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Course: Numerical methods and simulations

Tutor: Bengt Eliasson E-mail: bengt.eliasson@strath.ac.uk Homepage: http://www.strath.ac.uk/physics/staff/academic/bengteliasson/

Lecture 4: Solution of nonlinear systems of equations.

 \leq We will study Newton'a method to solve nonlinear systems of equations that cannot be solved analytically.

 \leq Nonlinear equations or systems of equations frequently occur in

- Dispersion relations (to solve ω as a function of k)
- Nonlinear boundary value problems, for example Poisson's equation with Boltzmann electrons

$$\frac{d^2\phi}{dx^2} = \exp(\phi) \quad n_i(x) \tag{1}$$

with the potential given at x = 0 and x = L.

- Implicit time-stepping methods (e.g. Crank-Nicholson's scheme) where one has to solve an equation in each time-step to obtain the solution.
- Newton's method is base on Taylor expansion of the nonlinear system. Assume that we want to solve the nonlinear equation $G(v^*) = 0$ to obtain the unknown value v^* . As an initial guess we use v, and we have that $v^* = v + \delta v$ where δv is the correction of v to obtain the real solution v^* . We can then Taylor expand G around v so that $G(v^*) = G(v + \delta v) \leftarrow G(v) + \delta v G'(v)$. Since $G(v^*) = 0$ we have $\delta v \leftarrow G(v)/G'(v)$, and in the Newton algorithm we have the new value $v^{new} \mid v^{\text{old}} = G(v^{\text{old}})/G'(v^{\text{old}})$.
- Example: Solve the nonlinear equation $x = \cos(x)$ numerically. First rewrite the equation as $G(x) = x \quad \cos(x) = 0$. Then G'(x) =

 $1 + \sin(x)$, and we have the correction term $\delta x = G(x)/G'(x) = (x \cos(x))/(1 + \sin(x))$. The Newton algorithm becomes $x^{new} \mid x^{\text{old}} (x^{\text{old}} \cos(x^{\text{old}}))/(1 + \sin(x^{\text{old}}))$. If we take the initial guess x = 1, the first few iterations give

1	$\delta x =$	0.24963613215976	x = 0.75036386784024
2	$\delta x =$	0.01125097692888	x = 0.73911289091136
3	$\delta x =$	0.00002775752607	x = 0.73908513338528
4	$\delta x =$	0.0000000017012	x = 0.73908513321516

After only 4 iterations, we have a solution with 10 digits accuracy! It is typical for the Newton method that the number of accurate digits in the solution doubles in each iteration. One can show that the correction term decreases as $\delta x^{\text{new}} = C \, \delta x^{\text{old } 2}$ for most problems, where C is a constant.

In multiple dimensions, we may want to solve the nonlinear system of equations $\vec{G}(\vec{v}^*) = \vec{0}$ to obtain the vector \vec{v}^* . With the initial guess \vec{v} we have $\vec{v}^* = \vec{v} + \delta \vec{v}$ where $\delta \vec{v}$ is the correction of \vec{v} to obtain \vec{v}^* . Taylor expanding \vec{G} around \vec{v} we have $\vec{G}(\vec{v}^*) = \vec{G}(\vec{v} + \delta \vec{v}) \leftarrow \vec{G}(\vec{v}) + \mathbf{J}\delta \vec{v}$ where

$$\mathbf{J} = \begin{bmatrix} \frac{\partial G_1}{\partial v_1} & \frac{\partial G_1}{\partial v_2} & \bullet \bullet \bullet & \frac{\partial G_1}{\partial v_N} \\ \frac{\partial G_2}{\partial v_1} & \frac{\partial G_2}{\partial v_2} & & & \\ \vdots & & \ddots & \vdots \\ \frac{\partial G_N}{\partial v_1} & & \bullet \bullet & \frac{\partial G_N}{\partial v_N} \end{bmatrix}$$
(2)

