

Assignment 5: The Resistor Problem

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March 3, 2020

0.1 Introduction

This report will discuss about the solver for the currents in a resistor and discusses about the current's dependency on the shape of the resistor and also discusses which part of the resistor is likely to get the hottest. Here we analyse the currents in a square copper plate to where a wire is soldered to its centre. It also discusses about how to find the stopping condition for the solver after certain iterations based on an error tolerance, and to model the errors obtained using Least Squares after analysing the actual errors in semilog and loglog plots. Finally we find the currents in the resistor after applying boundary conditions and analyse the vector plot of current flow and conclude which part of resistor will become hot.

- A wire is soldered to the middle of a copper plate and its voltage is held at 1 Volt. One side of the plate is rounded, while the remaining are floating. The plate is 1 cm by 1 cm in size.
- To solve for currents in resistor, we use following equations and boundary conditions mentioned below:
- Conductivity (Differential form of ohm's law)

$$\vec{J} = \sigma \vec{E} \quad (1)$$

- Electric field is the gradient of the potential

$$\vec{E} = -\nabla\phi \quad (2)$$

- Charge Continuity equation is used to conserve the inflow and outflow charges

$$\nabla \cdot \vec{J} = -\frac{\partial \rho}{\partial t} \quad (3)$$

- Combining the above equations above, we get

$$\nabla \cdot (-\sigma \nabla \phi) = -\frac{\partial \rho}{\partial t} \quad (4)$$

- Assuming that our resistor contains a material of constant conductivity, the equation becomes

$$\nabla^2 \phi = \frac{1}{\sigma} \frac{\partial \rho}{\partial t} \quad (5)$$

- For DC currents, the right side is zero, and we obtain

$$\nabla^2 \phi = 0 \quad (6)$$

- Here we use a 2-D plate so the Numerical solutions in 2D can be easily transformed into a difference equation. The equation can be written out in

$$\frac{\partial^2 \phi}{\partial x^2} + \frac{\partial^2 \phi}{\partial y^2} = 0 \quad (7)$$

$$\frac{\partial \phi}{\partial x(x_i, y_j)} = \frac{\phi(x_{i+1/2}, y_j) - \phi(x_{i-1/2}, y_j)}{\Delta x} \quad (8)$$

$$\frac{\partial^2 \phi}{\partial x^2(x_i, y_j)} = \frac{\phi(x_{i+1}, y_j) - 2\phi(x_i, y_j) + \phi(x_{i-1}, y_j)}{(\Delta x)^2} \quad (9)$$

- Using above equations we get

$$\phi_{i,j} = \frac{\phi_{i+1,j} + \phi_{i-1,j} + \phi_{i,j+1} + \phi_{i,j-1}}{4} \quad (10)$$

- Thus, the potential at any point should be the average of its neighbours. This is a very general result and the above calculation is just a special case of it. So the solution process is to take each point and replace the potential by the average of its neighbours. Keep iterating till the solution converges (i.e., the maximum change in elements of ϕ which is denoted by $error_k$ in the code, where 'k' is the no of iteration, is less than some tolerance which is taken as 10^{-8}).
- At boundaries where the electrode is present, just put the value of potential itself. At boundaries where there is no electrode, the current should be tangential because charge can't leap out of the material into air. Since current is proportional to the Electric Field, what this means is the gradient of ϕ should be tangential. This is implemented by requiring that ϕ should not vary in the normal direction
- Finally, we solve for currents in the resistor using all this information

