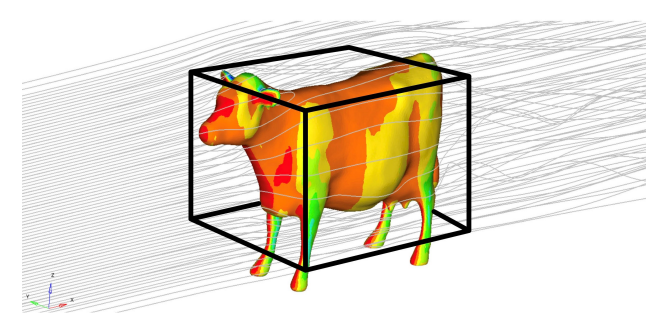
Project 1: Flow Over a Square

Aerospace 523: Computational Fluid Dynamics I Graduate Aerospace Engineering University of Michigan, Ann Arbor

> By: Dan Card*, dcard@umich.edu Date: October 9, 2020



Simplification of a irregular object for engineering approximations for engineering applications

^{*}Graduate Aerospace Engineering, AIAA Student Member, Aerospace Honors Society.



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1 Introduction

1.1 Overview

A flow that is incompressible and irrotational can be modeled by Laplace's equation for a scalar velocity potential. In two dimensions, an equivalent formulation is obtained using a stream function, $\psi(x, y)$, from which the velocity components are given by

$$u = \frac{\partial \psi}{\partial y}, \qquad v = -\frac{\partial \psi}{\partial x} \tag{1}$$

With this definition, continuity is automatically satisfied. Requiring the flow to be irrotational leads to Laplace's equation for ψ ,

$$\nabla^2 \psi = 0 \tag{2}$$

In this project, I will be solving for two-dimensional potential flow around a square, as illustrated in Figure 1.

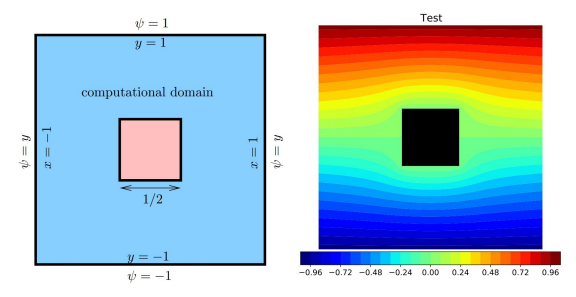


Figure 1: Solution of potential flow around a square, using a stream function, $\psi(x,y)$.

The computational domain is a square, $(x,y) \in [-1,1]^2$, and the boundary conditions are Dirichlet. On the bottom and top walls, ψ is set to -1 and +1, respectively, and on the sides, ψ is set to y. The inner square, of side length $\frac{1}{2}$ and placed in the center of the outer square, presents an obstacle to the flow, and by symmetry it corresponds to a streamline on which $\psi=0$.

1.2 Discretization

I will use a finite-difference method to solve Equation 2 on the computational domain. The grid consists of a lattice of $(N+1)^2$ points, as shown in Figure 2. Some of these points are on the boundary, and some are inside the inner square, hence outside the computational domain. It is up to me to decide how best to deal with these points. N is the number of intervals across the entire domain, so that the spacing is $\Delta x = \Delta y = h = 2/N$.

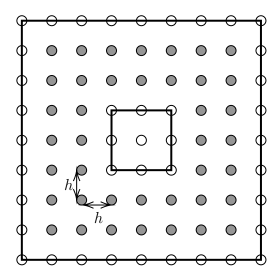


Figure 2: Finite difference grid for $p = 0 \rightarrow N = 8$.

To ensure that the grid conforms to the boundary of the inner square, use

$$N = 2^{p+3}, p = [0, 1, 2, \ldots]$$
 (3)

At each interior node, I will use a standard second-order five-point stencil to discretize the Laplacian in Equation 2. Note that the given Dirichlet conditions fully specify ψ on the boundaries.

1.3 Solvers

There are three types of solvers that I will implement to study the flow around the square.

1.3.1 Direct

The direct solver will build a *sparse* linear system of equations to solve for the nodal states. The system will take the form

$$\underline{\underline{A}\Psi} = \underline{F} \tag{4}$$

where $\underline{\Psi}$ is the unrolled state vector of unknowns. Solve this system using a sparse direct solver, such as the backslash operator in Matlab or scipy.sparse.linalg.spsolve().

1.3.2 Iterative Smoothers

For the other solver I will implement two iterative smoothers: under-relaxed Jacobi and overrelaxed Gauss-Seidel. For Gauss-Seidel, I will use the "red-black" ordering, in which the nodes are colored in checkerboard fashion and the smoother is applied first to the red nodes and then to the black nodes. The presence of the inner square does not change the red-black ordering: imagine a checkerboard with the center cut out. I will use under-relaxation for Jacobi and over-relaxation for Gauss-Seidel.

1.3.3 Multigrid

The last solver I will use will implement a V-cycle in which successively finer grids are obtained by increasing p in Equation 4. Use full-weighting for the restriction operator, I_{2h}^h , and interpolation for the prolongation operator, I_h^{2h} . On the down/up sweep of the V-cycle perform $\nu_1 = \nu_2 = 2$ pre/post smoothing iterations. On the coarsest grid, perform $\nu_c = 50$ smoothing iterations. For the initial conditions in all iterative runs, use $\psi = 0$ at the interior nodes. For the smoother, use Gauss-Seidel with an over-relaxation factor of $\omega = 1.5$. Again, implement these without any matrices whose size scales with the number of unknowns

1.4 Post-Processing

To visualize the flowfield, plot contours of the stream function, as shown in Figure 1. These contours are streamlines of the flow. The top and bottom of the domain represent impenetrable walls, since ψ is set to a constant there. Acceleration of the fluid around the square changes the pressure of the fluid, and this affects the force on the walls. Of interest will be the pressure coefficient distribution on the bottom wall,

$$c_p(x) = 1 - \frac{u^2}{U_{\infty}^2},\tag{5}$$

where the free-stream speed (speed of the flow without the inner square present) is $U_{\infty} = 1$ based on the boundary conditions. Note that u is obtained by differentiating ψ , according to Equation 1. Use a second-order one-sided finite difference of the ψ data to obtain u at the bottom boundary nodes.

