Scientific Computing: Solving Nonlinear Equations

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Fundamentals

• Simplest problem: Root finding in one dimension:

$$f(x) = 0$$
 with $x \in [a, b]$

• Or more generally, solving a square system of nonlinear equations

$$\mathbf{f}(\mathbf{x}) = \mathbf{0} \quad \Rightarrow f_i(x_1, x_2, \dots, x_n) = 0 \text{ for } i = 1, \dots, n.$$

- There can be no closed-form answer, so just as for eigenvalues, we need iterative methods.
- Most generally, starting from m ≥ 1 initial guesses x⁰, x¹,..., x^m, iterate:

$$x^{k+1} = \phi(x^k, x^{k-1}, \dots, x^{k-m}).$$

Order of convergence

- Consider one dimensional root finding and let the actual root be α , $f(\alpha) = 0$.
- A sequence of iterates x^k that converges to α has order of convergence p ≥ 1 if as k → ∞

$$\frac{\left|x^{k+1}-\alpha\right|}{\left|x^{k}-\alpha\right|^{p}} = \frac{\left|e^{k+1}\right|}{\left|e^{k}\right|^{p}} \to C = \text{const},$$

where the constant C is a **convergence factor**, C < 1 for p = 1.

- A method should at least converge **linearly** (p = 1), that is, the error should at least be reduced by a constant factor every iteration, for example, the number of accurate digits increases by 1 every iteration.
- A good method for root finding coverges **quadratically** (*p* = 2), that is, the number of accurate digits **doubles** every iteration!

Local vs. global convergence

- A good initial guess is extremely important in nonlinear solvers!
- Assume we are looking for a unique root a ≤ α ≤ b starting with an initial guess a ≤ x₀ ≤ b.
- A method has local convergence if it converges to a given root α for any initial guess that is sufficiently close to α (in the neighborhood of a root).
- A method has **global convergence** if it converges to the root for any initial guess.
- General rule: Global convergence requires a **slower** (careful) method **but is safer**.
- It is best to combine a global method to first find a good initial guess close to α and then use a faster local method.

Basics of Nonlinear Solvers

Conditioning of root finding

$$f(\alpha + \delta \alpha) \approx f(\alpha) + f'(\alpha)\delta \alpha = \delta f$$

$$|\delta \alpha| \approx \frac{|\delta f|}{|f'(\alpha)|} \quad \Rightarrow \kappa_{abs} = |f'(\alpha)|^{-1}.$$

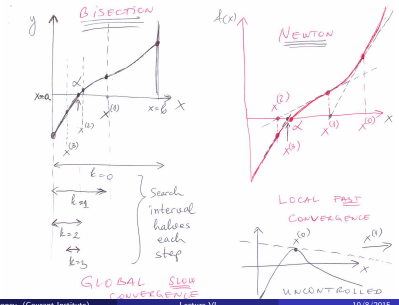
- The problem of finding a simple root is well-conditioned when |f'(α)| is far from zero.
- Finding roots with multiplicity m > 1 is ill-conditioned:

$$|f'(\alpha)| = \cdots = |f^{(m-1)}(\alpha)| = 0 \quad \Rightarrow \quad |\delta\alpha| \approx \left[\frac{|\delta f|}{|f^m(\alpha)|}\right]^{1/m}$$

• Note that finding **roots of algebraic equations** (polynomials) is a separate subject of its own that we skip.

One Dimensional Root Finding

The bisection and Newton algorithms



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Bisection

• First step is to **locate a root** by searching for a **sign change**, i.e., finding a^0 and b^0 such that

 $f(a^0)f(b^0) < 0.$

• The simply **bisect** the interval, for k = 0, 1, ...

$$x^k = \frac{a^k + b^k}{2}$$

and choose the half in which the function changes sign, i.e., either $a^{k+1} = x^k$, $b^{k+1} = b^k$ or $b^{k+1} = x^k$, $a^{k+1} = a^k$ so that $f(a^{k+1})f(b^{k+1}) < 0$.

