === Question 2(d) ===
39
                             Listing 10: Output of Question 3 program (q3.txt).
    === Question 3(b) ===
1
2
    Omega: 1.0
    Quarter grid:
      0.00
            3.96
                   8.56 15.00 15.00 15.00 15.00
4
                   9.09 15.00 15.00 15.00
      0.00
            4.25
                                            15.00
5
            3.96 8.56 15.00 15.00 15.00 15.00
      0.00
            3.03
                   6.18
                         9.25 10.29
      0.00
                                      10.55
                                             10.29
      0.00
            1.97
                   3.88
                         5.53
                                6.37
                                       6.61
                                              6.37
      0.00
           0.96
                  1.86
                         2.61
                                3.04
                                       3.17
                                              3.04
            0.00
                  0.00
                               0.00
10
     0.00
                         0.00
                                       0.00
                                             0.00
    Num iterations: 32
11
    Potential at (0.06, 0.04): 5.526 V
12
    Omega: 1.1
14
    Quarter grid:
           3.96
                   8.56 15.00 15.00 15.00 15.00
     0.00
15
      0.00
           4.25
                   9.09 15.00 15.00 15.00
                                            15.00
                         15.00
      0.00
            3.96
                   8.56
                               15.00
                                       15.00
                                             15.00
17
           3.03
18
      0.00
                   6.18
                         9.25 10.29
                                      10.55
                                             10.29
      0.00
           1.97
                   3.88
                         5.53
                                6.37
                                       6.61
                                              6.37
      0.00
            0.96
                   1.86
                          2.61
                                3.04
                                       3.17
                                              3.04
20
21
     0.00
           0.00
                   0.00
                         0.00
                                0.00
                                       0.00
                                              0.00
    Num iterations: 26
    Potential at (0.06, 0.04): 5.526 V
23
    Omega: 1.2
24
    Quarter grid:
25
           3.96
                  8.56 15.00 15.00 15.00 15.00
     0.00
26
27
      0.00
            4.25
                   9.09
                         15.00
                               15.00
                                      15.00
                                             15.00
      0.00 3.96 8.56 15.00 15.00 15.00
                                             15.00
28
           3.03
29
      0 00
                  6.18
                         9.25 10.29 10.55
                                             10.29
      0.00
            1.97
                   3.88
                         5.53
                                6.37
                                       6.61
30
                                              6.37
      0.00 0.96
                   1.86
                         2.61
                                3.04
                                       3.17
                                              3.04
31
      0.00 0.00
                   0.00
                         0.00
                               0.00
                                       0.00
                                              0.00
    Num iterations: 20
33
    Potential at (0.06, 0.04): 5.526 V
34
    Omega: 1.3
35
    Quarter grid:
36
                  8.56 15.00 15.00 15.00 15.00
37
     0.00
            3.96
      0.00
           4.25 9.09 15.00 15.00 15.00 15.00
38
      0.00
            3.96
                   8.56 15.00 15.00
                                      15.00
                                             15.00
39
      0.00
            3.03
                   6.18
                         9.25
                               10.29
                                      10.55
                                             10.29
40
           1.97
                                6.37
      0.00
                   3.88
                         5.53
                                       6.61
                                              6.37
41
      0.00
            0.96
                   1.86
                         2.61
42
                                3.04
                                       3.17
                                              3.04
43
     0.00
            0.00
                   0.00
                         0.00
                                0.00
                                       0.00
                                              0.00
    Num iterations: 14
44
    Potential at (0.06, 0.04): 5.526 V
45
    Omega: 1.4
46
    Quarter grid:
47
           3.96
      0.00
                   8.56 15.00 15.00 15.00 15.00
                               15.00
                   9.09
                         15.00
      0.00
            4.25
                                       15.00
49
      0.00
            3.96
                   8.56 15.00 15.00
                                      15.00
                                             15.00
50
      0.00
           3.03
                   6.18
                         9.25 10.29 10.55 10.29
51
```

0.00

52

53

1.97

0.00 0.96

3.88

1.86

0.00 0.00 0.00 0.00

5.53

2.61

6.37

3.04

0.00

6.61

3.17

0.00

6.37

3.04

0.00

```
55
    Num iterations: 16
    Potential at (0.06, 0.04): 5.526 V
     Omega: 1.5
57
    Quarter grid:
58
                   8.56 15.00 15.00 15.00 15.00
      0.00
            3.96
 59
      0.00
             4.25 9.09 15.00 15.00 15.00 15.00
60
            3.96 8.56 15.00 15.00 15.00 15.00
      0.00
61
62
      0.00
             3.03
                   6.18
                          9.25 10.29
                                       10.55
                                             10.29
            1.97
      0.00
                               6.37
                                       6.61
                                             6.37
63
                   3.88
                         5.53
                                      3.17
      0.00 0.96 1.86
                         2.61 3.04
                                              3.04
 64
            0.00
                   0.00
      0.00
                         0.00
                               0.00
                                       0.00
                                              0.00
65
    Num iterations: 20
66
    Potential at (0.06, 0.04): 5.526 V
    Omega: 1.6
68
69
    Quarter grid:
      0.00 3.96 8.56 15.00 15.00 15.00 15.00
70
                   9.09 15.00 15.00 15.00
            4.25
      0.00
                                             15.00
71
                   8.56 15.00 15.00
 72
      0.00
            3.96
                                      15.00
                                             15.00
73
      0.00
            3.03
                   6.18
                         9.25 10.29 10.55
            1.97
      0.00
                   3.88
                          5.53
                                6.37
                                       6.61
                                              6.37
 74
 75
      0.00
            0.96
                   1.86
                          2.61
                                 3.04
                                       3.17
                                              3.04
      0.00 0.00 0.00 0.00 0.00
                                      0.00
                                             0.00
76
77
    Num iterations: 27
    Potential at (0.06, 0.04): 5.526 V
78
    Omega: 1.7
79
    Quarter grid:
 80
      0.00
            3.96
                   8.56 15.00 15.00 15.00 15.00
81
                  9.09 15.00 15.00 15.00 15.00
      0.00
            4.25
82
      0.00 3.96 8.56 15.00 15.00 15.00 15.00
 83
      0.00
             3.03
                   6.18
                          9.25 10.29
                                      10.55
                                             10.29
84
            1.97
                         5.53
      0.00
85
                   3.88
                                6.37
                                       6.61
                                              6.37
      0.00 0.96 1.86
                         2.61
                               3.04
                                       3.17
                                              3.04
86
      0.00
            0.00
                   0.00
                         0.00
                               0.00
                                       0.00
                                              0.00
87
 88
    Num iterations: 39
    Potential at (0.06, 0.04): 5.526 V
89
90
    Omega: 1.8
     Quarter grid:
91
      0.00 3.96 8.56 15.00 15.00 15.00 15.00
92
      0.00
            4.25 9.09 15.00 15.00 15.00 15.00
93
94
      0.00
             3.96
                   8.56 15.00 15.00
                                      15.00
                                             15.00
      0.00 3.03 6.18
                         9.25 10.29 10.55
                                             10.29
95
            1.97
                               6.37
96
      0.00
                   3.88
                         5.53
                                       6.61
                                              6.37
      0.00
             0.96
                   1.86
                          2.61
                                 3.04
                                       3.17
97
                                              3.04
                         0.00 0.00
      0.00 0.00
                  0.00
                                       0.00
                                             0.00
98
    Num iterations: 60
    Potential at (0.06, 0.04): 5.526 V
100
    Omega: 1.9
101
    Quarter grid:
102
      0.00 3.96
0.00 4.25
                   8.56 15.00 15.00 15.00 15.00
9.09 15.00 15.00 15.00 15.00
103
104
      0.00
            3.96 8.56 15.00 15.00 15.00 15.00
105
      0.00
            3.03
                   6.18
                          9.25 10.29 10.55
                                             10.29
106
107
      0.00
             1.97
                   3.88
                          5.53
                                6.37
                                       6.61
                                              6.37
      0.00
            0.96
                   1.86 2.61
                                3.04
                                       3.17
                                              3.04
108
     0.00 0.00 0.00 0.00 0.00
109
                                       0.00
                                             0.00
    Num iterations: 127
110
    Potential at (0.06, 0.04): 5.526 V
111
112
    Best number of iterations: 14
113
    Best omega: 1.3
    === Question 3(c): SOR ===
114
    h: 0.02
    1/h: 50.0
116
    Num iterations: 14
117
    Potential at (0.06, 0.04): 5.526 V
    h: 0.01
119
120
    1/h: 100.0
121
    Num iterations: 59
    Potential at (0.06, 0.04): 5.351 V
122
    h: 0.005
123
    1/h: 200.0
124
```

```
125
    Num iterations: 189
    Potential at (0.06, 0.04): 5.289 V
126
    h: 0.0025
127
    1/h: 400.0
128
    Num iterations: 552
    Potential at (0.06, 0.04): 5.265 V
130
131
    h: 0.00125
132
    1/h: 800.0
    Num iterations: 1540
133
    Potential at (0.06, 0.04): 5.254 V
    h: 0.000625
135
    1/h: 1600.0
136
    Num iterations: 4507
    Potential at (0.06, 0.04): 5.247 V
138
139
    === Question 3(d): Jacobi ===
140
    h: 0.02
    Num iterations: 51
141
    Potential at (0.06, 0.04): 5.526 V
142
    h: 0.01
143
    Num iterations: 180
144
    Potential at (0.06, 0.04): 5.351 V
    h: 0.005
146
147
    Num iterations: 604
    Potential at (0.06, 0.04): 5.289 V
148
    h: 0.0025
149
    Num iterations: 1935
150
    Potential at (0.06, 0.04): 5.265 V
151
    h: 0.00125
152
    Num iterations: 5836
    Potential at (0.06, 0.04): 5.254 V
154
155
    h: 0.000625
    Num iterations: 16864
    Potential at (0.06, 0.04): 5.246 V
157
158
    Total runtime: 1724.82099986
     === Question 3(e): Non-Uniform Node Spacing ===
159
    Jacobi (for reference)
160
    Quarter grid:
161
     0.00 1.99
                   4.06 6.29 8.78 11.66 15.00 15.00 15.00 15.00 15.00 15.00
162
            2.03
                   4.14
      0.00
                          6.41 8.95 11.82 15.00 15.00 15.00 15.00 15.00 15.00
163
164
      0.00
             1.99
                   4.06
                          6.29
                                 8.78
                                       11.66
                                             15.00
                                                    15.00
                                                           15.00
                                                                  15.00
                                                                         15.00
      0.00
            1.87
                          5.89
                                 8.23 11.04 15.00 15.00 15.00 15.00
                   3.81
                                                                        15.00 15.00
165
      0.00
            1.69
                   3.42
                          5.24
                                 7.19
                                       9.28 11.33 12.14 12.50 12.66
                                                                        12.71 12.66
166
      0.00
                                        7.55
                                              8.90