Integrating and normalizing the pressure coefficient gives the lift coefficient on the bottom wall,

$$c_l(x) = \frac{1}{2} \int_0^2 c_p(x) \ dx \tag{6}$$

In this approximation I will use the trapezoidal method to perform this integration.

2 Tasks and Deliverables

In this project I will implement a finite-element solver to approximate the flow field around a square and verify its coefficients of lift and pressure. Additionally I will implement Jacobi and Gauss-Seidel smoothers and determine their convergences to theory. Lastly, I will implement a V-cycle multigrid method and will determine the best level so that the system converges to the appropriate solution.

2.1 Solving the Finite-Difference System

Since the boundary conditions have been specified, I will use a 5-point stencil about the interior nodes to approximate the global $\underline{\underline{A}}$ sparse matrix that will approximate the nodal values. After assembling the $\underline{\underline{A}}$ matrix I will reduce it to solve for the interior $\underline{\psi}$ nodal values. Where the 5-point stencil can be written to be,

$$\frac{1}{h^2} \begin{bmatrix} +u_{i,j+1} \\ +u_{i-1,j} - 4u_{i,j} + u_{i+1,j} \\ u_{i,j-1} \end{bmatrix} = f_{i,j}$$
 (7)

Using the 5-point stencil shown above in Equation 7, and the coefficients to form the global stiffness matrix $\underline{\underline{A}}$ and using the Dirichlet boundary conditions to form \underline{F} through Python I can solve directly for the approximated solution for ψ . After implementing the finite difference system shown in Algorithm 1 and using the direct solver the 9×9 matrix for p = 0, the re-shaped Ψ matrix becomes,

$\underline{\psi} =$	1.0000 0.7500 0.5000 0.2500 0.0000 -0.2500 -0.5000	1.0000 0.7336 0.4709 0.2208 -0.0000 -0.2208 -0.4709	1.0000 0.7137 0.4291 0.1625 -0.0000 -0.1625 -0.4291	1.0000 0.6920 0.3692 0.0000 0.0000 -0.3692	1.0000 0.6849 0.3558 0.0000 0.0000 -0.3558	1.0000 0.6920 0.3692 0.0000 0.0000 -0.3692	1.0000 0.7137 0.4291 0.1625 0.0000 -0.1625 -0.4291	1.0000 0.7336 0.4709 0.2208 0.0000 -0.2208 -0.4709	1.0000 0.7500 0.5000 0.2500 0.0000 -0.2500 -0.5000	
	$ \begin{bmatrix} -0.5000 \\ -0.7500 \\ -1.0000 \end{bmatrix} $	-0.4709 -0.7336 -1.0000	-0.4291 -0.7137 -1.0000	-0.3692 -0.6920 -1.0000	-0.3558 -0.6849 -1.0000	-0.3692 -0.6920 -1.0000	-0.4291 -0.7137 -1.0000	-0.4709 -0.7336 -1.0000	-0.5000 -0.7500 -1.0000	

2.2 Post-Processing Functions

Firstly, to approximate the coefficient of pressure about the bottom of the wall I will use a second-order one-sided finite difference to approximate u given through,

$$u = \frac{\partial \psi}{\partial y} \approx \frac{-\frac{3}{2}\psi_N + 2\psi_{N-1} - \frac{1}{2}\psi_{N-2}}{h^2} \tag{8}$$

Using Equation 8 above I can approximate the horizontal velocity to then approximate the coefficient of pressure using Equation 5 for $p \in [0, 2, 4]$. Doing this approximation with the aid of Algorithm 2 results in Figure 3 shown below.

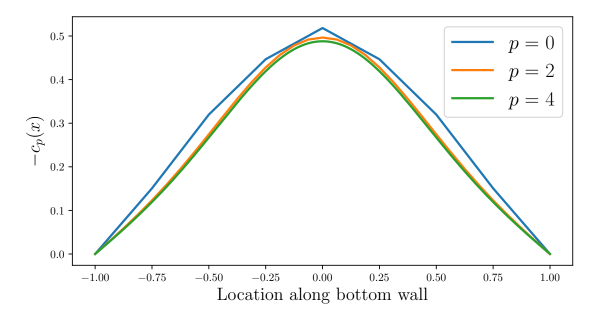


Figure 3: P-scaling coefficient of pressure $-c_p(x)$ along bottom wall.

Looking above to Figure 3, increasing p will simply "smooth" out the coefficient of pressure as it varies along x. Notable is that $-c_p(x)$ increases as it gets to the middle, where the middle of the square lies about the x-axis. This increase in $-c_p(x)$ indicates that the flow is increasing here as the coefficient of pressure indicates how the local flow is changing with respect to the ambient flow U_{∞} .

Nextly, after having approximated u, is to solve for the coefficient of lift c_l using Equation 6. However, to approximate this integral I will use the trapezoidal integration method shown below in Equation 9.

$$\int_{a}^{b} f(x) dx \approx \frac{\Delta x}{2} (f(x_N) + f(x_0)) + \Delta x \sum_{i=0}^{N} f(x_i)$$
 (9)

Using Equation 9 to numerically approximate the coefficient of lift through Equation 6, I can approximate the coefficient of lift per specific p value. Through this method I will run a simulation with p=7 to approximate an "exact" solution. Using this method I get the following results in Table 1 below.

Table 1: Coefficient of lift c_l values at p.

<i>p</i>	c_l
p=0	-0.2940765
p = 1	-0.2795586
p=2	-0.2706653
p = 3	-0.2665109
p=4	-0.2647269
p = 5	-0.2639878

Conducting this simulation again with p = 7 approximates the "exact" solution to be,

$$c_{l,exact}(p=7) = -0.2635660$$

Using this value to be the exact solution, I can then approximate the convergence rate of the simulations by taking the absolute value of the difference and comparing it to the p value and obtaining the rate of convergence as $r = \frac{\log_{10} \tau_{i+1}/\tau_i}{\log_{10} \Delta h_{i+1}/\Delta h_i}$. Conducting this convergence study gives that the approximated error is,

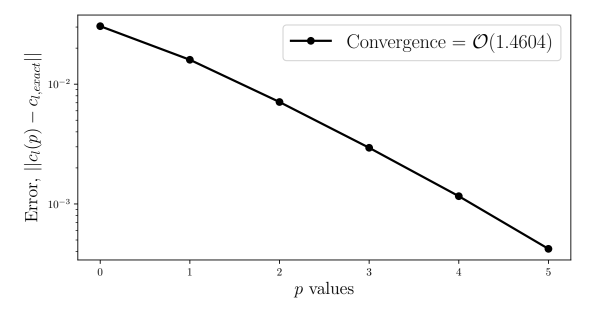


Figure 4: Approximated error and convergence while varying p.

