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ELECTRIC POTENTIALS AND FIELDS: LAPLACE'S EQUATION

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Abstract: This paper shows how to numerially calculate Laplace's equation in two-deimensional square. Then picture the electric field distribution and compared two approximately method-the relaxation method and SOR method. Finally, discuss the relationship between the parameter **alpha** and the number of boundary number in SOR method.

Keyword:electric firld, Laplcae's equation, numerially calculate, the SOR mehod, the relaxation method

BAKGROUND

In regions of space that do not contain any electric charges, the electric potential obeys Laplace's equution.

$$\frac{\partial^2 V}{\partial x^2} + \frac{\partial^2 V}{\partial y^2} + \frac{\partial^2 V}{\partial z^2} = 0$$

From a numerical perspective the situation here is a bit different from anything we have encountered so far, All of our problems to this point have involved differential equations for which several initial conditions were given, and we were able to use the **Euler method** or something like it to calculate the behavior for later times. However, with Laplace's equation we are generally given some boundary conditions for V, which specify its value on a surface in x-y-z space. Alternatively, the boundary conditions might be given in terms of the electric field, which is proportional to the gradient of **V**. In either case our problem is to find the function V(x,y,z) that satisfies both Laplace's equation and the specified

boundary conditions. Most importantly, satisfying the boundary conditions will not be as easy as with the ordinary differential equations encountered in previous chapters.

MODEL

As usual we discretize the independent variables, in this case x, y and z. Points in our space are then specified by integers i, j, and k, with x=i*delta x, y=j*delta y, z=k*delta k. Our goal is to determine the potential V(i, j, k) = V(x=i*delta x, y=j*delta y, z=k*delta k) on this lattice of points. The first step in reaching this goal is to rewrite

$$\frac{\partial^2 V}{\partial x^2} + \frac{\partial^2 V}{\partial y^2} + \frac{\partial^2 V}{\partial z^2} = 0$$

as a difference equation. We already know how to write a first derivative in finite-difference form. For example, at the point (i, j, k) the derivative with respect to x may be written as

$$\frac{\partial V}{\partial x} \approx \frac{V(i+1, j, k) - V(i, j, k)}{\Delta x}$$

$$V(i,j,k) = \frac{1}{6} [V(i+1,j,k) + V(i-1,j,k) + V(i,j+1,k) + V(i,j-1,k) + V(i,j,k+1) + V(i,j,k-1)]$$

where we have assumed that the step sizes along x, y, and z are all the same (**delta** x = **delta** y = **delta** z). In words, these simply says that the value of the potential at any point is the average of V at all of the neighboring points. The solution for V(i, j, k) is the function that manages to satisfy this condition at all points simultaneously.

From the symmetry it is natural to suppose

that the potential on these surfaces of the box will vary linearly with position as we move from $\mathbf{x=-1}$ to $\mathbf{x=+1}$. We assume further that the box is infinite in extent along $+-\mathbf{z}$, so $\mathbf{V(i,j,k)}$ is independent of \mathbf{k} . We thus have only a two-dimensional problem. Our goal is to find the potential function $\mathbf{V(i,j)}$ that satisfies

$$V(i, j, k) = \frac{1}{4} [V(i+1, j, k) + V(i-1, j, k) + V(i, j+1, k) + V(i, j-1, k)]$$

Finally, if we wish to obtain the electric field, this can be calculated by differentiating **V(i,j)**. To do this we use the fact that the component of **E** in the **x** direction is

$$E_x = -\frac{\partial V}{\partial x}$$

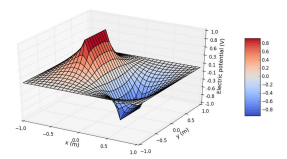
with corresponding relations for **E_y** and **E_x**. These dericatives can be estimated using our usual finite difference expressions. In this case we can use a symmetric form for this derivative

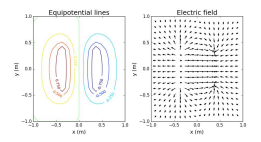
$$E_x(i, j) = -\frac{V(i+1, j, k) - V(i-1, j, k)}{2\Delta x}$$

but a lesss symmetric form would give essentially the same results. One note of caution is that **E** at the boundary needs to be calculated using a one-sided difference equation since, clearly, using values of **V** at sites beyond the boundary makes no sence.