                                                     9.73
                                                           10.20
                                                                  10.44
             1.46
                    2.95
                          4.47
                                 6.02
                                                                         10.51
167
            1.22
                                              6.99
                                                     7.69
      0.00
                   2.44
                          3.66
                                 4.87
                                       6.01
                                                           8.14
                                                                  8.38
                                                                         8.45
168
      0.00 0.96 1.92
                          2.87
                                 3.78
                                       4.63
                                              5.35
                                                     5.90
                                                            6.27
                                                                  6.48
                                                                         6.55
                                                                                6.48
      0.00
             0.71
                    1.42
                          2.11
                                 2.77
                                        3.37
                                               3.89
                                                     4.29
                                                            4.57
                                                                   4.73
                                                                         4.79
                                                                                4.73
170
      0.00
            0.47
                   0.94
                                 1.81
                                       2.20
                                              2.53
                                                     2.80
                                                            2.98
                                                                         3.13
171
                          1.39
                                                                  3.09
                                                                                3.09
      0.00
            0.23 0.46
                          0.69
                                0.90
                                       1.09
                                              1.25
                                                     1.38
                                                            1.47
                                                                  1.53
                                                                        1.55
                                                                                1.53
172
      0.00
            0.00
                   0.00
                          0.00
                                0.00
                                       0.00
                                              0.00
                                                     0.00
                                                            0.00
                                                                  0.00
                                                                         0.00
                                                                                0.00
173
174
    Num iterations: 106
     Potential at (0.06, 0.04): 5.351 V
175
    Uniform Mesh (same as Jacobi)
176
177
    Quarter grid:
      0.00 1.99
                   4.06
                         6.29
                                8.78 11.66 15.00 15.00 15.00 15.00 15.00 15.00
178
             2.03
                   4.14
                                 8.95 11.82 15.00 15.00 15.00 15.00
179
      0.00
                          6.41
                                                                        15.00 15.00
      0.00
             1.99
                    4.06
                          6.29
                                 8.78
                                       11.66
                                             15.00 15.00
                                                           15.00
                                                                  15.00
                                                                        15.00 15.00
180
      0.00
            1.87
                          5.89
                                 8.23 11.04 15.00 15.00 15.00 15.00
                    3.81
                                                                        15.00 15.00
181
            1.69
      0.00
                   3.42
                          5.24
                                 7.19
                                       9.28 11.33 12.14 12.50 12.66 12.71 12.66
182
      0.00
                    2.95
                          4.47
                                        7.55
                                              8.90
                                                     9.73
                                                           10.20
                                                                  10.44
183
             1.46
                                 6.02
                                                                         10.51
      0.00
            1.22
                   2.44
                          3.66
                                 4.87
                                       6.01
                                              6.99
                                                     7.69
                                                           8.14
                                                                  8.38
                                                                         8.45
184
      0.00
            0.96
                   1.92
                          2.87
                                 3.79
                                       4.63
                                              5.35
                                                     5.90
                                                            6.27
                                                                  6.48
                                                                         6.55
                                                                                6.48
185
      0.00
             0.71
                    1.42
                          2.11
                                 2.77
                                        3.37
                                               3.89
                                                     4.29
                                                            4.57
                                                                   4.73