0.2 Python Code :

0.2.1 Question 1

Part A

- Define the Parameters, The default parameter values taken for my particular code were $N_x = 25$ and $N_y = 25$ and No of iterations : 1500. The user can change the N_x and N_y if he or she wishes to.
- These values are taken to discuss about Stopping condition,etc
- To allocate the potential array $\phi = 0$.Note that the array should have N_y rows and N_x columns.
- To find the indices which lie inside the circle of radius 0.35 using `meshgrid()` by equation :

$$X^2 + Y^2 \leq 0.35^2 \quad (11)$$

- Then assign 1 V to those indices.
- To plot a contour plot of potential ϕ and to mark V=1 region in red

```

1  #importing the necessary libraries
2  from pylab import *
3  import numpy as np
4  from numpy import *
5  import matplotlib.pyplot as plt
6  import mpl_toolkits.mplot3d.axes3d as p3
7
8
9  Niter=1500
10 Nx=int(input('What do you want Nx value to be?')) #taking Nx value from the u
11 Ny=Nx
12 Radius=Nx*8/25
13 phi=np.zeros((Ny,Nx)) #initialising phi matrix
14 y=linspace(-0.5,0.5,Ny)
15 x=linspace(-0.5,0.5,Nx)
16 X,Y=np.meshgrid(x,y)# creates a coordinate system
17 ii = where(square(X) + square(Y) <= pow(0.35, 2))
18 phi[ii] = 1.0 # assigning V=1 for the circular region
19
20 #plotting the potential
21 plt.contourf(phi, cmap=cm.jet)#cmap gives the colour(cmap.jet=default) contour
22 plt.title("Figure 1 : Contour Plot of  $\phi$ ")
23 ax = gca()
24 plt.colorbar( ax=ax, orientation='vertical')
25 plt.xlabel("x->")
26 plt.ylabel("y->")
27 plt.show()

```

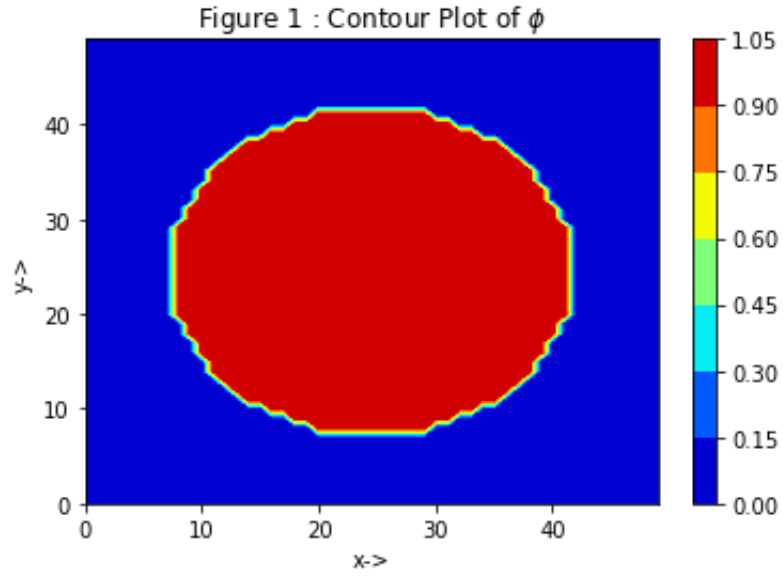


Figure 1: Contour plot of initial potential

Results and Discussion :

- The contour plot of potential becomes smoother i.e it almost becomes circular as we increase N_x and N_y , because we get more points, so the potential gradient is smoothed out between adjacent points since there are more points

Part B :

- To Perform the iterations
- To update the potential ϕ according to Equation below using vectorized code

$$\phi_{i,j} = \frac{\phi_{i+1,j} + \phi_{i-1,j} + \phi_{i,j+1} + \phi_{i,j-1}}{4} \quad (12)$$