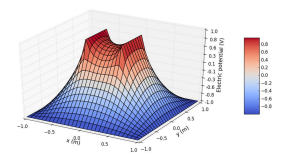
Another interesting use of the relaxation algorithm is the problem of the potential between two parallel capacitor plates. The case of infinite plates can, of course, be handled analytically using **Gauss' law**. However, here we are intereted in what

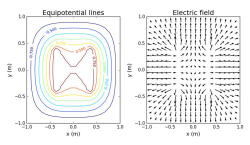
happens when the plates are finite in extent. Again, we can modify our earlier program to handle this case. All we need to do is set up the proper boundary conditions for V. We set the plates to V=+-1, and the square boundary defined by x=+-1, y=+-1 surrounding the plates is set to V=0. In analytic calculations we would generally apply the condition V=0 at **x,y** approximate to indinite, but that is usually not pracical in a numerical treatment. Here, for simplicity, we apply this condition on the square boundary just describled; we will have more to say about how to choose such boundary regions and their effects in the next section. After specifying the boundary conditions on V, the application of the relaxation algorithm is the same as that outlined above. The results are givem in





And we will also show in the following with V1=V2=1,V boundary=0





The electric field is seen to be largest between the two plates. In that region the field is approximately uniform (although this is hard to verify with the rather coarse scale used for this plot) and is directed from high to low potential. The fring fields at the edges of and outside the plates are also evident.

METHOD

We now require a numerical strategy for determining this function, assuming only that **V** is known at the boundaries. We can't just start at one of the boundaries and work our way into and across the system since according to:

$$V_0(i,j,k) \rightarrow V_1(i,j,k) \rightarrow \cdots \rightarrow V_n(i,j,k)$$

we need to know V at all of the neighbors in

order to calculate its value at any particular point. The approach we take is to begin with some initial guess for the solution; call it V_0(i,j,k). In general, unless we are extremely clever, the guess we make will not satisfy the above everywhere. To obtain an improved guess, we use it to calculate new values of V, using V_0 on the right-hand side. The guess V_0, together with it yields, a new and we hope improved guess, V_1(i,j,k). We then repeat the procedure with V_1 to obtain an even better guess, V_2, etc. This iterative process is continued until our result satisfies some convergence criteria, which we will discuss shortly. The general approach is called the relaxation method, and is a useful way to deal with several important classes of partial differential equations. There are different ways to implement the relaxation method, and some are much better than others with respect to speed of convergence, etc.

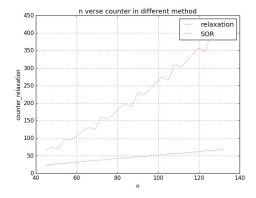
The problem with the slow convergence of the Jacobi and Gauss-Seidel methods can be overcome with a method known as simultaneous over-relaxation(SOR), which can be appreciated as follows. Let V(i,j) be the new value of the potential calculated using the Gauss-Seidel method. We can then think of

$$\Delta V(i,j) = V^*(i,j) - V_{old}(i,j)$$

as the change recommended

$$V_{new}(i,j) = \alpha \Delta V(i,j) + V_{old}(i,j)$$

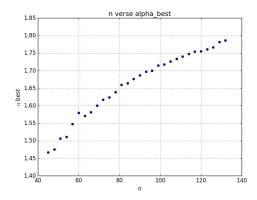
Where **alpha** is a factor that measures how much we "over-relax". So we compared the calculated numers of these methoods.



We found the **SOR method** is better compared with the **relaxation method** in calculated number or time.

Choosing alpha=1 yields the Gauss-Seidel method. It turns out that if alpha>=2, the method does not converge, while alpha<1 correspondings to "under-relaxtion." The algorithm known as simultaneous over-relaxation with 1<alpha<2. It remains for us to determine the best value of alpha; that is, the value that yields the fastest convergence.

We also found the relationship between alpha and n in following



For a problem on a two-dimensional square grid like the ones we have considered and with fixed-value boundary conditions (called **Dirichlet** boundary conditions) on all four sides, the best choice for alpha is

$$\alpha \approx \frac{2}{1 + \pi / L}$$

REFERENCE

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[2]Thanks for the code to ChenYangyao/computationalphysics_N20133 01020169