                                                                         4.79
186
      0.00
             0.47
                    0.94
                                 1.81
                                       2.20
                                              2.53
                                                     2.80
                                                            2.98
                                                                   3.09
                                                                         3.13
                                                                                3.09
187
                          1.39
      0.00
            0.23 0.46
                         0.69
                                0.90
                                       1.09
                                              1.25
                                                     1.38
                                                            1.47
                                                                   1.53
                                                                         1.55
                                                                                1.53
188
      0.00
            0.00
                    0.00
                          0.00
                                 0.00
                                       0.00
                                              0.00
                                                     0.00
                                                            0.00
                                                                   0.00
                                                                         0.00
189
    Num iterations: 209
190
    Potential at (0.06, 0.04): 5.351 V
191
    Jacobi potential: 5.35062156679 V, same as uniform potential: 5.35067998265 V
192
    Non-Uniform (clustered around (0.06, 0.04))
    Quarter grid:
194
```

```
4.08
                           6.33 11.61 13.25 15.00 15.00 15.00 15.00 15.00 15.00
195
       0.00
             2.00
       0.00
              2.04
                     4.17
                            6.45
                                 11.80
                                        13.37
                                               15.00
                                                      15.00
                                                             15.00
                                                                     15.00
                                                                            15.00
                                                                                  15.00
196
       0.00
              2.00
                     4.08
                            6.33
                                 11.61
                                        13.25
                                               15.00 15.00
                                                             15.00
                                                                    15.00
                                                                            15.00 15.00
197
       0.00
              1.89
                     3.84
                            5.93
                                 10.90
                                        12.71
                                               15.00
                                                      15.00
                                                             15.00
                                                                    15.00
                                                                            15.00 15.00
198
       0.00
              1.71
                     3.45
                            5.28
                                  9.27
                                         10.26
                                                11.15
                                                       11.74
                                                              12.14
                                                                     12.66
                                                                            12.71
                                                                                  12.66
199
       0.00
              1.21
                     2.43
                            3.66
                                   6.06
                                         6.57
                                                7.03
                                                       7.42
                                                              7.75
                                                                     8.38
                                                                            8.45
200
                                                       6.52
201
       0.00
             1.09
                     2.18
                           3.26
                                  5.35
                                         5.78
                                                6.18
                                                              6.81
                                                                     7.41
                                                                            7.48
                                                                                   7.41
202
       0.00
              0.96
                     1.92
                            2.87
                                   4.66
                                         5.04
                                                 5.38
                                                       5.67
                                                               5.93
                                                                      6.48
                                                                             6.55
                                                       4.87
       0.00
              0.84
                    1.67
                            2.48
                                   4.01
                                         4.33
                                                4.62
                                                              5.09
                                                                            5.65
203
                                                                     5.59
                                                                                   5.59
       0.00
              0.71
                    1.42
                           2.11
                                   3.39
                                         3.65
                                                3.89
                                                       4.11
                                                               4.29
                                                                      4.72
                                                                            4.77