- Observe that each step we need one function evaluation, $f(x^k)$, but only the sign matters.
- The convergence is essentially linear because

$$|x^k - \alpha| \le \frac{b^k}{2^{k+1}} \quad \Rightarrow \frac{|x^{k+1} - \alpha|}{|x^k - \alpha|} \le 2.$$

Newton's Method

- Bisection is a slow but sure method. It uses no information about the value of the function or its derivatives.
- Better convergence, of order $p = (1 + \sqrt{5})/2 \approx 1.63$ (the golden ratio), can be achieved by using the value of the function at two points, as in the **secant method**.
- Achieving second-order convergence requires also evaluating the **function derivative**.
- Linearize the function around the current guess using Taylor series:

$$f(x^{k+1}) \approx f(x^k) + (x^{k+1} - x^k)f'(x^k) = 0$$

$$x^{k+1} = x^k - \frac{f(x^k)}{f'(x^k)}$$

One Dimensional Root Finding

Convergence of Newton's method

Use Taylor series with remainder and divide by $f'(x^k) \neq 0$:

$$\exists \xi \in [x_n, \alpha]: \quad f(\alpha) = 0 = f(x^k) + (\alpha - x^k)f'(x^k) + \frac{1}{2}(\alpha - x^k)^2 f''(\xi) = 0,$$

$$\begin{bmatrix} x^{k} - \frac{f(x^{k})}{f'(x^{k})} \end{bmatrix} - \alpha = -\frac{1}{2}(\alpha - x^{k})^{2} \frac{f''(\xi)}{f'(x^{k})}$$
$$x^{k+1} - \alpha = e^{k+1} = -\frac{1}{2} \left(e^{k}\right)^{2} \frac{f''(\xi)}{f'(x^{k})}$$

which shows second-order convergence

$$\frac{\left|x^{k+1}-\alpha\right|}{\left|x^{k}-\alpha\right|^{2}} = \frac{\left|e^{k+1}\right|}{\left|e^{k}\right|^{2}} = \left|\frac{f''(\xi)}{2f'(x^{k})}\right| \to \left|\frac{f''(\alpha)}{2f'(\alpha)}\right|$$

Newton's method **converges quadratically if we start sufficiently close to a simple root**, more precisely, if

$$|\mathbf{x}^{0} - \alpha| = |\mathbf{e}^{0}| \lesssim \left|\frac{2f'(\alpha)}{f''(\alpha)}\right|$$

Fixed-Point Iteration

 Another way to devise iterative root finding is to rewrite f(x) = 0 in an equivalent form

$$x = \phi(x)$$

• Then we can use fixed-point iteration

$$x^{k+1} = \phi(x^k)$$

whose fixed point (limit), if it converges, is $x \to \alpha$.

It can be proven that the fixed-point iteration x^{k+1} = φ(x^k) converges if φ(x) is a contraction mapping:

$$|\phi'(x)| \leq K < 1 \quad \forall x \in [a, b]$$

Stopping Criteria

- A good library function for root finding has to implement careful termination criteria.
- An obvious option is to terminate when the residual becomes small

$$\left|f(x^{k})\right| < \varepsilon,$$

which is only good for very well-conditioned problems, $|f'(\alpha)| \sim 1$.

• Another option is to terminate when the increment becomes small

$$\left|x^{k+1}-x^k\right|<\varepsilon.$$

For fixed-point iteration

$$x^{k+1} - x^k = e^{k+1} - e^k pprox \left[1 - \phi'(lpha)
ight] e^k \quad \Rightarrow \quad \left|e^k\right| pprox rac{arepsilon}{\left[1 - \phi'(lpha)
ight]},$$

so we see that the increment test works for rapidly converging iterations ($\phi'(\alpha) \ll 1$).

In practice

- A robust but fast algorithm for root finding would **combine bisection** with Newton's method.
- Specifically, a method like Newton's that can easily take huge steps in the wrong direction and lead far from the current point must be **safeguarded** by a method that ensures one does not leave the search interval and that the zero is not missed.
- Once x^k is close to α, the safeguard will not be used and quadratic or faster convergence will be achieved.
- Newton's method requires first-order derivatives so often other methods are preferred that require **function evaluation only**.
- Matlab's function *fzero* combines bisection, secant and inverse quadratic interpolation and is "fail-safe".

