Project 2 in FYS3150

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https://github.com/UlrikSeip/Projects/tree/master/prosjekt2

1 ABSTRACT

In this project we have implemented Jacobian algorithm, to find eigenvectors, and their corresponding eigenvalues in tridiagonal matrices. We then used this to model a harmonic oscillator problem in three dimensions, with one and two electrons. This turned out to be a computationally heavy, but relatively accurate method.

2 INTRODUCTION

Finding eigenvectors analytically is complicated, can be tedious, and can sometimes be impossible, and this is why it is much more convenient to do so numerically. A common way of doing this is by the application of the Jacobian method. Essentially we rotate one matrix element at a time, always taking the one with the highest absolute value, until all but the diagonal elements are essentially zero. All transformations are also applied to an identity matrix that then turns into our eigenvectors.

The modelling of an electron in a harmonic oscillator potential is one of the most classical problems in quantum mechanics. This has a simple analytical solution, however the interaction between multiple electrons can be harder to solve analytically, but does follow similar principles. Therefore any good numerical solution of the first problem, should very easily be able to calculate the interaction between two or more electrons. What we have done in this experiment is to customize the method described above to solve for one electron, and then used that same method on two electrons.

3 METHOD

3.a Implementing the Jacobian algorithm

The implementation follows a standard recipe:[1] We start with the relation

$$\cot 2\theta = \tau = \frac{a_{ll} - a_{kk}}{2a_{kl}}.$$

This can be used to find the angle θ that makes the non-diagonal matrix elements of the transformed matrix $a_{kl} = 0$. The quadratic equation is obtained using $\cot 2\theta = 1/2(\cot \theta - \tan \theta)$.

$$t^2 + 2\tau t - 1 = 0 (1)$$

Which gives us

$$t = -\tau \pm \sqrt{1 + \tau^2} \tag{2}$$

c and s are then obtained by

$$c = \frac{1}{\sqrt{1+t^2}}\tag{3}$$

and

$$s = tc (4)$$

We then use the rotational factors c and s to rotate every other element in the matrix according to their position, giving us a new diagonal, that is slightly closer to the eigenvalues, and new matrix elements elsewhere, slightly closer to 0. The same is done for a matrix that tarts out as an identity matrix, with the purpose of transforming it into the original matrix' eigenvectors. This is all done in the programming language Julia for high efficiency. See the documentation in section 6.a for further explanation.

3.b Testing the code

For testing the algorithm we have implemented two tests. One for checking if the largest element in the matrix is correctly located, and one for testing if the resulting eigenvalues are correct. The first one is more useful for development purposes, whilst the second one is essential for validating that our implementation works correctly. See section 6.b for the test functions.

3.c Quantum dots in three dimensions, one electron

Now that we had a general algorithm we used it to model a electron that moves in a three-dimensional harmonic oscillator potential. In other words, we looked for the solution of the radial part of Schroedinger's equation for one electron, which reads

$$-\frac{\hbar^2}{2m} \left(\frac{1}{r^2} \frac{d}{dr} r^2 \frac{d}{dr} - \frac{l(l+1)}{r^2} \right) R(r) + V(r) R(r) = ER(r). \tag{5}$$

This problem also has analytical solutions, so we can test how accurate our algorithm is.

We decided to limit our experiment to the ground state of l=0. After introducing the constant $\alpha=\left(\frac{\hbar^2}{mk}\right)^{1/4}$, and the variables $\lambda=\frac{2m\alpha^2}{\hbar^2}E$ and $\rho=r/\alpha$, the Schroedinger's equation becomes

$$-\frac{d^2}{d\rho^2}u(\rho) + \rho^2 u(\rho) = \lambda u(\rho). \tag{6}$$

See section 6.c for futher details.

Since we are working in radial coordinates we have $\rho \in [0, \infty)$. Since we can't represent infinity on a computer we have to find an approximation, which we will come back to later. For now we define $\rho_{min} = 10^{-6}$ and ρ_{max} to represent the minimum and maximum values of ρ . We used 10^{-6} instead of 0 to avoid any potential divisions by zero.