is the Jacobian matrix. Since $\vec{G}(\vec{v}^*) = 0$ we can solve for $\delta \vec{v}$ to obtain $\delta \vec{v} \leftarrow \mathbf{J}^{-1} \vec{G}(\vec{v})$ and in the Newton algorithm we have the new value $\vec{v}^{new} = \vec{v} \quad \mathbf{J}^{-1} \vec{G}(\vec{v})$. The iterations are stopped when the norm of the correction term, $\delta \mathbf{v}$, is smaller than some small value ϵ . The norm is a measure of the "length" of the vector. For example the Euclidian norm (norm(x,2) in Matlab), is defined as $\vec{x}_2 = \sqrt{x_1^2 + x_2^2 + \ldots + x_N^2}$, and the max norm (norm(x,inf) in Matlab) is defined as $\delta \vec{x}_{\infty} = \max(x_1, x_2, \ldots, x_N)$.

As an example, we will now make a program to solve the system of two coupled equations

$$x = y + \cos(x + y) \tag{3}$$

$$y = x + \sin(xy) \tag{4}$$

We first rewrite the system in the form $\vec{G}(x, y) = 0$, or

$$\vec{G} = \begin{bmatrix} G_1 \\ G_2 \end{bmatrix} = \begin{bmatrix} x & y & \cos(x+y) \\ y & x & \sin(xy) \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \end{bmatrix}$$
(5)

and the Jacobian then becomes

$$\mathbf{J} = \begin{bmatrix} \partial G_1 / \partial x & \partial G_1 / \partial y \\ \partial G_2 / \partial x & \partial G_2 / \partial y \end{bmatrix} = \begin{bmatrix} 1 + \sin(x+y) & 1 + \sin(x+y) \\ 1 & y \cos(xy) & 1 & x \cos(xy) \end{bmatrix}$$
(6)

We now write the main program "main.m" for the solution of the two coupled equations. Select from the Matlab menu *New Script*, then from the menu of the new window, choose *Save / Save as*, and File Name: main.m

We now write the program in the file:

Then choose from the menu Save.

Now we write the Newton subroutine: From the Matlab menu, choose *New Script*, then choose from the menu of the new file *Save / Save as*, and File Name: Newton.m

In the new file, we write

```
% The Newton algorithm: Newton.m
% As input we have the initial guess.
function v=Newton(v)
  dv=1; % dv is the norm of the correction term.
  epsilon=1e-6; % The accuracy limit.
  counter=0;
  %Iterate the Newton algorithm until
  %the correction is smaller than epsilon.
  while(norm(dv,inf)>epsilon)
    dv=-Jacobian(v) \setminus G(v); % Solve the linear equation system J*delta=-F.
    v=v+dv; % Update the solution
    disp(['|dv|=' num2str(norm(dv,inf),'%0.10g') ' x=' ...
      num2str(v(1,1),'%0.10g') ' y=' num2str(v(2,1),'%0.10g')]);
    counter=counter+1;
    \% Check if we iterate more than 20 times, then no convergence.
    if counter>20
      disp('No convergence!')
      return;
    end
  end
```

Choose Save.

Next, we write the function that defines the system of nonlinear equations. From the Matlab menu, choose *New Script*, and then *Save / Save as* and File Name: G.m

In the new file, we write

% Definition of the nonlinear system G(x,y)=0;

```
function G=G(v)
x=v(1,1);
y=v(2,1);
% The system of nonlinearly coupled equations
G(1,1)=x-y-cos(x+y);
G(2,1)=y-x-sin(x*y);
```

Choose Save.

Finally, we write the function that defines the Jacobian for our system of nonlinear equations. From the Matlab menu, choose *New Script*, and then *Save / Save as* and File Name: Jacobian.m

In the new file, we write

% The Jacobian matrix dG/dv.

function Jacobian=Jacobian(v)

```
x=v(1,1);
y=v(2,1);
Jacobian(1,1)=1+sin(x+y); % dG1/dv1
Jacobian(1,2)=-1+sin(x+y); % dG1/dv2
Jacobian(2,1)=-1-y*cos(x*y); % dG2/dv1
Jacobian(2,2)=1-x*cos(x*y); % dG2/dv2
```

Choose Save.