Find zeros of $a\sin(x) + b\exp(-x^2/2)$ in MATLAB

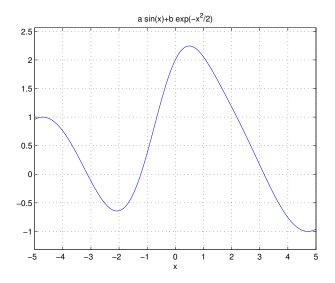
% f=0mfile uses a function in an m-file

% Parameterized functions are created with:

$$a = 1; b = 2;$$

 $f = @(x) a*sin(x) + b*exp(-x^2/2) ; % Handle$
figure(1)
 $ezplot(f, [-5, 5]);$ grid
 $x1=fzero(f, [-2, 0])$
 $[x2, f2]=fzero(f, 2.0)$
 $x1 = -1.227430849357917$
 $x2 = 3.155366415494801$
 $f2 = -2.116362640691705e-16$

Figure of f(x)



Multi-Variable Taylor Expansion

- It is convenient to focus on one of the equations, i.e., consider a scalar function f(x).
- The usual Taylor series is replaced by

$$f(\mathbf{x} + \Delta \mathbf{x}) = f(\mathbf{x}) + \mathbf{g}^{T} (\Delta \mathbf{x}) + \frac{1}{2} (\Delta \mathbf{x})^{T} \mathbf{H} (\Delta \mathbf{x})$$

where the gradient vector is

$$\mathbf{g} = \boldsymbol{\nabla}_{\mathbf{x}} f = \left[\frac{\partial f}{\partial x_1}, \frac{\partial f}{\partial x_2}, \cdots, \frac{\partial f}{\partial x_n}\right]^T$$

and the Hessian matrix is

$$\mathbf{H} = \boldsymbol{\nabla}_{\mathbf{x}}^2 f = \left\{ \frac{\partial^2 f}{\partial x_i \partial x_j} \right\}_{ij}$$

Vector Functions of Vectors

• We are after solving a square system of nonlinear equations for some variables x:

$$\mathbf{f}(\mathbf{x}) = \mathbf{0} \quad \Rightarrow f_i(x_1, x_2, \dots, x_n) = 0 \text{ for } i = 1, \dots, n.$$

• The first-order Taylor series is

$$\mathbf{f}\left(\mathbf{x}^{k}+\Delta\mathbf{x}\right)\approx\mathbf{f}\left(\mathbf{x}^{k}\right)+\left[\mathbf{J}\left(\mathbf{x}^{k}\right)\right]\Delta\mathbf{x}=\mathbf{0}$$

where the Jacobian **J** has the gradients of $f_i(\mathbf{x})$ as rows:

$$\left[\mathbf{J}\left(\mathbf{x}\right)\right]_{ij} = \frac{\partial f_i}{\partial x_j}$$

Newton's Method for Systems of Equations

- It is much harder if not impossible to do globally convergent methods like bisection in higher dimensions!
- A good initial guess is therefore a must when solving systems, and Newton's method can be used to refine the guess.
- The basic idea behind Newton's method is to **linearize** the equation around the current guess:

$$\mathbf{f}\left(\mathbf{x}^{k}+\Delta\mathbf{x}\right)\approx\mathbf{f}\left(\mathbf{x}^{k}\right)+\left[\mathbf{J}\left(\mathbf{x}^{k}\right)\right]\Delta\mathbf{x}=\mathbf{0}$$

$$\left[{{\mathbf{J}}\left({{\mathbf{x}}^k}
ight)}
ight]\Delta {\mathbf{x}} = - {\mathbf{f}}\left({{\mathbf{x}}^k}
ight)$$
 but denote ${\mathbf{J}} \equiv {\mathbf{J}}\left({{\mathbf{x}}^k}
ight)$

$$\mathbf{x}^{k+1} = \mathbf{x}^k + \Delta \mathbf{x} = \mathbf{x}^k - \mathbf{J}^{-1} \mathbf{f} \left(\mathbf{x}^k \right).$$

• Newton's method converges **quadratically** if started sufficiently close to a root **x***at which the **Jacobian is not singular**.

$$\|\mathbf{e}^{k+1}\| \le \frac{\|\mathbf{J}^{-1}\| \|\mathbf{H}\|}{2} \|\mathbf{e}^{k}\|^{2}$$

- Newton's method requires solving **many linear systems**, which can become complicated when there are many variables.
- It also requires computing a whole **matrix of derivatives**, which can be expensive or hard to do (differentiation by hand?)
- Newton's method converges fast if the Jacobian **J**(**x**^{*}) is well-conditioned, otherwise it can "blow up".
- For large systems one can use so called **quasi-Newton** methods:
 - Approximate the Jacobian with another matrix \widetilde{J} and solve $\widetilde{J}\Delta x=f(x^k).$
 - Damp the step by a step length $\alpha_k \lesssim 1$,

$$\mathbf{x}^{k+1} = \mathbf{x}^k + \alpha_k \Delta \mathbf{x}.$$

• Update \widetilde{J} by a simple update, e.g., a rank-1 update (recall homework 2).