Function 6 is an differential equation that can be modeled similarly to earlier. If we have N mesh points we get a step length

$$h = \frac{\rho_{max} - \rho_{min}}{N}. (7)$$

The value of ρ at a point i is then

$$\rho_i = \rho_0 + ih \qquad i = 1, 2, \dots, N. \tag{8}$$

We can rewrite the Schroedinger equation for a value ρ_i as

$$-\frac{u_{i+1} - 2u_i + u_{i-1}}{h^2} + \rho_i^2 u_i = \lambda u_i). \tag{9}$$

We then introduced

$$d_i = \frac{2}{h^2} + \rho_i^2, (10)$$

and

$$e_i = -\frac{1}{\hbar^2},\tag{11}$$

which gives us

$$d_i u_i + e_{i-1} u_{i-1} + e_{i+1} u_{i+1} = \lambda u_i. (12)$$

We then wrote the latter equation as a matrix eigenvalue problem

$$\begin{bmatrix} d_0 & e_0 & 0 & 0 & \dots & 0 & 0 \\ e_1 & d_1 & e_1 & 0 & \dots & 0 & 0 \\ 0 & e_2 & d_2 & e_2 & 0 & \dots & 0 \\ \dots & \dots & \dots & \dots & \dots & \dots & \dots \\ 0 & \dots & \dots & \dots & e_{N-1} & d_{N-1} & e_{N-1} \\ 0 & \dots & \dots & \dots & \dots & e_N & d_N \end{bmatrix} \begin{bmatrix} u_0 \\ u_1 \\ \dots \\ u_N \end{bmatrix} = \lambda \begin{bmatrix} u_0 \\ u_1 \\ \dots \\ \dots \\ u_N \end{bmatrix}.$$

$$(13)$$

To solve the problem we implemented the matrix in Julia, and ran it through the algorithm in section 6.a, with a tolerance of 10^{-4} . The implementation of the matrix can be seen in section 6.d. What we then did was test different approximations for ρ_{max} and N to find some values that are accurate and don't require a extemly large N. Our goal was to reproduce the analytical eigenvalues with four leading digits after the decimal point for the lowest eigenvalue. We looked both at the accuracy of the lowest eigenvalue, and used the Julia function Statistics.std() on the difference between our numerical approximation and the analytical solutions, to get an idea of how close the results were overall. std() is an function that finds the standar diviation for an array.

3.d Quantum dots in three dimensions, two electrons

We then wanted to study two electrons in a harmonic oscillator well which also interact via a repulsive Coulomb interaction. The Schroedinger equation for two electrons with no repulsive Coulomb interaction is

$$\left(-\frac{\hbar^2}{2m}\frac{d^2}{dr_1^2} - \frac{\hbar^2}{2m}\frac{d^2}{dr_2^2} + \frac{1}{2}kr_1^2 + \frac{1}{2}kr_2^2\right)u(r_1, r_2) = E^{(2)}u(r_1, r_2),$$
(14)

and the repulsive Coulomb interaction between two electrons is

$$V(r_1, r_2) = \frac{\beta e^2}{|\mathbf{r}_1 - \mathbf{r}_2|} = \frac{\beta e^2}{r},$$
(15)

with $\beta e^2 = 1.44 eV nm$. We introduce the relative coordinate $\mathbf{r} = \mathbf{r}_1 - \mathbf{r}_2$, and the center-of-mass coordinate $\mathbf{R} = 1/2(\mathbf{r}_1 + \mathbf{r}_2)$. The Schroedinger equation now reads

$$\left(-\frac{\hbar^2}{m}\frac{d^2}{dr^2} - \frac{\hbar^2}{4m}\frac{d^2}{dR^2} + \frac{1}{4}kr^2 + kR^2\right)u(r,R) = E^{(2)}u(r,R).$$
(16)

The equations for r and R can be separated so that $u(r,R)=\psi(r)\phi(R)$ and the energy is given by the sumof the relative energy E_r and the center-of-mass energy E_R , that is $E^{(2)}=E_r+E_R$. After introducing the variables $\rho=r/\alpha$ and $\omega_r^2=\frac{1}{4}\frac{mk}{\hbar^2}\alpha^4$, the constants $\alpha=\frac{\hbar^2}{m\beta e^2}$ and $\lambda=\frac{m\alpha^2}{\hbar^2}E$, and only looking at $\psi(r)$ we are left with

$$-\frac{d^2}{d\rho^2}\psi(\rho) + \omega_r^2 \rho^2 \psi(\rho) + \frac{1}{\rho} = \lambda \psi(\rho), \tag{17}$$

see section 6.e for more details. Function 17 can be solved in almost the same way as for when we had one electron, only that the diagonal values d_i now are $d_i = \frac{2}{h^2} + \omega_r^2 \rho^2 + 1/\rho$.

