

# Assignment 4

[https://github.com/jchryssanthacopoulos/quantum\\_information/tree/main/assignment\\_4](https://github.com/jchryssanthacopoulos/quantum_information/tree/main/assignment_4)

## Quantum Information and Computing AA 2022–23

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# Quantum Harmonic Oscillator



- Goal is to solve 1D Schrodinger equation,  $\hat{H}|\psi_n\rangle = E_n|\psi_n\rangle$
- Hamiltonian was discretized using finite difference. For  $m = \omega = \hbar = 1$ , this becomes eigenvalue problem

$$\begin{pmatrix} \frac{x_1^2}{2} + \frac{1}{\Delta x^2} & -\frac{1}{2\Delta x^2} & 0 & \dots & 0 \\ -\frac{1}{2\Delta x^2} & \frac{x_2^2}{2} + \frac{1}{\Delta x^2} & -\frac{1}{2\Delta x^2} & \dots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & \dots & 0 & -\frac{1}{2\Delta x^2} & \frac{x_N^2}{2} + \frac{1}{\Delta x^2} \end{pmatrix} |\psi_n\rangle = E_n |\psi_n\rangle$$

where  $\{x_i\}$  are points between  $[-x_{\max}, x_{\max}]$  and  $\Delta x = 2x_{\max}/N$

```
! get spacing
dx = (xmax - xmin) / (npoints - 1)

! discretize Hamiltonian
H_off_diag = -1 / (2 * dx ** 2)

do ii = 1, npoints
  x_grid(ii) = xmin + (ii - 1) * dx
  H_diag(ii) = 1 / (dx ** 2) + 0.5 * x_grid(ii) ** 2
end do

! compute eigenvalues and eigenvectors
call dsteqr("I", npoints, H_diag, H_off_diag, eigenvectors, npoints, work, info)

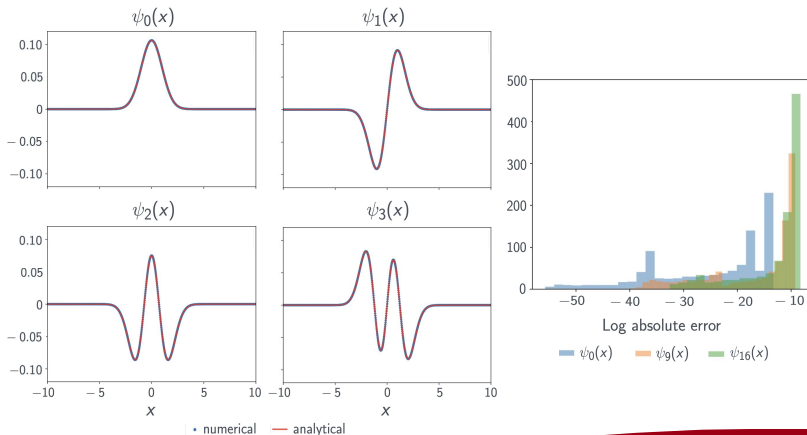
if (info .eq. 0) then
  print *, "Success!"
else
  print *, "Failed to obtain eigenvalues and eigenvectors"
end if
```

dsteqr computes eigenvalues and eigenvectors for tridiagonal matrix, results saved in file

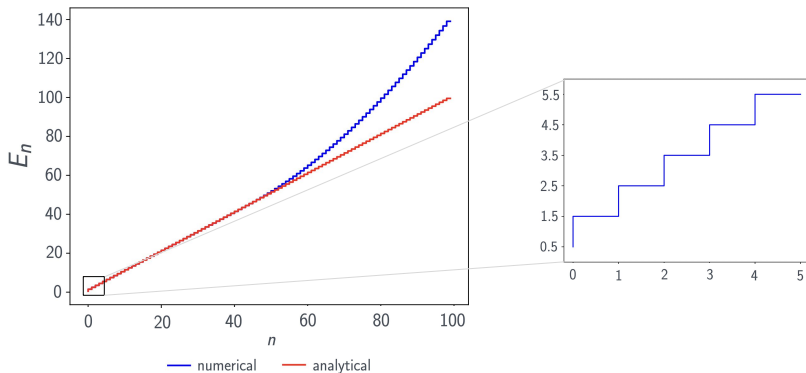
```
for N in 100 1000 5000; do
  for xmax in 2.5 5 10 15; do
    echo "Running for N = $N, xmin = -$xmax, xmax = $xmax ..."
    compiled/eigen_schrodinger \
      --output_filename data/solution_${N}_${xmax}.txt \
      --npoints $N \
      --xmin -$xmax \
      --xmax $xmax
  done
done
```

Program accepts discretization parameters and output filename, results analyzed in Jupyter notebooks

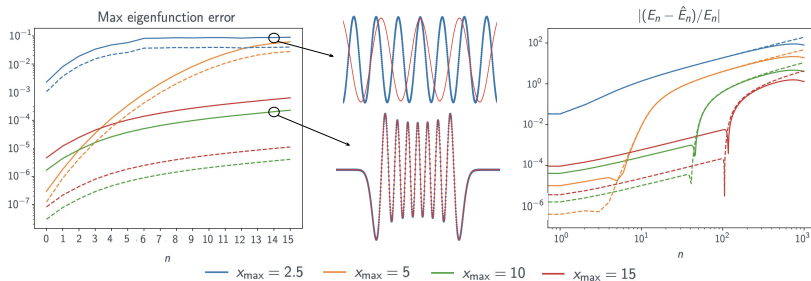
- Eigenfunctions given by  $\psi_n(x) = \frac{1}{\sqrt{2^n n!}} \left(\frac{1}{\pi}\right)^{1/4} \exp(-x^2/2) H_n(x)$
- Good match to expected values using  $N = 1000$  and  $x_{\max} = 10$ , but error increases with  $n$ , particularly around edges of domain boundaries



- Eigenvalues given by  $E_n = n + \frac{1}{2}$
- Good match to expected values, but again, error increases with  $n$



- Error decreases as  $N$  increases (solid lines are  $N = 1000$ , dashed lines are  $N = 5000$ )
- Error decreases as  $x_{\max}$  increases, but then increases again. Best overall results found for  $x_{\max} = 10$



- **Correctness.** Results closely match analytical solutions for  $n < \mathcal{O}(100)$ , but start to diverge for higher values. Loss of accuracy is due to discretizing continuous problem
- **Stability.** Code is stable, and `dsteqr` returns with `info = 0`. Results are reproducible across runs
- **Accurate Discretization.** Accuracy can be improved by making  $\Delta x$  smaller (i.e., increasing  $N$ ), since error scales as  $\Delta x^2$
- **Flexibility.** Other potentials can easily be substituted in. It is harder to accommodate more dense discretization schemes, and extend to multidimensional and time-dependent problems
- **Efficiently.** Hamiltonian is tridiagonal, so diagonalization is efficient. Times on 10-core M1 Macbook Pro 32 GB below

$N$	Run time (s)
$1.0 \times 10^3$	0.582
$2.5 \times 10^3$	9.269
$5.0 \times 10^3$	70.66
$1.0 \times 10^4$	536.8