Looking above to Figure 4 the slope of the convergence is $\mathcal{O}(\Delta h^2)$ hence second-order accuracy. This order of accuracy is consistent with the second-oder one-sided finite difference and the 5-point stencil and is the expected slope. This graph confirms the order of accuracy.

2.3 Implementing Jacobi and Gauss-Seidel Smoothers

In this task, I will implement both a Jacobi iterative smoother and a Gauss-Seidel smoother. Taking an iterative solution approach allows for easier implementation, but as a result will display a slower convergence. This convergence history will be highlighted here.

2.3.1 Jacobi Iteration Smoother

In order to implement the Jacobi Iteration Smoother I will use the expression for the next iteration state approximation shown below as,

$$u_{i,j}^{n+1} = \frac{1}{4} \left(u_{i-1,j}^n + u_{i+1,j}^n + u_{i,j-1}^n + u_{i,j-1}^n + \Delta h^2 f_i \right)$$
(10)

Using Equation 10 above, and implementing into Python code shown in Appendix 3, I can use this to approximate the next iteration state values and iterate several times until the approximation matches that of the analytical. Choosing the iteration values I choose to pick a high value of iterations until the approximated solution matched that of Figure 1. Then from here I conducted an L_2 residual norm and its convergence shown below.

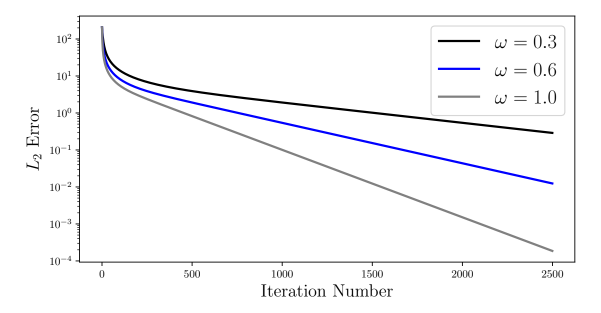


Figure 5: L_2 residual norm convergence history for varying ω .

As shown above in Figure 5, the over-relaxation factor plays a large role into how fast the solution converges. Further analysis on the y-log plot shows that it appears as if the multiple of the over-relaxation factor of ω denotes what its L_2 error will be as $\omega = 0.6$ has an L_2 residual norm that is approximately twice the magnitude that of $\omega = 0.3$ residual norm.

2.3.2 Gauss-Seidel Smoother

The Gauss-Seidel smoother is very similar in implementation to the Jacobi iteration smoother, however it differs in how the state updates. Gauss-Seidel will update half of the state first and then use the updated state to further smooth the approximated states in that given iteration. This method is called the "checker-board" or black-red update. This essentially means that every other node/adjacent node will get updated and then use these updated nodes to make a better approximation. Implementing this in Python can be shown in Appendix 4, but with the L_2 residual norm error shown below for differing values of over-relaxation.

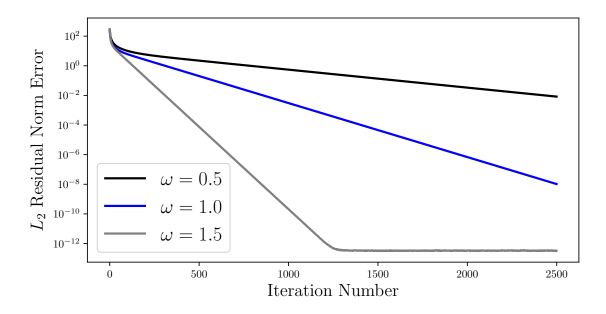


Figure 6: L_2 residual norm convergence history for varying ω .

Shown above in Figure 6, is the L_2 residual norm error through its iteration values. This shows again the the over-relaxation value does converge faster with a higher value. However, this may be an artifact of machine precision but it does appear that $\omega = 1.5$ will converge to a given solution whereas $\omega = 0.5$ or $\omega = 1.0$ may be able to converge to a more precise solution but would take longer to do so.

Across both the Jacobi iteration smoother and the Gauss-Seidel smoother, the rates at which they converge depend on the over-relaxation value. Having a higher over-relaxation value will allow the iterative solver to coverge faster. However, notable is that the Gauss-Seidel smoother can have a over-relaxation value greater than 1 or $\omega > 1$ and still converge. When Jacobi iteration smoother has an over-relaxation value greater than 1 it will diverge from the analytical solution whereas Gauss-Seidel continues to converge. Running Jacobi iteration with $\omega = 1.1$ gave that the final value was $\mathcal{O}(10^{140})$ which is in agreement that an over-relaxation value greater than 1 will diverge.

2.4 Implementing V-Cycle Multigrid Method

The method of the V-Cycle multigrid is computing the residual at a fine mesh, restricting it to a smaller grid then repeating until reaching the coarsest mesh grid size. At this coarsest meshgrid the restricted residual will be smoothed and then the prolongation scheme will start. In this prolongation the error is passed up through each prolongation and smoother and ultimately added to the initial guess of the state matrix.