- To apply Boundary Conditions where there is no electrode, the gradient of ϕ should be tangential. This is implemented by Equation given below , basically potential should not vary in the normal direction so we equate the last but row or column to outermost row or column correspondingly when applying boundary conditions for a side of plate,implemented using Vectorized code

$$\frac{\partial \phi}{\partial n} = 0 \quad (13)$$

- To plot the errors in semilog and loglog and observe how the errors are evolving.

```
1  # function to create Matrix for finding the Best fit using lstsq
2  # with no_of rows, columns by default and vector x as arguments
3
4  def createAmatrix(nrow, x):
5      A = zeros((nrow, 2)) # allocate space for A
6      A[:, 0] = 1
7      A[:, 1] = x
8      return A
9
10
11 # function to find best fit errors using lstsq
12 def fitForError(errors, x):
13     A = createAmatrix(len(errors), x)
14     return A, np.linalg.lstsq(A, np.log(errors))[0]
15
16 # Function to compute function back from Matrix and Coefficients A and B
17 def computeErrorFit(M, c):
```

```

18     return np.exp(dot(M,c))
19
20 errors = zeros(Niter)  # initialise error array to zeros
21 iterations = []  # array from 0 to Niter used for findind lstsq
22
23 for k in range(Niter):
24     # copy the old phi
25     oldphi = phi.copy()
26     # Updating the potential
27     phi[1:-1, 1:-1] = 0.25 *(phi[1:-1, 0:-2]+phi[1:-1, 2:]+phi[0:-2, 1:-1]+phi[2:, 1:-1])
28     # applying boundary conditions: constant in the normal direction
29     phi[:, 0] = phi[:, 1]  # Left edge
30     phi[:, -1] = phi[:, -2]  # right edge
31     phi[0, :] = phi[1, :]  # Top edge
32     # Bottom edge is grounded so no boundary conditions
33     # Assign 1 V to electrode region
34     phi[ii] = 1.0
35     # Appending errors for each iterations
36     errors[k] = (abs(phi-oldphi)).max()
37     iterations.append(k)
38
39 plt.semilogy(iterations[0::50], errors[0::50], 'go', markersize=8, label="Original Error")
40 plt.semilogy(iterations, errors, markersize=8, label="Original Error cont",color='red')
41 plt.legend()
42 plt.title("Figure 2a : Error Vs No of iterations (Semilog)")
43 plt.xlabel("Niter ->")
44 plt.ylabel("Error ->")
45 plt.grid()
46 plt.show()
47
48 plt.loglog(iterations[0::50], errors[0::50], 'go', markersize=8, label="Original Error")
49 plt.loglog(iterations, errors, markersize=8, label="Original Error",color='black')
50 plt.legend()
51 plt.title("Figure 2b : Error Vs No of iterations (Loglog)")
52 plt.xlabel("Niter ->")