We now have an additional unknown value in ω_r . We then studied the cases of $\omega_r = 0.01$, $\omega_r = 0.5$, $\omega_r = 1$, and $\omega_r = 5$, and use the values of ρ_{max} and N we found worked best for one electron.

4 RESULTS

4.a The implementation

We see a fairly linear correlation between the number of matrix elements n^2 and the required amount similarity transformations. The computation time for each matrix was also proportional to n^2 In figure 1 the tolerance for deviation from 0 in the non diagonal elements was 1e - 4. We found this to be the best balance between accuracy and efficiency.

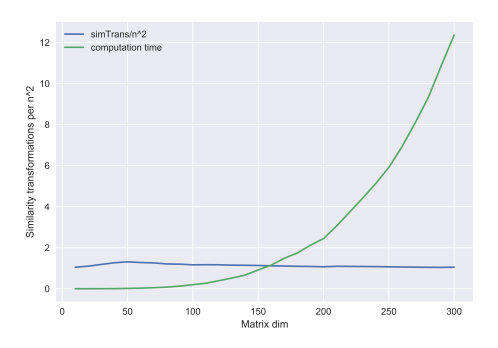


Figure 1: A plot of the required number of rotations as a function of n: $rotations/n^2$, and the time used for calculating the eigenvalues of a $n \cdot n$ matrix.

4.b Quantum dots in three dimensions, one electron

We firstly tested with ρ_{max} equal to all the whole numbers beteen 1 and 100 with N=100. We found that $\rho_{max}=4$ had the closest lowest eigenvalue, with 2.9995 to the analytical 3. This was within our desired four leading digits after the decimal point, however the standard deviation was quite high, as the error increased rapidly. You can see this in figure 2. The lowest std was for $\rho_{max}=14$, but for it the lowest eigenvalue was only 2.99374, so we had to increase N.

We then tested for values of ρ_{max} between 2 and 25, with N=200. Again $\rho_{max}=4$ had the closest lowest eigenvalue, but now $\rho_{max}=15$ had the lowest std. Still, the lowest eigenvalue for $\rho_{max}=15$ was only 2.99824.

We the increased N to 400, which took a lot of processing time. The ρ_{max} with the lowest std was now 23, but it had a lowest eigenvalue of 2.99897, which still wasn't close enough. We also ran the algorithm for $\rho_{max}=23$ and N=500, but it took a really long time, so we didn't run it for any other values. This time the lowest eigenvalue was 2.99934, which was within our desired limit. Due to the long calculation time we chose to use $\rho_{max}=23$ and N=400 when we calculated the interaction between two electrons. A plot of the eigenvalues for $\rho_{max}=23$ can be seen in figure 3, with both N=400 and N=500.

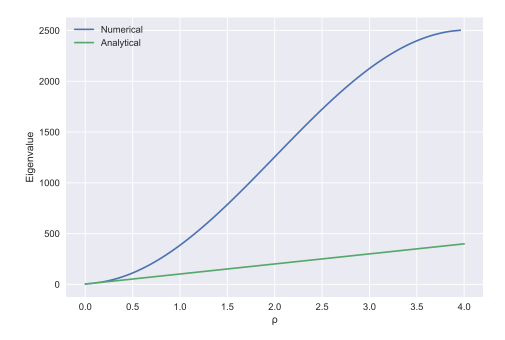


Figure 2: A plot of the numerical solution against the analytical solution, when $\rho_{max} = 4$ and N = 100

4.c Quantum dots in three dimensions, two electrons

The plots of the eigenvales we found with different ω_r can be seen in 4.