2.4.1 V-Cycle Multigrid Method Convergence

Restriction Restricting the multigrid can be done by simply iterating by multiples of 2 throughout the x and y directions and taking a weighted average of the surrounding nodes such that the restriction can be written below in Equation 11,

$$r_{i,j} = \frac{1}{4} \underbrace{r_{i,j}}_{\text{Center node}} + \frac{1}{8} \underbrace{(r_{i\pm 1,j} + r_{i,j\pm 1})}_{\text{Up/Down nodes}} + \frac{1}{16} \underbrace{(r_{i\pm 1,j\pm 1} + r_{i\pm 1,j\mp 1})}_{\text{Corner nodes}}$$
(11)

Prolongation Prolongation is similar to restriction except that it is acting in the opposite direction. Instead of weighting adjacent nodes to a center node it applies one node to several. In my implementation I would iterate fully through the finer mesh but I would conduct integer division to use values from every other node and then weight them accordingly. The expression that I used for prolongation can be shown below in Equation 12

$$\begin{bmatrix} e_{i-1_h,j+1_h} & e_{i_h,j+1_h} & e_{i+1_h,j+1_h} \\ e_{i-1_h,j_h} & e_{i_h,j_h} & e_{i+1_h,j_h} \\ e_{i-1_h,j-1_h} & e_{i_h,j-1_h} & e_{i+1_h,j+1_h} \end{bmatrix} = \begin{bmatrix} \frac{1}{4}e_{i_{2h},j_{2h}} & \frac{1}{2}e_{i_{2h},j_{2h}} & \frac{1}{4}e_{i_{2h},j_{2h}} \\ \frac{1}{2}e_{i_{2h},j_{2h}} & e_{i_{2h},j_{2h}} & \frac{1}{2}e_{i_{2h},j_{2h}} \\ \frac{1}{4}e_{i_{2h},j_{2h}} & \frac{1}{2}e_{i_{2h},j_{2h}} & \frac{1}{4}e_{i_{2h},j_{2h}} \end{bmatrix}$$
(12)

Implementing Multigrid Using Equations 11, 12 and implementing into the Python environment with Algorithm 5, I ultimately arrive to find the convergence rates for the V-Cycle multigrid. Running several iterations with differing p values I get that the L_2 residual norms with $\nu_1 = \nu_2 = 2$ and $\nu_c = 50$ are shown in Figure 7 on the following page.

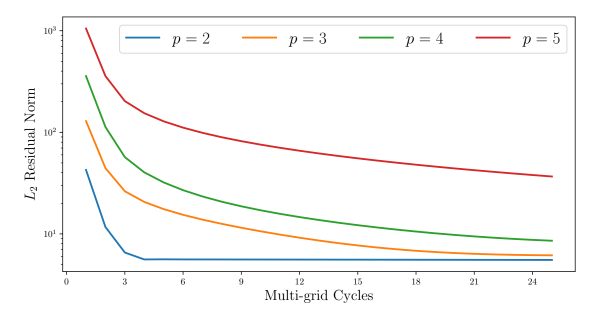


Figure 7: The V-Cycle multigrid method L_2 convergence while varying p.

Shown above in Figure 7, is the convergence for the V-cycle multigrid L_2 residual norm. Looking to the convergence rates, as you increase the size of the meshgrid (increase of p) the longer it will take to converge to the solution. This can be shown as the horizontal asymptote reached first from p=2. Noting that the L_2 residual norm is relatively large this is most likely due to the fact that the smoothing values ν_1, ν_2, ν_c are all relatively small compared to the actual smoothing iterations that I used for Jacobi and Gauss-Seidel with $\mathcal{O}(2500)$ iterations. These small smoothing values result in a larger residual but a converged solution nonetheless.

2.4.2 V-Cycle Multigrid Smoothing Iterations

In order to choose an efficient V-cycle setting to create accurate solutions that require lower computing power relative to other solutions will require "tweaking" of the smoothing iterations until values are found that result in accurate solutions that require less computing than other alternatives. To relate V-Cycle multigrid to Gauss-Seidel I will be using Work Units to compare the two. The units for "work units" can be defined below in Equation 13,

1 V Cycle =
$$\sum_{l=0}^{n_{\text{level}}-1} (\nu_1 + \nu_2) 2^{-l}$$
 Work Units (13)

After some trial and error and adjusting ν_1, ν_2, ν_c , and the number of iterations plotting V-Cycle multigrid and Gauss-Seidel against their work units arrives to Figure 8 shown below.

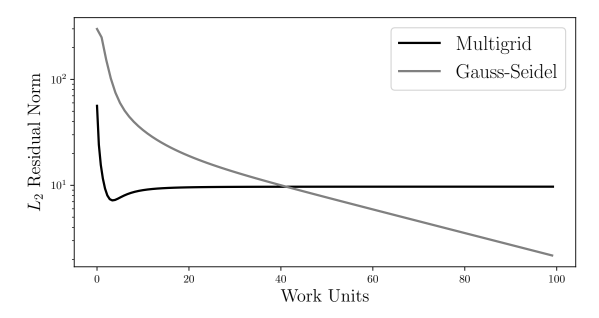


Figure 8: Adjusting V-Cycle multigrid parameters to compare computational costs between V-Cycle multigrid and Gauss-Seidel smoother.

Looking above to Figure 8, I found that the V-cycle was ultimately more accurate for lower work units but at higher work units Gauss-Seidel would become more accurate as V-cycle would start to bring diminishing returns at higher work units. I believe that this is due to the fact that this V-cycle multigrid would converge quickly to its solvers solution whereas the Gauss-Seidel methods solution is more accurate but for better accuracy costs significant more computational power(multigrid converged to its solution at ≈ 10 work units whereas it took Gauss-Seidel ≈ 40 work units). For quick convergence these are the V-Cycle multigrid smoothing parameters that I recommend.

$$\mathbf{k} = \mathbf{50}, \quad
u_1 = \mathbf{10}, \quad
u_2 = \mathbf{10}, \quad
u_c = \mathbf{1000}$$

