## 1 — PHY 494: Makeup assignment 1 (25 points total)

Due Saturday, April 30, 2016, 5pm.

This is a **optional Makeup Assignment**. If you choose to hand it in by the deadline, it will be graded like a normal homework assignment. If its grade is better than your worst homework grade then it will replace that grade.

Submission is to your private GitHub repository.

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

cd assignments-2016-YourGitHubUsername
bash ./scripts/update.sh

It should create three subdirectories makeup\_01/Submission, makeup\_01/Grade, and makeup\_01/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 makeup\_01/Submission directory. Commit any other additional files exactly as required in the problems.

Failure to adhere to the following requirements may lead to homework being returned ungraded with 0 points for the problem.

- Only submit code.
- All code should be in a file makeup01.py.
- Code will be tested against the unit tests in test\_makeup01.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. (Failing tests for the Bonus problem can be ignored.)

## 1.1 Factorial function (10 points)

Write a function factorial (n) that

- 1. calculates the factorial n! for any integer number  $n = 0, 1, 2, \ldots$
- 2. raises ValueError if n is negative
- 3. raises TypeError if n is not an integer.

<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.

## 1.2 ODE with Scipy (15 points)

Use the Scipy function scipy.integrate.odeint() to solve the following ordinary differential equation

$$-\frac{1}{2}\frac{d^2\psi_n(x)}{dx^2} + \left[\frac{1}{2}x^2 - E_n\right]\psi_n(x) = 0 \tag{1}$$

$$\psi_n(0) = 1; \quad \frac{d\psi_n(0)}{dx} = 0 \tag{2}$$

$$E_n = n + \frac{1}{2}, \quad n = 0, 2, 4, 6, \dots$$
 (3)

for the real function  $\psi_n(x)^2$  and the three values n = 0, n = 2, and n = 8. The code should contain the following functions:

1. ode\_qmhosc() solves the ODE Eq. 1:

```
psi = ode_qmhosc(x, psi0, dpsidx0, n=n)
```

that takes as arguments

- all the values x at which the solution should be evaluated (the first value must be the one for which the initial conditions are given, i.e., x = 0 for this problem),
- as initial conditions the function value and first derivative at x=0 (Eq. 2), and
- the value of n that determines  $E_n$  (Eq. 3).

It should return the function values  $\psi_n(x)$  as a numpy array.

2. **f\_qmhosc()** is needed to transform the ODE Eq. 1 into standard form so that it can be solved with scipy.integrate.odeint().

$$f_qmhosc(y, t, E=0)$$

should produce the ODE standard  $form^3$  of the derivative vector  $\mathbf{f}$  when provided with

- the current values of y
- $\bullet$  the current value of t
- and the parameter  $E_n$  (see Eq. 1).

$$\frac{d\mathbf{y}}{dt} = \mathbf{f}$$

and in the case of Eq. 1, we would have t=x,  $y_0=\psi_n(x)$ , and  $y_1=\psi_n'(x)\equiv\frac{d\psi_n}{dx}$ .

<sup>&</sup>lt;sup>2</sup>If you have done Quantum Mechanics 1 (PHY 314) then you should recognize it as the Schröinger equation for the simple harmonic oscillator.

<sup>&</sup>lt;sup>3</sup>Remember that the ODE standard form is

- (a) Numerically solve Eq. 1 on a lattice with  $\Delta x = 0.01$  for  $0 \le x < 6$  for
  - n = 0
  - n = 2
  - n = 8
  - (i) The tests in test\_makeup01.py should all pass to show that you correctly implemented a solution. [12 points]
  - (ii) Plot all your solutions in one figure and include the figure as a PDF named qmhosc.pdf. [3 points]
- (b) Bonus: Correct physical<sup>4</sup> solutions for Eq. 1 are symmetric  $(\psi(x) = \psi(-x))$  for even n and antisymmetric  $(\psi(x) = -\psi(-x))$  for odd n. This implies that the initial conditions for  $n = 1, 3, 5, \ldots$  are different from the ones for even n (Eq. 2). Choose appropriate initial conditions<sup>5</sup> and write a function

```
phi0, dphidx0 = initial_values_qmhosc(n)
```

that produces the correct initial values depending on the value of n.

Using your initial values, solve Eq. 1 for n = 3 and plot the solution (include a PDF named qmhosc\_odd.pdf). [bonus +4\*]

<sup>&</sup>lt;sup>4</sup>Physical solutions for the wavefunction  $\psi_n(x)$  have to be normalizable, i.e., they have to decay to zero for  $x \to \pm \infty$ . The numerical solutions only fulfill this requirement for small x and even for moderately large x they start to diverge. It would be better to use specialized algorithms that have the normalizability requirement built in.

<sup>&</sup>lt;sup>5</sup>Choose  $\psi \ge 0$ ,  $\frac{d\psi}{dx} \ge 0$  and either use values of 0 or 1 because this is the convention employed in the