5 CONCLUSIONS

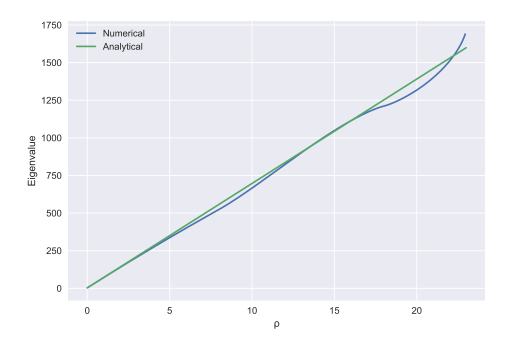
With an accuracy of more digits than our editor cared to print out the Jacobi method, with a tolerance for non diagonal values of up to 1e-4, seems to be an efficient and precise algorithm for computing eigenvectors and eigenvalues. The computation time is proportional to the number of matrix elements, and so the realistic limit for matrix size should be around $10^4 \cdot 10^4$. With $300 \cdot 300$ taking 12 seconds, $10^4 \cdot 10^4$ should take about

$$\frac{(10^4)^2}{300^2} \cdot 12s \approx 4h \tag{18}$$

on a normal laptop. On a supercomputer this would of course be different, and i presume our implementation could have been further vectorised for greater efficiency.

When modeling the electrons we found that values for ρ_{max} that had a low difference between the analytical and numerical solution for the lowest eigenvalue, often had a high standard deviation, and wise versa. From plot 3 we do see that numerical and analytical solutions are very close, while the opposite is true for 2. Overall having a lower standard deviation is probally more important than having a small difference between the analytical and numerical solution for the lowest eigenvalue, but maybe a better compromise could have been achieved.

For the models of two electrons, we see that they vaguely follow a straigth line (or an s-curve maybye). If this is because of ω_r or something else is unknown.



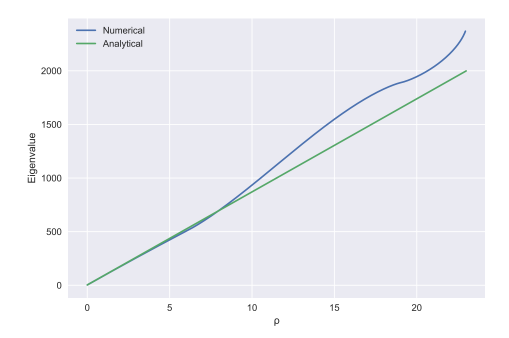


Figure 3: A plot of the numerical solution against the analytical solution, when $\rho_{max}=23$, and N=400 and N=500

6 APENDICES

6.a Integration loop from rotator.jl

```
function maxKnotL(a) #finds larges element that is not on the diagonal on matrix a
    max = 0
    kl = [1,1]
    n = Int64(length(a[1,:]))
    for l = 1:n
        for k = l+1:n
            ma = abs(a[k, 1])
        if (ma > max)
            max = ma
            kl = [k, 1]
        end
```