Appendices

A Python Direct Solver Implementation

Algorithm 1: Driving Code to Implement Direct Solver.

```
def directsol(p):
       import numpy as np
3
       from scipy import sparse
4
       from scipy.sparse import linalg
       N = 2**(p + 3) \# P-value scaling
h = float(2/N) \# Step size
6
       xlin = np.linspace(-1, 1, N+1, endpoint=True) # linspace over domain
9
       xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
10
11
       A = \text{sparse.lil\_matrix}(((N+1)**2,(N+1)**2)) \# \text{Pre-allocate sparse Matrix}
12
       q = np.zeros((N+1)**2)
                                                     # Pre-allocate q vector
13
       for iy in range(N+1):
14
           for ix in range(N+1):
               15
16
17
18
19
               if ylin[iy,ix] == 1: # Top Boundary
20
                   q[i] = 1
A[i,i] = 1
21
22
               elif ylin[iy,ix] == -1: # Bottom Boundary
23
                   q[i] = -1
                   A[i,i] = 1
24
25
               elif abs(xlin[iy,ix]) == 1: # Left/Right Boundary
26
                   q[i] = ylin[iy,ix]
A[i,i] = 1
27
28
               elif abs(xlin[iy,ix]) <= 0.25 and abs(ylin[iy,ix]) <= 0.25: # Interior</pre>
                   Boundary
                   q[i] = 0
29
                  \bar{A}[i,i] = 1
30
               else: # Interior Domain
31
32
                   q[i] = 0
                   \bar{A}[i,i] = -4*h**-2
33
34
                   A[i,iL] = h**-2
                   A[i,iR] = h**-2
35
36
                   A[i,iD] = h**-2
37
                   A[i,iU] = h**-2
38
       phiv = linalg.spsolve(sparse.csr_matrix(A), q) # Solve for Phi
       phi = np.reshape(phiv, (N+1, N+1))
39
                                             # Re-shape for plotting
40
41
       return xlin, ylin, phi
```

B Coefficient of Pressure Implementation

Algorithm 2: Python Implemention to Approximate Coefficient of Pressure.

```
def calc_cp(ylin, phi):
 1
2
3
         import numpy as np
 4
         h = ylin[0,0] - ylin[1,0]
         num = max(np.shape(ylin)) - 1
cp = np.zeros(num + 1)
 5
 \frac{6}{7}
         for i in range(num+1):
    u = (-3/2*phi[num,i] + 2*phi[num-1,i] - 1/2*phi[num-2,i])*h**-1
 8
 9
10
             cp[i] = 1 - u**2
11
12
         return cp
13
     def calc_cl(ylin, cp):
14
15
         import numpy as np
16
         h = ylin[0,0] - ylin[1,0]
17
18
         num = max(np.shape(ylin)) - 1
         cl = 0
19
20
21
22
         for i in range(1, num):
         cl += h*cp[i]
cl += h/2*(cp[0] + cp[num])
23
24
25
         cl *= 1/2
26
         return cl
```

C Jacobi Iteration Smoother

Algorithm 3: Python Implemention of Jacboi Iteration Smoother.

```
def jacobisol(omega):
         import numpy as np
 3
         import math
 4
         p = 3
N = 2**(p + 3) # P-value scaling
h = float(2/N) # Step size
 5
 6
         xlin = np.linspace(-1, 1, N+1, endpoint=True) # linspace over domain
xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
 8
 9
10
11
         # Variables for Jacobi Iteration
12
         resid_mat = np.zeros((N+1, N+1))
         num_iters = 2500
13
14
         resid = np.zeros(num_iters)
15
16
         # Boundary Conditions / Intial Guess
         U = np.zeros((N+1, N+1))

U[0,:] = 1; U[N,:] = -1

U[:,0] = ylin[:,0]; U[:,N] = ylin[:,0]
17
18
19
         u_temp = U.copy()
20
\frac{1}{21}
         for k in range(num_iters):
23
              for j in range(1, N):
24
                  for i in range(1, N):
    if abs(xlin[j,i]) <= 0.25 and abs(ylin[j,i]) <= 0.25: # Interior</pre>
25
                            Boundary
                           u_{temp}[j,i] = 0
                       resid_mat[j,i] = 0
else: # Interior Domain
27
28
29
                           u_{temp}[j,i] = (U[j+1, i] + U[j-1, i] + U[j,i+1] + U[j,i-1])/4
30