```



```
53 plt.ylabel("Error ->")
54 plt.grid()
55 plt.show()
```

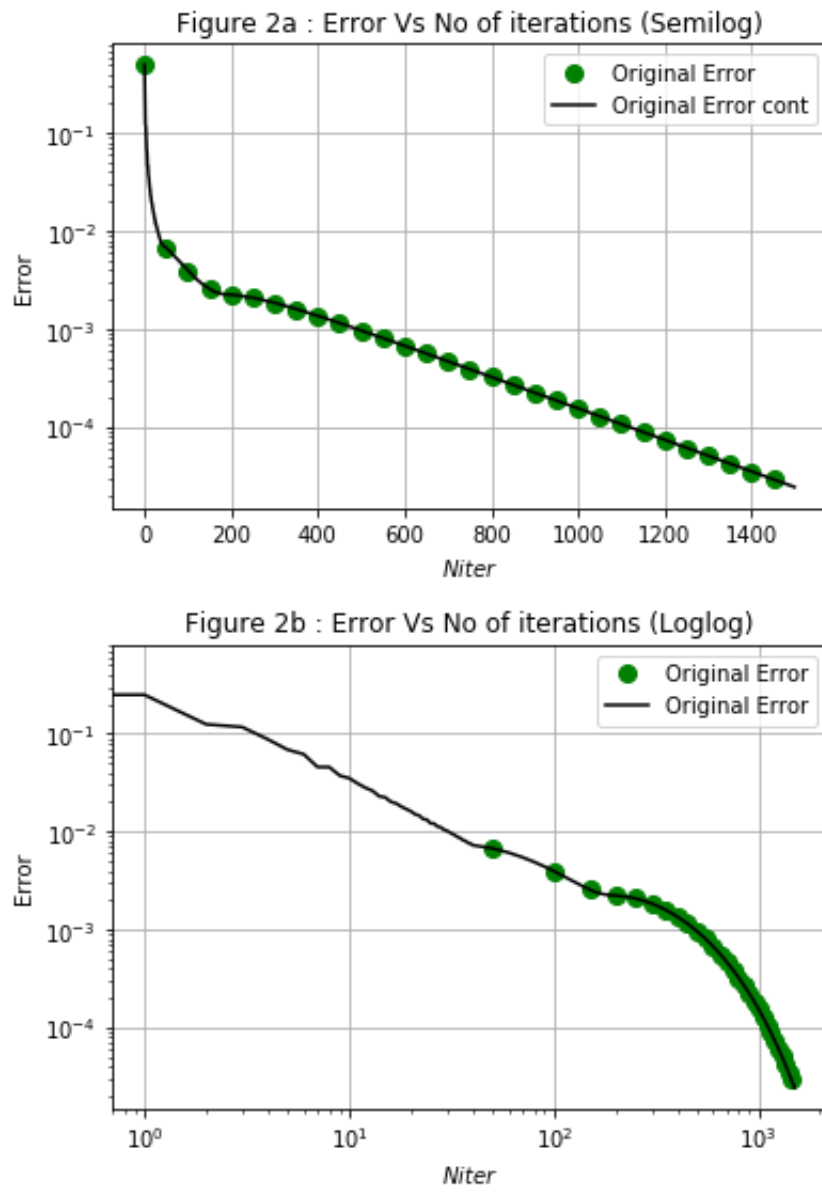


Figure 2: Normal and Log-Log plot of Error vs No.of Iterations

Results and Discussion:

- As we observe the Figure 2a that error decreases linearly for higher no of iterations,so from this we conclude that for large iterations error decreases exponentially with No of iterations i.e it follows Ae^{Bx} as it is a semilog plot
- And if we observe loglog plot the error is almost linearly decreasing for smaller no of iterations so it follows a^x form since it is loglog plot and follows some other pattern at larger iterations.
- So to conclude the error follows Ae^{Bx} for higher no of iterations(≈ 500) and it follows a^x form for smaller iterations which can be seen from figure 2a & 2b respectively

Part C :

- To find the fit using Least squares for all iterations named as **fit1** and for iterations ≥ 500 named as **fit2** separately and compare them.
- As we know that error follows Ae^{Bx} at large iterations, we use equation given below to fit the errors using least squares

$$\log y = \log A + Bx \quad (14)$$

- To find the time constant of error function obtained for the two cases using lstsq and compare them
- To plot the two fits obtained and observe them

```
1  # to find the coefficients of fit1 and fit2
2  # M1 and M2 are matrices and c1 and c2 are coefficients
3  M1, c1 = fitForError(errors, iterations) # fit1
4  M2, c2 = fitForError(errors[500:], iterations[500:]) # fit2
5
6  print("Fit1 : A = %g , B = %g" % ((np.around(np.exp(c1[0]),4), np.around(c1[1],4)
7  print("Fit2 : A = %g , B = %g" % ((np.around(np.exp(c2[0]),4), np.around(c2[1],4)
8
9  print("The time Constant (1/B) all iterations considered: %g" % (np.around(1/c1[1],4)
10 print("The time Constant (1/B) for higher iterations (from 500) : %g" % (np.around(1/c2[1],4)
11
12
13 # Calculating the fit using Matrix M and Coefficients C obtained
14 error_fit1 = computeErrorFit(M1, c1) # fit1
15 M2new = createAmatrix(len(errors), iterations)
16
17 # Error calculated for all iterations using coefficients found using lstsq
18 error_fit2 = computeErrorFit(M2new, c2) # fit2 calculated
19
20 # Plotting the estimated error_fits using lstsq
21 # plotted for every 200 points for fit1 and fit2
22
23 plt.semilogy(iterations[0::200], error_fit1[0::200], 'ro', markersize=8, label=
```

```
24 plt.semilogy(iterations[0::200], error_fit2[0::200], 'bo', markersize=6, label=
25 plt.semilogy(iterations, errors, 'k', markersize=6, label="Actual Error")
26 plt.legend()
27 plt.title(r"Figure 3 : Error Vs No of iterations (Semilog)")
28 plt.xlabel("Niter ->")
29 plt.ylabel("Error ->")
30 plt.grid()
31 plt.show()
```