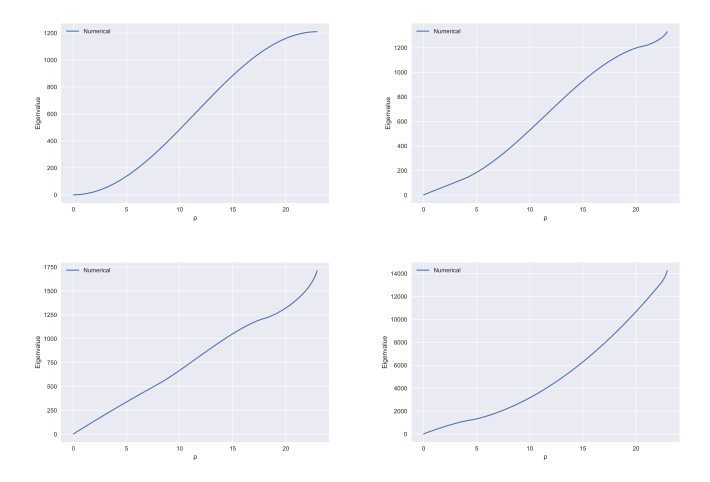


Figure 4: A plot of the numerical solution of numerical solution of two electrons in a harmonic oscillator well which also interact via a repulsive Coulomb interaction with ω_r equal to 0.01, 0.5, 1, and 5 respectively

```
end
   end
   return kl[1], kl[2]
                                       #k, 1
end
function rotate(a, tol)
                                  #the actual rotation function. a = the matrix, tol = tolerance for non diagonal element
                                  #initiates n for later use
   n = Int64(length(a[1,:]))
   r = Matrix{Float64}(I, n, n)
                                 #initialising eigenvector matrix
   counter = 0
                                  #teller antall "similarity transformaitons"
   k, 1 = maxKnotL(a)
                                  #finds indices of matrix element with highest value
   countermax = n^3
   while (abs(a[k ,1]) > tol) && (counter < countermax) #this is the actual loop
       rotation(a, r, k, l, n)
       counter += 1
       k, 1 = maxKnotL(a)
   end
                                     #a = newA, r = egenvektorer, n = dim(a), counter = n(simTrans) og tol = max value
   return a, r, n, counter, tol
end
function rotation(a, r, k, l, n)
   if (a[k, 1] != 0.0)
       #kl = maxKnotL(a)
       tau = (a[1, 1] - a[k, k])/(2*a[k, 1]) #blir ikke dette alltid null?
       if (tau > 0)
          t = -tau + sqrt(1.0 + tau^2)
       else
           t = -tau - sqrt(1.0 + tau^2)
       end
       c = 1/sqrt(1.0 + t^2)
       s = c*t
   #end
```

```
else
      c = 1.0
       s = 0.0
   end
   a_k = a[k, k]
   a_11 = a[1, 1]
   #doing stuff with indices k and 1
   c2 = c^2
   s2 = s^2
   csak12 = 2.0*c*s*a[k, 1]
   a[k, k] = c2*a_kk - csakl2 + s2*a_l1
   a[1, 1] = s2*a_kk + csakl2 + c2*a_l1
   a[k, 1] = 0
   a[1, k] = 0
   #doing stuff with the remaing matrix elements
   for i = 1:n
       if (i != k) && (i != 1)
          a_{ik} = a[i, k]
          a_{il} = a[i, 1]
          a[i, k] = c*a_ik - s*a_il
           a[k, i] = a[i, k]
           a[i, 1] = c*a_il + s*a_ik
           a[1, i] = a[i, 1]
       end
       #calculating eigenvectors
       r_{ik} = r[i, k]
       r_{il} = r[i, 1]
       r[i, k] = c*r_ik - s*r_il
       r[i, 1] = c*r_il + s*r_ik
   end
   k, 1 = maxKnotL(a)
   #println(a[k, 1])
end
```

6.b Test functions

```
function test_maxKnotL()
   test_matrix = ones(Float64, 10, 10)
   test_matrix[1, 4] = 3
   test_matrix[4, 1] = 3
   k, 1 = maxKnotL(test_matrix)
   if test_matrix[k, 1] < 3</pre>
       error("maxKnotL returns wrong value")
   end
end
function test_eigenvalues()
   #creating testable values
   n = Int64(5)
   a = zeros(Float64, n, n)
   for i in range(1, step = 1, length = n)
       for j in range(1, step = 1, length = n)
           if ((j == i+1) || (j == i-1))
              a[i, j] = 2
                                            # b = 2
           end
           if (i == j)
                                            \# a = 5
              a[i, j] = 5
           end
       end
   test_values = [5-2*sqrt(3), 3, 5, 7, 5+2*sqrt(3)]
   #calculating values
   a_, thing1, thing2, thing3, thing4 = rotate(a, 1e-5)
   tol = 1e-3
   #sorting values into usable format
   values = zeros(5)
   for i = 1:5
       values[i] = a_[i,i]
```

```
end
values = sort(values)
#tests each element
for i = n:5
    if abs(values[i]-test_values[i]) > tol
        error("Eigentest failed: expected: $test_values, got $values")
    end
end
end
test_eigenvalues()
```

6.c Math for Quantum dots in three dimensions, one electron

We begin with

$$-\frac{\hbar^2}{2m} \left(\frac{1}{r^2} \frac{d}{dr} r^2 \frac{d}{dr} - \frac{l(l+1)}{r^2} \right) R(r) + V(r)R(r) = ER(r).$$
 (19)