                           rij = 0 - h**-2*(4*u_temp[j,i] - 4*U[j,i])
31
                           resid_mat[j,i] = rij
32
33
              U[1:N, 1:N] = U[1:N, 1:N]*(1.-omega) + u_temp[1:N, 1:N]*omega
34
              for i in range(N+1):
35
                  for j in range(N+1):
36
                       resid[k] += resid_mat[j,i]**2
              resid[k] = math.sqrt(resid[k]/(N+1)**2)
37
38
39
         return resid
```

D Gauss-Seidel Iteration Smoother

Algorithm 4: Python Implemention of Gauss-Seidel Iteration Smoother.

```
def gausssol(num_iters, p, omega):
        import numpy as np
 3
        import math
 4
        # Variables for Jacobi Iteration
 5
       N = 2**(p + 3) \# P-value scaling
       h = \frac{1}{1000} # Step size
 6
 7
       xlin = np.linspace(-1, 1, N+1, endpoint=True) # linspace over domain
 8
       xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
9
       resid_mat = np.zeros((N+1, N+1))
10
       resid = np.zeros(num_iters)
11
12
        # Boundary Conditions / Intial Guess
13
       U = np.zeros((N+1, N+1))
       U[0,:] = 1; U[N,:] = -1; U[:,0] = ylin[:,0]; U[:,N] = ylin[:,0]
14
15
        for k in range(num_iters):
           for j in range(1, N): # Red Nodes
16
17
               if j\%2 == 0:
18
                   for i in range(1, N, 2):
                       if abs(xlin[j,i]) <= 0.25 and abs(ylin[j,i]) <= 0.25: # Interior</pre>
19
                           Boundary
20
                          resid_mat[j,i] = 0
                       else: # Interior Domain
21
22
                          u_{u} = (U[j+1, i] + U[j-1, i] + U[j,i+1] + U[j,i-1])/4
                          rij = 0 - h**-2*(4*u_unew - 4*U[j,i])
resid_mat[j,i] = rij
23
24
25
                          U[j,i] = U[j,i]*(1.-omega) + u_unew*omega
26
               else:
27
                   for i in range(2, N-1, 2):
28
                       if abs(xlin[j,i]) \le 0.25 and abs(ylin[j,i]) \le 0.25: # Interior
                           Boundary
29
                          resid_mat[j,i] = 0
30
                       else: # Interior Domain
                          u_u = (U[j+1, i] + U[j-1, i] + U[j,i+1] + U[j,i-1])/4
31
32
                          rij = 0 - h**-2*(4*u\_unew - 4*U[j,i])
                          resid_mat[j,i] = rij
33
                          U[j,i] = U[j,i]*(1.-omega) + u_unew*omega
34
35
           for j in range(1, N): # Black Nodes
36
               if j\%2 == 0:
37
                   for i in range(2, N-1, 2):
                       if abs(xlin[j,i]) <= 0.25 and abs(ylin[j,i]) <= 0.25: # Interior</pre>
38
                           Boundary
39
                          resid_mat[j,i] = 0
40
                       else: # Interior Domain
                          u_{u} = (U[j+1, i] + U[j-1, i] + U[j,i+1] + U[j,i-1])/4
41
                          rij = 0 - h**-2*(4*u_unew - 4*U[j,i])
42
43
                          resid_mat[j,i] = rij
                          U[j,i] = U[j,i]*(1.-omega) + u_unew*omega
44
45
               else:
46
                   for i in range(1, N, 2):
                       if abs(xlin[j,i]) \le 0.25 and abs(ylin[j,i]) \le 0.25: # Interior
47
                           Boundary
48
                          resid_mat[j,i] = 0
49
                       else: # Interior Domain
                          u_u = (U[j+1, i] + U[j-1, i] + U[j,i+1] + U[j,i-1])/4
50
                          rij = 0 - h**-2*(4*u\_unew - 4*U[j,i])
51
52
                          resid_mat[j,i] = rij
53
                          U[j,i] = U[j,i]*(1.-omega) + u_unew*omega
           for i in range(N+1):
54
55
               for j in range(N+1):
56
                   resid[k] += resid_mat[j,i]**2
           resid[k] = math.sqrt(resid[k]/(N+1)**2)
57
58
        return resid
```