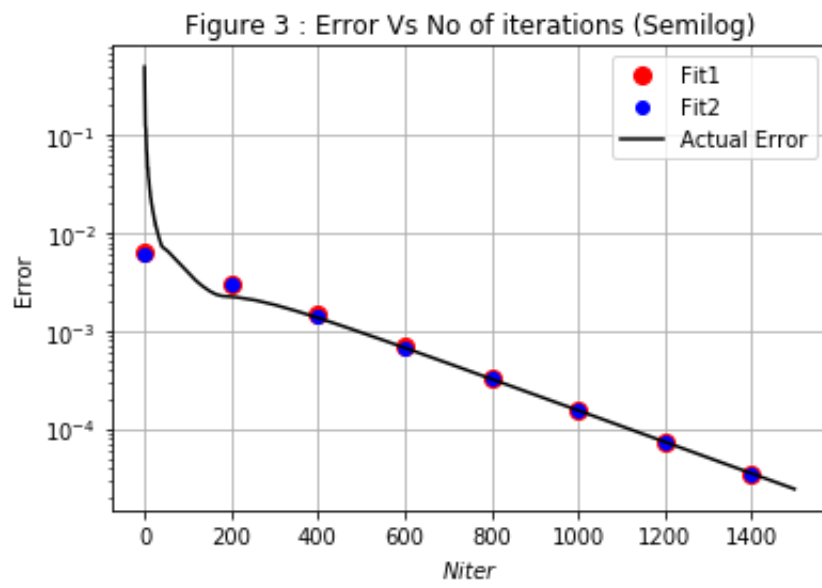


Figure 3: Semilog plot of Error vs No.of Iterations

Results and Discussion:

- Fit1 : $A = 0.0065$, $B = -0.0038$
Fit2 : $A = 0.0062$, $B = -0$
- As we observe the Fit1's time constant and Fit2's time constant, Fit2's is slightly higher than Fit1's time constant, so the error decreases slowly at larger iterations compared to fit1.
- Ideally the time constant for Fit2 should be larger than Fit1 with good margin, when we take less no of points i.e stepsize N_x and N_y being less, we get less difference between their time constants. If we increase the N_x and N_y to 100,100 respectively, we get these results (Niter=6000):
 - Time Constant for Fit1 : 1120.02s
 - Time Constant for Fit2 (higher iterations from 500) : 1189.62s
- We see that there is a significance difference between them.
- So the time constant increase with increase in N_x and N_y

Stopping Condition :

- To find the cumulative error for all iterations and compare them with some error tolerance to stop the iteration.
- So to find the cumulative error, we add all the absolute values of errors for each iteration since worst case is, all errors add up
- So we use the equations given below:

$$Error = \sum_{N+1}^{\infty} error_k \quad (15)$$

- The above error is approximated to

$$Error \approx -\frac{A}{B} \exp(B(N + 0.5)) \quad (16)$$

where N is no of iteration

```
1  #function calculating cumulative error
2  def cumerror(error, N, A, B):
3      return -(A/B)*exp(B*(N+0.5))
4
5  #finds the stopping condition
6  def findStopCondn(errors, Niter, error_tol):
7      cum_error = []
8      for n in range(1, Niter):
9          cum_error.append(cumerror(errors[n], n, exp(c1[0]), c1[1]))
10         if(cum_error[n-1] <= error_tol):
11             print("last per-iteration change in the error is %g"
12                   % (np.around(cum_error[-1]-cum_error[-2],4)))
13             return cum_error[n-1], n    #reduces the number of iterations base
14
15
16
17     print("last per-iteration change in the error is %g"
18           % (np.abs(cum_error[-1]-cum_error[-2])))
19     return cum_error[-1], Niter
```



```
20
21 #error tolerance
22 error_tol = pow(10, -8)
23 cum_error, Nstop = findStopCondN(errors, Niter, error_tol)
24 print("Stopping Condition N: %g and Error is %g" % (Nstop, cum_error))
```

Results and Discussion :

- From running the code, we get stopping condition as N : 6000 and the total cumulative error till that iteration is 0.00821977
- And the last per iteration change in error: $7.3422 * 10^{-6}$
- So we observe that the profile was changing very little every iteration, but it was continuously changing. So the cumulative error was still large.
- So that is why this method of solving Laplace's Equation is known to be one of the worst available. This is because of the very slow coefficient with which the error reduces.

Part D: Surface Plot of Potential

- To do a 3-D surface plot of the potential.
- To plot contour plot of potential
- And analyse them and to comment about flow of currents

```
1  fig1 = plt.figure()  # open a new figure
2  ax1= p3.Axes3D(fig1)  # Axes3D is the means to do a surface plot
3  plt.title("Figure 4: 3-D surface plot of the potential  $\phi$ ")
4  surf = ax1.plot_surface(X, Y, phi, rstride=1, cstride=1, cmap=cm.jet)
5  ax1.set_xlabel('x')
6  ax1.set_ylabel('y')
7  ax1.set_zlabel('z')
8  cax = fig1.add_axes([1, 0, 0.1, 1])
9  fig1.colorbar(surf, cax=cax, orientation='vertical')
10 plt.show()
11
12
13 ##### Contour Plot of the Potential:
14 plt.contourf(X, Y, phi, cmap=cm.jet)
15 plt.title("Figure 6 : Contour plot of Updated potential  $\phi$ ")
16 ax = gca()
17 plt.colorbar(plt6, ax=ax, orientation='vertical')
18 plt.xlabel("x ->")
```

```
19 plt.ylabel("y ->")
20 plt.grid()
21 plt.show()
```

