## 11 — PHY 494: Assignment 11 (20 points total)

Due Wednesday, April 24, 2019, 11:59pm.

Submission is to your **private GitHub repository**.

Enter the repository and run the script scripts/update.sh (replace *YourGitHubUsername* with your GitHub username):

cd assignments-2019-YourGitHubUsername
bash ./scripts/update.sh

It should create three subdirectories assignment\_11/Submission, assignment\_11/Grade, and assignment\_11/Work and also pull in the PDF of the makeup and an additional file.

To submit your makeup assignment, commit and push Python code inside the assignment\_11/Submission directory. Commit any other additional files exactly as required in the problems.

Code will be tested against the unit tests in test\_capacitor.py. The grade will be approximately proportional to the number of tests that pass successfully so your code *must* be able run under the tests.

## 11.1 Numerical solution of the plate capacitor potential (20 points)

Compute the electrostatic potential of a plate capacitor in a grounded box, shown in Fig. 1. Solve the problem in 2D. The square box has sides of length 100. Consider the capacitor plates to be conducting, very thin sheets of metal. The plates are held at an electric potential of +100 V and -100 V. The plates are w = 50 units wide and located at a distance d = 20. The capacitor is centered inside the box.

Submit your code as a python module capacitor.py. In particular, you should have the following functions:<sup>2</sup>

<sup>&</sup>lt;sup>1</sup>If the script fails, file an issue in the Issue Tracker for PHY494-assignments-skeleton and just create the directories manually.

<sup>&</sup>lt;sup>2</sup>See also Lesson 16 and in particular the notebook 16\_PDEs-2.ipynb.

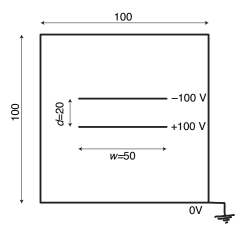


Figure 1: Schematic of the capacitor in the grounded box problem (2D). The capacitor is centered in the box. The capacitor plates are thin, conducting sheets of metal that are held at potential +100 V and -100 V, respectively.

**set\_boundaries (Phi)** Given the potential on the lattice, enforce the boundary conditions by setting the appropriate lattice points to the prescribed values.

calculate\_phi\_capacitor(Max\_iter=10000, tol=1e-3) This function should calculate the potential  $\Phi$  on the lattice. It needs to take at least the two keyword arguments Max\_iter to set the maximum number of iterations and tol to set the convergence criterion.

You must implement a convergence check: consider the solution converged when the Frobenius norm

$$||\Phi^{\text{new}} - \Phi^{\text{old}}|| := \sqrt{\sum_{i=1}^{M} \sum_{j=1}^{N} \left| \Phi_{ij}^{\text{new}} - \Phi_{ij}^{\text{old}} \right|^2}$$
 (1)

is less than the set tolerance tol. Report the number of iterations

that were required. Also report if convergence was not achieved within Max\_iter iterations.

The function must return the potential  $\Phi$  as a number array of dimensions (100, 100).

It must be possible to run these functions as

## import capacitor

```
Phi = capacitor.calculate_phi_capacitor(Max_iter=2000, tol=1)
```

Start from the skeleton code in capacitor.py to ensure that your functions adhere to the prescribed interface.<sup>3</sup>

Specifically, you need to fullfil the following objectives:

(a) Your code must produce correct results, as tested with test\_capacitor.py. You can run these tests yourself with

```
pytest -v test_capacitor.py
(in the same directory as your capacitor.py). [15 points]
```

- (b) Converge your results to tol = 1e-3. Include in your submission:
  - a) Report how many iterations you required for convergence. Simply write the number in a *textfile* iterations.txt [3 points]
  - b) Make a 3D plot of the potential (you can use capacitor.plot\_phi(Phi)) [1 points]
  - c) Make a 2D plot of the initial potential (boundary conditions) and of the final converged solution (you can use capacitor.plot\_panel(Phi)) [1 points]

Note: You don't have to submit any notebooks or written text. It will be sufficient to submit

<sup>&</sup>lt;sup>3</sup>The skeleton code contains a optimized function Laplace\_Gauss\_Seidel\_odd\_even() which replaces the Gauss-Seidel code from the lecture. It runs 50 times faster so you are advised to use it...but you may use the slower code, if you prefer. The original code (as developed in class) is in the function Laplace\_Gauss\_Seidel().

- capacitor.py
- iterations.txt
- capacitor\_potential\_2d.pdf and capacitor\_potential\_3d.pdf (these are the default filenames when using the functions in capacitor.py).