E V-Cycle Multigrid

Algorithm 5: Python Implemention of V-Cycle Multigrid.

```
import numpy as np
  2
            import math
  3
            from direct_sol import directsol
            def residual(U, F):
  5
                       N = U.shape[0] - 1; h = 2.0/N
                       R = U.copy()
  7
  8
  9
                       for iy in range(1, N):
10
                                  for ix in range(1, N):
                                             R[iy, ix] = F[iy, ix] - (U[iy+1, ix] + U[iy-1, ix] + U[iy, ix+1] + U[i
11
                                                           ix-1] - 4*U[iy, ix])*h**-2
12
                       return R
13
            def restrict(r, N):
                       rc = np.zeros([int(N/2)+1, int(N/2)+1])
14
15
16
                       ix = 0; iy = 0
                       for j in range(1, N+1, 2):
17
                                   for i in range(1, N+1, 2):
18
                                             rc[iy,ix] = 1/8.*(r[j+1,i] + r[j-1,i] + r[j,i+1] + r[j,i-1]) + 1/16.*(r[j+1,i] + r[j,i+1]) + 1/16.*(r[j+1,i] + r[j+1,i] + r[j+1,i]) + 1/16.*(r[j+1,i] + r[j+1,i] + r[j+1,i]) + 1/16.*(r[j+1,i] + r[j+1,i] + r[j+1,i]) + 1/16.*(r[j+1,i] + r[j+1,i] + r
19
                                                          +1,i+1 + r[j-1,i-1] + r[j-1,i+1] + r[j+1,i-1]) + 1/4.*r[j,i]
20
                                              ix += 1
                                              if ix >= int(N/2):
21
                                                        ix = 0
22
23
                                  iy += 1
24
                       return rc
25
            def prolongate(e2h, N):
26
                       eh = np.zeros([int(2*N)+1, int(2*N)+1])
27
                       xlin = np.linspace(-1, 1, int(2*N)+1, endpoint=True) # linspace over domain
28
                       xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
29
                       for j in range(1, int(2*N), 2):
30
                                  for i in range(1, int(2*N), 2):
31
                                                         if abs(xlin[j,i]) <= 0.25 and abs(ylin[j,i]) <= 0.25: # Interior</pre>
32
                                                                     Boundary
33
                                                                   pass
34
                                                        else: # Interior Domain
35
                                                                    # Self Weight
36
                                                                    eh[j, i] = e2h[math.floor(j/2.), math.floor(i/2.)]
37
38
39
                                                                   eh[j+1, i] += 0.5*e2h[math.floor(j/2.), math.floor(i/2.)]
                                                                    eh[j-1, i] += 0.5*e2h[math.floor(j/2.), math.floor(i/2.)]
40
                                                                    eh[j, i+1] += 0.5*e2h[math.floor(j/2.), math.floor(i/2.)]
41
42
                                                                   eh[j, i-1] += 0.5*e2h[math.floor(j/2.), math.floor(i/2.)]
43
44
                                                                   # Corner Nodes
                                                                   eh[j+1, i+1] += 0.25*e2h[math.floor(j/2.), math.floor(i/2.)]
45
                                                                    eh[j-1, i-1] += 0.25*e2h[math.floor(j/2.), math.floor(i/2.)]
46
47
                                                                    eh[j-1, i+1] += 0.25*e2h[math.floor(j/2.), math.floor(i/2.)]
48
                                                                    eh[j+1, i-1] += 0.25*e2h[math.floor(j/2.), math.floor(i/2.)]
49
                       return eh
50
            def multigrid(U, F, p, pmax, viter, nu1, nu2, nuc):
51
52
                       12err = np.zeros(viter)
53
54
55
                        print("V-Cycle_Method(p=", pmax, ")\n----")
56
                       for k in range(viter):
57
                                 print("Iteration:□", k)
58
59
                                  N = U.shape[0] - 1
60
                                  griditer = 0
61
                                  fmat = np.zeros([U.shape[0], U.shape[0], pmax-p+1])
62
63
                                  # Sweep Down
```

```
64
             utemp = smooth(U, fh, nu1)
             while N > 2**(p + 3):
 65
 66
                 rh = residual(utemp, fh)
 67
                 f2h = restrict(rh, N)
 68
                 griditer += 1
 69
                 \tilde{N} = int(N/2)
 70
 71
                 fmat[0:(N+1), 0:(N+1), griditer] = f2h
 72
                 utemp = smooth(np.zeros([N+1, N+1]), f2h, nu1)
 73
 74
 75
76
             utemp = smooth(np.zeros([N+1, N+1]), fmat[0:(N+1), 0:(N+1), griditer], nuc)
 77
             # Sweep Up
 78
             while N \le 2**(pmax + 2):
 79
                 utemp = prolongate(utemp, N)
 80
 81
                 N = int(2*N)
 82
                 griditer -= 1
 83
                 utemp = smooth(utemp, fmat[0:(N+1), 0:(N+1), griditer], nu2)
 84
 85
             U += utemp
 86
             resid = residual(U, F)
 87
 88
             for j in range(N + 1):
 89
                 for i in range(N + 1):
 90
                     12err[k] += resid[j,i]**2
 91
             12err[k] = np.sqrt(12err[k]/(N + 1)**2)
 92
 93
         return U, 12err
 94
     def vcyclesol(p, pmax, viter, nu1, nu2, nuc):
 95
         N = 2**(pmax + 3)
 96
         U = np.zeros([N+1, N+1])
         F = \overline{U.copy}()
 97
         U[:, 0] = np.flip(np.linspace(-1, 1, N+1))
U[:, N] = np.flip(np.linspace(-1, 1, N+1))
 98
 99
100
         U[0,:] = 1; U[N,:] = -1
101
102
         U, 12err = multigrid(U, F, p, pmax, viter, nu1, nu2, nuc)
103
104
         return U, 12err
     def smooth(U, F, nu):
105
106
         N = U.shape[0]-1; h = 2.0/N
107
         omega = 1.5
108
109
         xlin = np.linspace(-1, 1, N+1, endpoint=True) # linspace over domain
110
         xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
111
         for k in range(nu):
112
             for iy in range(1, N): # Red Nodes
113
                 if iy%2 == 0:
                     for ix in range(1, N, 2):
114
                         if abs(xlin[iy,ix]) <= 0.25 and abs(ylin[iy,ix]) <= 0.25: #</pre>
115
                              Interior Boundary
116
                             U[iy, ix] = 0
                         else: # Interior Domain
117
                             unew = 0.25*(U[iy+1, ix] + U[iy-1, ix] + U[iy, ix-1] + U[iy, ix+1] - F[iy, ix]*h**2)
118
119
                             U[iy, ix] = U[iy, ix]*(1.0 - omega) + unew*omega
120
                 else:
121
                     for ix in range(2, N-1, 2):
                         if abs(xlin[iy,ix]) <= 0.25 and abs(ylin[iy,ix]) <= 0.25: #
122
                              Interior Boundary
                             U[iy, ix] = 0
123
                         else: # Interior Domain
124
125
                             unew = 0.25*(U[iy+1, ix] + U[iy-1, ix] + U[iy, ix-1] + U[iy,
                             ix+1] - F[iy, ix]*h**2)
U[iy, ix] = U[iy, ix]*(1.0 - omega) + unew*omega
126
             for iy in range(1, N): # Black Nodes
127
128
                 if iy%2 == 0:
```

```
for ix in range(2, N-1, 2):
    if abs(xlin[iy,ix]) <= 0.25 and abs(ylin[iy,ix]) <= 0.25: #</pre>
129
130
                               Interior Boundary
                          U[iy, ix] = 0
else: # Interior Domain
131
132
                              unew = 0.25*(U[iy+1, ix] + U[iy-1, ix] + U[iy, ix-1] + U[iy,
133
                              ix+1] - F[iy, ix]*h**2)
U[iy, ix] = U[iy, ix]*(1.0 - omega) + unew*omega
134
135
                  else:
                      for ix in range(1, N, 2):
    if abs(xlin[iy,ix]) <= 0.25 and abs(ylin[iy,ix]) <= 0.25: #</pre>
136
137
                              Interior Boundary
                          U[iy, ix] = 0
else: # Interior Domain
138
139
                              140
141
142
         return U
```