                                                                                    4.72
204
       0.00
              0.23
                     0.47
                            0.69
                                   1.10
                                          1.19
                                                 1.26
                                                        1.33
                                                               1.39
                                                                      1.54
                                                                             1.56
                                                                                    1.54
205
                                                       0.00
                                                              0.00
                                                                            0.00
       0.00
             0.00
                     0.00
                           0.00
                                  0.00
                                         0.00
                                                0.00
                                                                     0.00
                                                                                   0.00
206
     Num iterations: 385
207
     Potential at (0.06, 0.04): 5.378 V
208
     Non-Uniform (more clustered around (0.06, 0.04))
209
210
     Quarter grid:
                           6.41 13.24 14.65 15.00 15.00 15.00 15.00 15.00 15.00
       0.00
             2.03
                    4.14
211
                           6.53 13.40
212
       0.00
              2.07
                     4.22
                                        14.68
                                               15.00
                                                      15.00
                                                             15.00
                                                                    15.00
                                                                            15.00
                                                                                  15.00
       0.00
              2.03
                     4.14
                            6.41 13.24 14.65 15.00 15.00 15.00
                                                                    15.00
                                                                            15.00 15.00
213
       0.00
             1.92
                     3.90
                           6.02 12.55 14.45
                                               15.00
                                                      15.00
                                                             15.00
                                                                    15.00
                                                                            15.00
                                                                                  15.00
214
215
       0.00
              1.73
                     3.51
                            5.36
                                 10.40
                                         11.09
                                                11.24
                                                       11.38
                                                              11.86
                                                                     12.65
                                                                            12.71
                                                                                  12.65
       0.00
              1.10
                     2.19
                           3.28
                                  5.90
                                         6.21
                                                6.29
                                                       6.36
                                                              6.62
                                                                     7.44
                                                                            7.51
                                                                                   7.44
216
217
       0.00
             1.00
                    1.99
                           2.97
                                  5.28
                                         5.56
                                                5.62
                                                       5.69
                                                              5.92
                                                                     6.69
                                                                            6.75
                                                                                   6.69
       0.00
              0.97
                     1.94
                            2.89
                                   5.13
                                          5.40
                                                 5.46
                                                        5.52
                                                              5.75
                                                                      6.50
218
                                                                             6.57
                                                                                    6.50
       0.00
              0.94
                    1.88
                            2.81
                                   4.98
                                          5.24
                                                5.30
                                                       5.36
                                                              5.58
                                                                      6.32
                                                                            6.38
                                                                                    6.32
219
220
       0.00
              0.84
                    1.68
                           2.50
                                  4.39
                                         4.62
                                                4.68
                                                       4.73
                                                              4.92
                                                                      5.60
                                                                            5.66
                                                                                   5.60
                     0.47
       0.00
              0.24
                            0.70
                                   1.21
                                          1.28
                                                 1.29
                                                        1.31
                                                               1.36
                                                                      1.56
                                                                             1.57