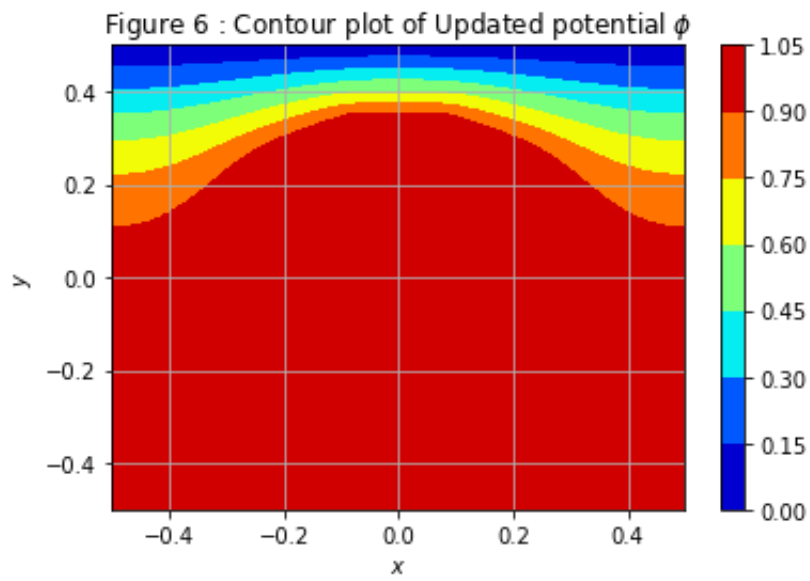
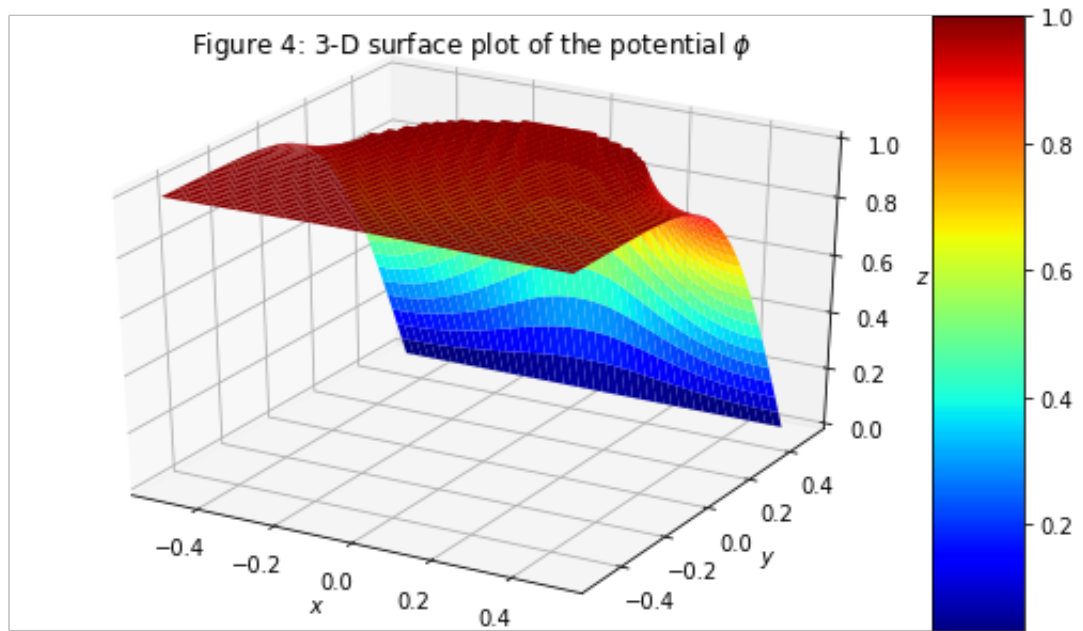


Figure 4: 3-D Surface potential plot and Contour plot of potential

Results and Discussion:

- As we observe that the surface plot we conclude that after updating the potential, the potential gradient is higher in down part of the plate since, the down side is grounded and the electrode is at 1 V, so there is high potential gradient from electrode to grounded plate.
- And the upper part of the plate is almost 1 V since they didn't have forced Voltage and they were floating, so while applying updating we replaced all points by average of surrounding points so the potential is almost 1 V in the upper region of the plate.
- Same observation we see using contour plot in 2 dimensions, we note that there are gradients in down part of the plate and almost negligible gradient in upper part of the plate.

Part E : Vector Plot of Currents :

- To obtain the currents by computing the gradient.
- The actual value of σ does not matter to the shape of the current profile, so we set it to unity. Our equations are

$$J_x = -\frac{\partial\phi}{\partial x} \quad (17)$$

$$J_y = -\frac{\partial\phi}{\partial y} \quad (18)$$

- To program this we use these equations as follows:

$$J_{x,ij} = \frac{1}{2}(\phi_{i,j-1} - \phi_{i,j+1}) \quad (19)$$

$$J_{y,ij} = \frac{1}{2}(\phi_{i-1,j} - \phi_{i+1,j}) \quad (20)$$

```
1 Jx = zeros((Ny, Nx))
2 Jy = zeros((Ny, Nx))
3 #from differential equation for current
4 Jx[1:-1, 1:-1] = 0.5*(phi[1:-1, 0:-2] - phi[1:-1, 2:])
5 Jy[1:-1, 1:-1] = 0.5*(phi[2:, 1:-1] - phi[0:-2, 1:-1])
6
7
8 # ##### To Plot the current density using quiver, and mark the electrode via
9 plt.scatter(x[ii[0]], y[ii[1]], color='r', s=10, label="V = 1 region")#s is f
10 plt.quiver(X, Y, Jx[:, :-1, :], Jy[:, :-1, :])
11 plt.xlabel('x ->')
12 plt.ylabel('y ->')
13 plt.legend()
14 plt.title("The Vector plot of the current flow")
15 plt.show()
```