We now run the program. In the Matlab window, type in main (Enter). The result is

||dv||=1.308454231 x=0.8442009784 y=2.308454231
||dv||=0.4010851826 x=0.9036509561 y=1.907369048

```
||dv||=0.0539587817 x=0.9576097378 y=1.927521852
||dv||=0.006653695218 x=0.9557371965 y=1.920868157
||dv||=3.898306889e-005 x=0.9557422362 y=1.920829174
||dv||=3.816997014e-010 x=0.9557422366 y=1.920829173
x=0.9557422366 y=1.920829173
```

We see that we have 10 significant digits after 6 iterations! Also here it is typical for the Newton method that the number of accurate digits in the solution doubles in each iteration. One can show that when $\delta \mathbf{v}$ is small enough, the correction term decreases as $\delta \mathbf{v}^{\text{new}} = C \ \delta \mathbf{v}^{\text{old } 2}$ for most problems, where C is a constant.

It is necessary to give an initial guess that is close enough to the solution that we want. For example, the initial guess x = y = 10 gives different solution from that above, and the initial guess x = 10, y = -10 leads to no convergence!

In Matlab, there also exists a command *roots* to find all roots of a polynomial. This is very useful to find solutions of dispersion relations in fluid plasmas. Example: to find the complex roots of the polynomial

$$w^4 + 2w^3 + 3w^2 + w + 4 = 0 \tag{7}$$

we do the command in Matlab

roots([1 2 3 1 4])

The result is

```
ans =
```

-1.3030 + 1.3062i -1.3030 - 1.3062i 0.3030 + 1.0409i 0.3030 - 1.0409i

i.e., the complex roots are

 $w_1 = 1.3030 + 1.3062i, w_2 = 1.3030 \quad 1.3062i, w_3 = 0.3030 + 1.0409i$ and $w_4 = 0.3030 \quad 1.0409i$ Example 2/Assignment (Nonlinear boundary value problem):

Write a program that solves the nonlinear Poisson equation

$$\frac{\partial^2 \phi}{\partial x^2} = \exp(\phi) \quad n_i(x) \tag{8}$$

where $n_i = 1 + \exp(-3(x-5)^2)$, on the domain $0 \approx x \approx L$ with the boundary conditions $\phi(0) = 0$ and $\phi(L) = 0$ with L = 10.

First we write the equation as

$$\frac{\partial^2 \phi}{\partial x^2} \quad \exp(\phi) + n_i(x) = 0. \tag{9}$$

Then discretize the space into discrete points $x = x_0, x_1, \ldots, x_{Nx}$ and represent the space in a column vector

$$\vec{x} = \begin{bmatrix} x_0 \\ x_1 \\ \vdots \\ x_{Nx} \end{bmatrix}$$
(10)

Then discretize the solution as $\phi = \phi_0, \phi_1, \ldots, \phi_{Nx}$, where $\phi_j \leftarrow \phi(x_j)$. Order the solution in a column vector as

$$\vec{\phi} = \begin{bmatrix} \phi_0 \\ \phi_1 \\ \vdots \\ \phi_{Nx} \end{bmatrix} \tag{11}$$

In order to obtain the $N_x + 1$ unknowns ϕ_j , we need $N_x + 1$ equations which we symbolically denote $\vec{G}(\vec{\phi}, x) = 0$, where the equations are ordered into a column vector

$$\vec{G} = \begin{bmatrix} G_0 \\ G_1 \\ \vdots \\ G_{Nx} \end{bmatrix}$$
(12)

We now find the equations. The left-hand boundary condition is $\phi_0 = 0$ so that $G_0 = \phi_0$, while the right-hand boundary condition is $\phi_{Nx} = 0$ so that $G_{Nx} = \phi_{Nx}$. Between the boundaries, we have the nonlinear Poisson equation, which we approximate with a centered difference method so that

$$\frac{\phi_{j-1} \quad 2\phi_j + \phi_{j+1}}{\Delta x^2} \quad \exp(\phi_j) + n_i(x_j) = 0, \quad j = 1, \dots, N_x \quad 1, (13)$$

hence

$$G_j = \frac{\phi_{j-1} - 2\phi_j + \phi_{j+1}}{\Delta x^2} \quad \exp(\phi_j) + n_i(x_j), \quad j = 1, \dots, N_x \quad 1.$$
(14)