                                                                                    1.56
221
       0.00
             0.00
                    0.00
                                  0.00
                                         0.00
                                                0.00
                                                       0.00
                                                              0.00
                                                                     0.00
                                                                            0.00
                                                                                   0.00
                           0.00
222
     Num iterations: 1337
     Potential at (0.06, 0.04): 5.461 V
224
     Non-Uniform (clustered near outer conductor)
225
     Quarter grid:
226
       0.00
             4.38
                                                8.97
                                                       9.82 10.43
                     7.21 10.30 13.47
                                         7.42
                                                                    10.80
                                                                           10.86
                                                                                   7.63
227
                           10.46 13.55 15.00 15.00 15.00
228
       0.00
             4.46
                     7.34
                                                             15.00
                                                                    15.00
                                                                            15.00
                                                                                  15.00
       0.00
                    7.21
                           10.30 13.47 15.00 15.00 15.00
             4.38
                                                             15.00
                                                                    15.00
                                                                            15.00 15.00
229
                                               15.00
                                                      15.00
       0.00
             4.19
                     6.91
                           9.94 13.24
                                        15.00
                                                             15.00
                                                                    15.00
                                                                            15.00 15.00
230
       0.00
              3.95
                     6.50
                           9.37
                                 12.69
                                         15.00
                                               15.00
                                                      15.00
                                                             15.00
                                                                     15.00
                                                                            15.00 15.00
231
       0.00
              3.61
                     5.91
                            8.39
                                 10.87
                                        11.93
                                               12.87 13.10
                                                             13.22
                                                                    13.30
                                                                            13.33 13.30
232
                           7.16
                                  8.96
       0.00
                                         9.63
                                               10.73 11.09
233
              3.18
                     5.15
                                                             11.29
                                                                    11.43
                                                                           11.49 11.43
234
       0.00
              2.67
                     4.27
                            5.84
                                   7.16
                                         7.66
                                                8.66
                                                       9.03
                                                              9.27
                                                                     9.44
                                                                            9.51
       0.00
              1.89
                     3.00
                           4.05
                                          5.24
                                                5.99
                                                       6.29
                                                              6.49
                                                                            6.71
                                   4.91
                                                                     6.64
                                                                                   6.64
235
236
       0.00
             1.50
                    2.36
                           3.17
                                   3.83
                                         4.09
                                                4.69
                                                       4.94
                                                              5.11
                                                                     5.23
                                                                            5.29
                                                                                   5.23
       0.00
                     1.44
                                          2.49
                                                 2.86
                                                        3.02
                                                              3.13
                                                                      3.21
237
              0.92
                           1.93
                                   2.33
                                                                             3.25
                                                                                    3.21
                                                0.00
                                                       0.00
                                                              0.00
       0.00
             0.00
                     0.00
                           0.00
                                  0.00
                                         0.00
                                                                     0.00
                                                                            0.00
                                                                                   0.00
238
     Num iterations: 222
    Potential at (0.06, 0.04): 5.243 V
240
```