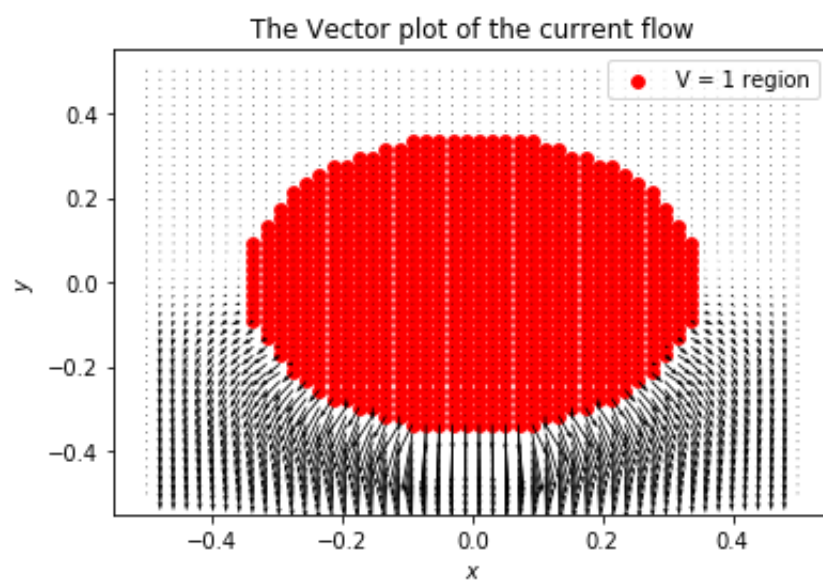


Figure 5: Vector plot of current flow

Results and Discussion:

- So as we noted that the potential gradient was higher in down region of the plate, and we know that Electric field is the gradient of the potential as given below

$$\vec{E} = -\nabla\phi \quad (21)$$

- So \vec{E} is larger where there is potential gradient is high and is inverted since it is negative of the gradient!, So it is higher in down region which is closer to bottom plate which is grounded
- And we know that

$$\vec{J} = \sigma\vec{E} \quad (22)$$

- So \vec{J} is higher and perpendicular to equipotential electrode region i.e "Red dotted region" so the current is larger in down part of the plate and perpendicular to the red dotted electrode region since $I = \vec{J} \cdot \vec{A}$
- So because of this most of the current flows from electrode to the bottom plate which is grounded because of higher potential gradient.
- And there is almost zero current in upper part of the plate since there is not much potential gradient as we observed from the surface and contour plot of the potential ϕ

0.3 Conclusion :

- To conclude , Most of the current is in the narrow region at the bottom. So that is what will get strongly heated.
- Since there is almost no current in the upper region of plate, the bottom part of the plate gets hotter and temperature increases in down region of the plate.
- And we know that heat generated is from $\vec{J} \cdot \vec{E}$ (ohmic loss) so since \vec{J} and \vec{E} are higher in the bottom region of the plate, there will more heat generation and temperature rise will be present.
- So overall we looked the modelling of the currents in resistor in this report ,and we observe that the best method to solve this is to increase N_x and N_y to very high values(100 or ≥ 100) and increase the no of iterations too, so that we get accurate answers i.e currents in the resistor.

- But the tradeoff is this method of solving is very slow even though we use vectorized code because the decrease in errors is very slow w.r.t no of iterations.