

# CIS and TDHF - Homework

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## Data Availability

The source code of this project as well as the example input files are available at the corresponding [Github page](#). There one could also find the example output files and the detailed instructions on how to run and compile the code. The example output files are the outputs of the CIS and TDHF calculations. The source code of this project is written in **Fortran90**.

## Code Overview

### Input

In order to run the code, one should provide two input files, namely **wavepacket** and **potential**. It's important to name the files this way.

The **wavepacket** file should contain the following parameters: number of lattice points, initial position of the wavepacket, the initial width, propagation time step, total number of steps, snapshot frequency, total number of coefficients and the coefficients themselves.

The **potential** file should specify the type of a potential, e.g. harmonic and double well are supported, as well as the angular frequency in case of the harmonic potential and the barrier height in case of the double well potential. In both input files, each parameter should be written on a separate line. The example input files can be found in **harmonic/wavepacket** and **harmonic/potential**, respectively.

## Wavepacket Initialisation and Normalisation

In this project, instead of using a gaussian wavepacket, the wavepacket is initialised as a superposition of the eigenstates of the harmonic oscillator. At  $t = 0$ , the wavepacket is given by the following expression:

$$\Psi(x, t = 0) = \sum_{n=0}^{\infty} c_n \Phi_n(x) \quad (1)$$

Where the eigenstates of the quantum harmonic oscillator are calculated according to the following formula:

$$\Phi_n(x) = \frac{1}{\sqrt{2^n n!}} \left( \frac{m\omega}{\pi} \right)^{1/4} e^{-m\omega x^2/2} H_n(x\sqrt{m\omega}) \quad (n = 0, 1, 2, \dots) \quad (2)$$

$H_n$  is the Hermite polynomial of order  $n$ . We'll touch upon the implementation of the Hermite polynomials in the next section. While initialising the wavepacket according to the Equation 1 and Equation 2 as shown in Listing 1, the calculation of a norm of the wavepacket is performed alongside. The code snippets for the wavepacket initialisation are provided below.

```

104 norm = 0.0d0
105 nmax = size(coeff) - 1
106 do i = -npoints/2 + 1, npoints/2
107     x = dble(i) * dx
108     if (i > 0) then
109         j = i
110     else
111         j = i + npoints
112     endif
113     psi0(j) = 0.0d0
114     do n = 0, nmax
115         call factorial(n, fact)
116         psi0(j) = psi0(j) + &
117             coeff(n+1) * &
118             (1.0d0 / sqrt(2.0d0**n * fact)) * &
119             (mass * angfreq / pi)**0.25d0 * &
120             exp(-mass * angfreq * alpha * (x-x0)**2.0d0 / 2.0d0) * &
121             hermite(n, (x-x0) * sqrt(mass * angfreq))
122     end do
123     norm = norm + abs(psi0(j))**2.0d0 * dx
124 end do

```

Listing 1: Initialisation of the wavepacket in the `initpsi` subroutine.

As soon as the norm of the wavepacket is calculated, the wavepacket is normalised as shown in Listing 2. The normalisation is performed by dividing each element of the wavepacket by the square root of the norm. This ensures that the wavepacket is properly normalised.

```

126 norm = 1.0d0 / sqrt(norm)
127 do i = -npoints/2 + 1, npoints/2
128     x = dble(i) * dx
129     if (i > 0) then
130         j = i
131     else
132         j = i + npoints
133     endif
134     psi0(j) = norm * psi0(j)
135 end do

```

Listing 2: Normalisation of the wavepacket in the `initpsi` subroutine.

Due to the need of computing the factorials, the corresponding subroutine is implemented as shown in Listing 3. The factorial is calculated up to the  $n$ th order.

```

148 result = 1.0d0
149 do i = 1, n
150     result = result * i
151 end do

```

Listing 3: Calculation of the factorial up to the  $n$ th order in the `factorial` subroutine.

## Hermite Polynomials

The Hermite polynomials are calculated using the following recurrence relation:

$$H_n(y) = 2yH_{n-1}(y) - 2(n-1)H_{n-2}(y) \quad (H_0(y) = 1, H_1(y) = 2y) \quad (3)$$

Where  $H_0$  isn't calculated explicitly. The reason to use Hermite polynomials in the wavepacket initialisation is due to the fact that the eigenstates of the quantum harmonic oscillator are expressed in terms of the Hermite polynomials as shown in [Equation 2](#). Specifically, the physicist's Hermite polynomials are used in this project. The code snippet for the calculation of the Hermite polynomials is provided in [Listing 4](#).

```
166 if (n == 0) then
167   Hn = 1.0d0
168 elseif (n == 1) then
169   Hn = 2.0d0 * y
170 else
171   H0 = 1.0d0
172   H1 = 2.0d0 * y
173   do i = 2, n
174     H_prev = H1
175     H1 = 2.0d0 * y * H1 - 2.0d0 * (i - 1) * H0
176     H0 = H_prev
177   end do
178   Hn = H1
179 endif
```

Listing 4: Calculation of the Hermite polynomials up to the  $n$ th order in the `hermite` function.

## Results and Discussion

The wavepackets were propagated at the bottom of the harmonic potential with the angular frequency  $\omega = 0.2825 \text{ fs}^{-1}$ . Moreover they were propagated for 5000 steps with a 0.1 fs time step. The initial width of the wave packet was equal to  $1.0 \text{ Bohr}^{-2}$  and the snapshots were taken each 10th step. The wavepackets were propagated with different number of eigenstates, namely 1, 2, 3, and 5. The time evolution of the wavepackets is shown in ??.

The wavepackets with a single eigenstate are stationary, whilst the wavepackets with multiple eigenstates are oscillating. The oscillations are due to the fact that the wavepacket is a superposition of the eigenstates of the harmonic oscillator. The more eigenstates are included in the wavepacket, the more oscillations are observed. The wavepacket with 5 eigenstates oscillates the most. The oscillations are periodic and the period of the probability density is calculated below.

The period of the probability density for the harmonic oscillator case can be calculated from the angular frequency as follows:

$$T = \frac{2\pi}{\omega} \quad (4)$$

Therefore the periods  $T$  of the corresponding probability densities are equal to 22.23 fs.

Speaking of the symmetry, the wavepackets with the even number of eigenstates have symmetric probability densities of the wavefunction. On the other hand, the odd number of eigenstates results in the probability densities being asymmetric as can be seen in ??.

The underlying reason for this is the wavefunction itself. The solutions alternate between even and odd functions around the  $x = 0$  resulting in different symmetries of the probability densities [? ].

While the particle with one eigenstate spends the majority of time at the bottom of the potential, the particles with multiple eigenstates oscillate. The chance of finding the particle outside of the potential increases as well. For instance, for a particle with one eigenstate this probability is around 16 % [? ].

## Generative AI Usage

In this work, [ChatGPT](#) large language model was used to check grammar and spelling of the main body of text, whilst [Perplexity](#) was utilised for the sake of literature search.

## Acknowledgments

This project is based on data and instructions provided by APina Romaniello available at the aforementioned [Github page](#).