Continuation methods

- To get a good initial guess for Newton's method and ensure that it converges fast we can use **continuation methods** (also called **homotopy methods**).
- The basic idea is to solve

$$\tilde{\mathbf{f}}_{\lambda}\left(\mathbf{x}\right) = \lambda \mathbf{f}\left(\mathbf{x}\right) + (1 - \lambda) \mathbf{f}_{\mathbf{a}}\left(\mathbf{x}\right) = \mathbf{0}$$

instead of the original equations, where $0 \le \lambda \le 1$ is a parameter.

- If λ = 1, we are solving the original equation f (x) = 0, which is hard because we do not have a good guess for the initial solution.
- If $\lambda = 0$, we are solving $\mathbf{f}_{a}(\mathbf{x}) = \mathbf{0}$, and we will assume that this is easy to solve. For example, consider making this a linear function,

$$f_{a}\left(x\right) =x-a,$$

where **a** is a vector of parameters that need to be chosen somehow. One can also take a more general $f_a\left(x\right) = Ax - a$ where A is a matrix of parameters, so that solving $f_a\left(x\right) = 0$ amounts to a linear solve which we know how to do already.

Path Following

- The basic idea of continuation methods is to start with $\lambda = 0$, and solve $\tilde{\mathbf{f}}_{\lambda}(\mathbf{x}) = 0$. This gives us a solution \mathbf{x}_{0} .
- Then increment λ by a little bit, say λ = 0.05, and solve f
 _λ (x) using Newton's method starting with x₀ as an initial guess.
 Observe that this is a good initial guess under the assumption that the solution has not changed much because λ has not changed much.
- We can repeat this process until we reach $\lambda = 1$, when we get the actual solution we are after:
 - Choose a sequence $\lambda_0 = 0 < \lambda_1 < \lambda_2 < \cdots < \lambda_{n-1} < \lambda_n = 1$.
 - For k = 0 solve $\mathbf{f}_{\mathbf{a}}(\mathbf{x}_0) = \mathbf{0}$ to get \mathbf{x}_0 .
 - For k = 1, ..., n, solve a nonlinear system to get \mathbf{x}_k ,

$$\widetilde{\mathbf{f}}_{\lambda_{k}}\left(\mathbf{x}_{k}
ight)=\mathbf{0}$$

using Newton's method starting from x_{k-1} as an initial guess.

Path Following

- Observe that if we change λ very slowly we have hope that the solution will trace a **continuous path of solutions**.
- That is, we can think of $\mathbf{x}(\lambda)$ as a continuous function defined on [0, 1], defined implicitly via

$$\lambda \mathbf{f} (\mathbf{x} (\lambda)) + (1 - \lambda) \mathbf{f}_{\mathbf{a}} (\mathbf{x} (\lambda)) = \mathbf{0}.$$

- This rests on the assumption that this path will not have turning points, bifurcate or wonder to infinity, and that there is a solution for every λ.
- It turns out that by a judicious choice of f_a one can insure this is the case. For example, choosing a random a and taking $f_a(x) = x a$ works.
- The trick now becomes how to choose the sequence λ_k to make sure λ changes not too much but also not too little (i.e., not too slowly), see HOMPACK library for an example.

In practice

- It is much harder to construct general robust solvers in higher dimensions and some **problem-specific knowledge** is required.
- There is no built-in function for solving nonlinear systems in MATLAB, but the **Optimization Toolbox** has *fsolve*.
- In many practical situations there is some continuity of the problem so that a previous solution can be used as an initial guess.
- For example, **implicit methods for differential equations** have a time-dependent Jacobian **J**(*t*) and in many cases the solution **x**(*t*) evolves smootly in time.
- For large problems specialized sparse-matrix solvers need to be used.
- In many cases derivatives are not provided but there are some techniques for **automatic differentiation**.

Conclusions/Summary

- Root finding is well-conditioned for **simple roots** (unit multiplicity), ill-conditioned otherwise.
- Methods for solving nonlinear equations are always iterative and the order of convergence matters: second order is usually good enough.
- A good method uses a higher-order unsafe method such as **Newton method** near the root, but **safeguards** it with something like the **bisection** method.
- Newton's method is second-order but requires derivative/Jacobian evaluation. In **higher dimensions** having a **good initial guess** for Newton's method becomes very important.
- **Quasi-Newton** methods can aleviate the complexity of solving the Jacobian linear system.