F Main Python Driving Code

Algorithm 6: Python Implemention of V-Cycle Multigrid.

```
from matplotlib import pyplot as plt
    from matplotlib.ticker import MaxNLocator
   import numpy as np
    import math
   from direct_sol import directsol
    from calc_coefs import calc_cp, calc_cl
    from jacobi_sol import jacobisol
    from gauss_sol import gausssol
    from vcycle_sol import vcyclesol
10
    plt.rc('text', usetex=True)
plt.rc('font', family='serif')
11
12
13
14
    def export_phi(phi):
        f = open('9by9_mat', "w") # Filename
output = ''
15
16
17
        for j in range(9):
18
             for i in range(9):
                 output += str.format('\{0:.4f\}',phi[j,i]) + r'\&_{\perp}'
19
20
             if i == 8:
21
                 output += r'\\' # Output results to LaTeX environment
22
        f.write(output)
23
        f.close()
24
25
    def export_cl(cl):
26
        f = open('cl_vals', "w") # Filename
27
        output = ''
28
        for i in range(6):
            output += r'$p_=,$\(\frac{1}{2}\)' + str.format('\{0:.0f\}',i)+ r'\&_{\Box}' + str.format('\{0:.7f\}')
29
                  ,cl[i]) + r'\\'
30
        f.write(output)
31
        f.close()
32
    def gen_grids(p):
    N = 2**(p + 3) # P-value scaling
33
34
        xlin = np.linspace(-1, 1, N+1, endpoint=True) # linspace over domain
xlin, ylin = np.meshgrid(xlin, np.flip(xlin)) # Meshgrid values
35
36
37
38
        return xlin, ylin
39
40
    def run_q1():
        xlin, ylin, phi = directsol(0)
41
42
        export_phi(phi)
43
44
    def run_q2():
        plt.figure(figsize=(8,4))
45
46
        for i in np.array([0, 2, 4]):
47
            xlin, ylin, phi = directsol(i)
48
            cp = calc_cp(ylin, phi)
49
50
            plot_label = r'$p$_{\square}=_{\square}' + str(i)
            plt.plot(xlin[0,:], -cp, lw = 2, label = plot_label)
51
        plt.xlabel(r'Location_along_bottom_wall', fontsize = 16)
52
53
        plt.ylabel(r'$-c_p(x)$', fontsize = 16)
        plt.legend(fontsize = 18)
54
55
        plt.savefig('figs/cp_runs.pdf', bbox_inches = 'tight')
56
        plt.show()
57
58
        xlin, ylin, phi = directsol(7)
        cl_exact = calc_cl(ylin, calc_cp(ylin, phi))
f = open('cl_exact', "w")
59
60
        output = str.format('{0:.7f}',cl_exact)
61
62
        f.write(output)
63
        f.close()
64
        cl_vals = np.zeros(6); h_vals = np.zeros(6); cl_err = np.zeros(6)
```

```
66
                      for i in range(6):
                               xlin, ylin, phi = directsol(i)
  67
                               cl = calc_cl(ylin, calc_cp(ylin, phi))
  68
  69
                               cl_vals[i] = cl
  70
                              h_vals[i] = ylin[0,0] - ylin[1,0]
                               cl_err[i] = abs(cl - cl_exact)
   71
   72
                     export_cl(cl_vals)
  73
   74
                     rate = math.log10(cl_err[4]/cl_err[5])/math.log10(h_vals[4]/h_vals[5])
   75
                     plot_label = r^{\bar{}}Convergence_{\sqcup} = \frac{1}{2} \cdot \frac{1}{2
   76
                     plt.figure(figsize=(8,4))
  77
                     plt.plot(range(6), cl_err, color = 'black', marker = 'o', lw = 2, label =
                                plot_label)
  78
                     plt.xlabel(r'$p$_values', fontsize = 16)
                     plt.ylabel(r'Error,_{\square}$||_{\square}c_1(p)_{\square}-_{\square}c_{1,_{\square}exact}||$', fontsize = 16)
  79
  80
                     plt.yscale('log')
  81
                     plt.legend(fontsize = 18)
  82
                     plt.savefig('figs/cl_err.pdf', bbox_inches = 'tight')
  83
                     plt.show()
  84
  85
             def run_q3():
  86
                     plt.figure(figsize=(8,4))
                     plt.plot(range(2500), jacobisol(0.3), color = 'black', lw = 2, label = r'$\omega \( \_=\_0.3\$'); \( \text{print}('Omega \( \_=\_0.3\_-\_\Done') \)
  87
                     plt.plot(range(2500), jacobisol(0.6), color = 'blue', lw = 2, label = r'$\omega_\
=\u0.6$'); print('Omega_\=\u0.6\u0.6\u0.1\u0.00ne')
  88
  89
                      plt.plot(range(2500), jacobisol(1.0), color = 'gray', lw = 2, label = r'$\omega_\( \)
                                 = 1.0; print('Omega_=1.0,-Done')
                     plt.xlabel(r'Iteration_Number', fontsize = 16)
  90
                     plt.ylabel(r'$L_2$_Residual_Norm_Error', fontsize = 16)
  91
  92
                     plt.yscale('log')
                     plt.legend(fontsize = 18)
  93
  94
                     plt.savefig('figs/jacobi_12.pdf', bbox_inches = 'tight')
                     plt.show()
  95
  96
  97
                     plt.figure(figsize=(8,4))
                     plt.plot(range(2500), gausssol(2500, 3, 0.5), color = 'black', lw = 2, label = r '$\omega_{\pu}_0.5\$'); print('Omega_{\pu}_0.5_{\pu}_0.5_{\pu}')
  98
  99
                      plt.plot(range(2500), gausssol(2500, 3, 1.0), color = 'blue', lw = 2, label = r'
                                 $\omega_=\_1.0$'); print('Omega_=\_1.0\_-\_Done')
100
                      plt.plot(range(2500), gausssol(2500, 3, 1.5), color = 'gray', lw = 2, label = r'
                                 $\omega_=_1.5$'); print('Omega_=_1.5_-_Done')
101
                     plt.xlabel(r'Iteration_Number', fontsize = 16)
102
                     plt.ylabel(r'$L_2$\_Residual\_Norm\_Error', fontsize = 16)
                     plt.yscale('log')
103
                     plt.legend(fontsize = 18)
104
                     plt.savefig('figs/gauss_12.pdf', bbox_inches = 'tight')
105
106
                     plt.show()
107
108
             def run_q4():
109
                     p = 1; viter = 25
                     ax = plt.figure(figsize=(10,5)).gca()
110
                      ax.xaxis.set_major_locator(MaxNLocator(integer=True))
111
112
                      for pval in range(2, 6):
                              U, 12_2 = vcyclesol(p, pval, viter, 2, 2, 50)
plotlabel = r'$p$_{\square}_{\square}' + str(pval)
113
114
                     plt.plot(range(1, viter+1), 12_2, lw = 2, label = plotlabel)
plt.xlabel(r'Multi-grid_Cycles', fontsize = 16)
115
116
                     plt.ylabel(r'$L_2$\_Residual\_Norm', fontsize = 16)
117
                     plt.yscale('log')
118
119
                     plt.legend(fontsize = 18, ncol = 4)
120
                     plt.savefig('figs/vcyc_12_err.pdf', bbox_inches = 'tight')
121
                     plt.show()
122
123
124
                     p = 0; pmax = 4; viter = 50
125
                     nu1 = 10; nu2 = 10; nuc = 1000
126
                     U, resid_norm = vcyclesol(p, pmax, viter, nu1, nu2, nuc)
127
                     workunits = np.zeros(viter)
128
                     for 1 in range(viter-1):
129
                               workunits[1+1] = workunits[1] + 2**(pmax*1/viter + 3)/(nu1 + nu2)
```

```
130
        plt.figure(figsize=(8,4))
131
       132
133
        plt.xlabel(r'Work_Units', fontsize = 16)
134
       plt.ylabel(r'$L_2$\_Residual\_Norm', fontsize = 16)
plt.yscale('log')
135
136
        plt.legend(fontsize = 18)
137
138
        plt.savefig('figs/vcyc_gs.pdf', bbox_inches = 'tight')
    plt.show()
def main():
139
140
141
        run_q1()
142
        run_q2()
       run_q3()
143
144
        run_q4()
    if __name__ == "__main__":
    main()
145
146
```