Then the Jacobain matrix becomes

$$\mathbf{J} = \begin{bmatrix} \frac{\partial G_0}{\partial \phi_0} & \frac{\partial G_0}{\partial \phi_1} & \bullet \bullet & \frac{\partial G_0}{\partial \phi_N} \\ \frac{\partial G_1}{\partial \phi_0} & \frac{\partial G_1}{\partial \phi_1} & & \\ \vdots & \ddots & \vdots \\ \frac{\partial G_{Nx}}{\partial \phi_0} & \bullet \bullet \bullet & \frac{\partial G_{Nx}}{\partial \phi_{Nx}} \end{bmatrix}$$

$$= \begin{bmatrix} 1 & 0 & \bullet \bullet & & 0 \\ \frac{1}{\Delta x^2} & \frac{2}{\Delta x^2} & \exp(\phi_1) & \frac{1}{\Delta x^2} & \\ 0 & \frac{1}{\Delta x^2} & \frac{2}{\Delta x^2} & \exp(\phi_2) & \frac{1}{\Delta x^2} & \vdots \\ \vdots & \ddots & \ddots & \ddots & 0 \\ 0 & \frac{1}{\Delta x^2} & \frac{2}{\Delta x^2} & \exp(\phi_{Nx-1}) & \frac{1}{\Delta x^2} \\ 0 & \bullet \bullet \bullet & & 0 \end{bmatrix}$$
(15)

We now write the program to solve the nonlinear system of equations. Fitst the main program, *main.m*:

% Main program to solve a boundary value problem: main.m clear

```
L=10;
Nx=100;
dx=L/Nx;
x=(0:Nx)'*dx;
ni=1+exp(-3*(x-5).^2);
% Initial guess
phi=log(ni);
% Solve with the Newton algorithm
phi=Newton(phi,x,ni,dx,Nx);
%Plot the result
subplot(3,1,1) plot(x,phi) title('\phi') subplot(3,1,2)
plot(x,exp(phi)) title('n_e') subplot(3,1,3) plot(x,ni) title('n_i')
```

```
while(norm(dphi,inf)>epsilon)
% Solve the linear equation system J*delta=-F.
```

```
dphi=-Jacobian(phi,x,ni,dx,Nx)\G(phi,x,ni,dx,Nx);
```

phi=phi+dphi; % Update the solution

disp(['||dphi||=' num2str(norm(dphi,inf),'%0.10g')]);

counter=counter+1;

```
% Check if we iterate more than 20 times, then no convergence.
if counter>20
  disp('No convergence!')
  return;
end
```

end

The system of equations \vec{G} is calculated in the file G.m:

% Definition of the nonlinear system: G.m

function G=G(phi,x,ni,dx,Nx)

```
% The system of nonlinearly coupled equations
G(1,1)=phi(1,1);
G(2:Nx,1)=(phi(1:Nx-1,1)-2*phi(2:Nx,1)+phi(3:Nx+1,1))/dx<sup>2</sup> ...
-exp(phi(2:Nx,1))+ni(2:Nx,1);
G(Nx+1,1)=phi(Nx+1,1);
```

Finally, the Jacobian matrix is calculated in the file Jacobian.m

% The Jacobian matrix dG/dphi: Jacobian.m

function Jacobian=Jacobian(phi,x,ni,dx,Nx)

Jacobian=sparse(0); % Store only non-zero elements

```
Jacobian(1,1)=1;
for j=2:Nx
    Jacobian(j,j-1)=1/dx^2;
    Jacobian(j,j)=-2/dx^2-exp(phi(j,1));
    Jacobian(j,j+1)=1/dx^2;
end
Jacobian(Nx+1,Nx+1)=1;
```

Now try to run the program by the command **main** in matlab. The printout on screen shows the norm of the correction term, which is a measure of the convergence of the method.

Printout:

||dphi||=0.3246604088
||dphi||=0.02132031524
||dphi||=0.0001206819599
||dphi||=4.499908369e-009

After 4 iterations, we have the solution with about 10 digits accuracy. The plot of the solution is