Firstly we had that l = 0. We then substitued R(r) = u(r)/r, and got

$$-\frac{\hbar^2}{2m}\frac{d^2}{dr^2}u(r) + V(r)u(r) = Eu(r).$$
 (20)

In our case V(r) is equal to $(1/2)kr^2$, where $k = m\omega^2$. We now introduce a dimensionless variable, $\rho = r/\alpha$, which means $V(\rho) = (1/2)k\alpha^2\rho^2$. This gives us

$$-\frac{\hbar^2}{2m\alpha^2}\frac{d^2}{d\rho^2}u(\rho) + \frac{k}{2}\alpha^2\rho^2u(\rho) = Eu(\rho)$$
(21)

$$-\frac{d^2}{d\rho^2}u(\rho) + \frac{mk}{\hbar^2}\alpha^4\rho^2u(\rho) = \frac{2m\alpha^2}{\hbar^2}Eu(\rho).$$
 (22)

Finally we scale α so that $\frac{mk}{\hbar^2}\alpha^4 = 1$, and define $\lambda = \frac{2m\alpha^2}{\hbar^2}E$, which gives us

$$-\frac{d^2}{d\rho^2}u(\rho) + \rho^2 u(\rho) = \lambda u(\rho). \tag{23}$$

6.d Implementing a matrix in Julia

```
#function for making the matrix
function make_mat(rho_minn, rho_max, n)
   h = (rho_max-rho_minn)/n #the number of steps
   e = -1/h^2 #the non-diagonal matrix element
   d = zeros(n) #array to hold the diagonal matrix element
   #finds the diagonal matrix element for each rho
       d[r] = 2/h^2 + (rho_minn + r*h)^2
   a = zeros(Float64, n, n) #the matrix, we now have to fill in the diagonal values
   a[1,1] = 2/h^2 + rho_minn^2 #the first elements
   a[1,1+1] = e
   for i=2:n-1 #all the other elements
       a[i,i-1] = e
       a[i,i] = d[i]
       a[i,i+1] = e
   a[n,n-1] = e #the last elements
   a[n,n] = 2/h^2 + rho_max^2
   return a
end
```

6.e Math for Quantum dots in three dimensions, two electrons

$$\left(-\frac{\hbar^2}{m}\frac{d^2}{dr^2} - \frac{\hbar^2}{4m}\frac{d^2}{dR^2} + \frac{1}{4}kr^2 + kR^2\right)u(r,R) = E^{(2)}u(r,R).$$
(24)

Firstly we split u(r,R) into $\psi(r)$ and $\phi(R)$, with $E^{(2)}=E_r+E_R$. We are only interested in E_r , so we can remove everything related to R. We then add the repulsive Coulomb interaction $V(r_1,r_2)$, giving us

$$\left(-\frac{\hbar^2}{m}\frac{d^2}{dr^2} + \frac{1}{4}kr^2 + \frac{\beta e^2}{r}\right)\psi(r) = E_r\psi(r).$$
(25)

We then introduced the dimensionless vaiable $\rho = r/\alpha$, giving us

$$-\frac{d^2}{d\rho^2}\psi(\rho) + \frac{1}{4}\frac{mk}{\hbar^2}\alpha^4\rho^2\psi(\rho) + \frac{m\alpha\beta e^2}{\rho\hbar^2}\psi(\rho) = \frac{m\alpha^2}{\hbar^2}E_r\psi(\rho). \tag{26}$$

We then define $\omega_r^2 = \frac{1}{4} \frac{mk}{\hbar^2} \alpha^4$, $\frac{m\alpha\beta e^2}{\hbar^2} = 1$ and $\lambda = \frac{m\alpha^2}{\hbar^2} E$, giving us

$$-\frac{d^2}{d\rho^2}\psi(\rho) + \omega_r^2 \rho^2 \psi(\rho) + \frac{1}{\rho} = \lambda \psi(\rho). \tag{27}$$

7 REFERENCES

References

[1] Computational Physics, Lecture Notes Fall 2015, Morten Hjort-Jensen p.215-220