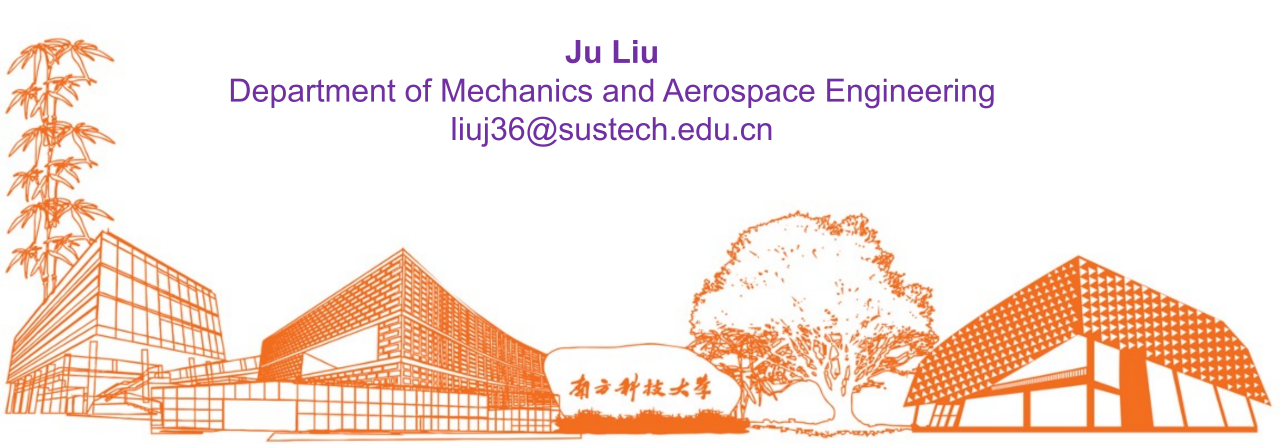
MAE 5032 High Performance Computing: Methods and Practices

Lecture 12: Numerical Analysis Basics



Floating-point arithmetic

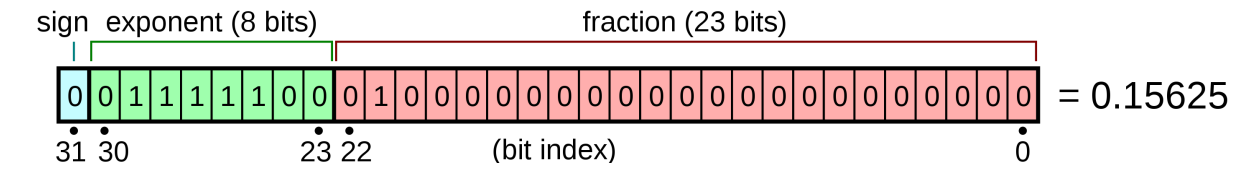
 Computers use a finite number of bits to represent numbers. Thus only a finite number of numbers can be represented.

$$x = \pm \sum_{i=0}^{t-1} d_i \beta^{-i} \beta^e = (1 \times 2^{-0} + 1 \times 2^{-1} + 0 \times 2^{-2} + \dots \times 2^{-23}) \times 2^1$$

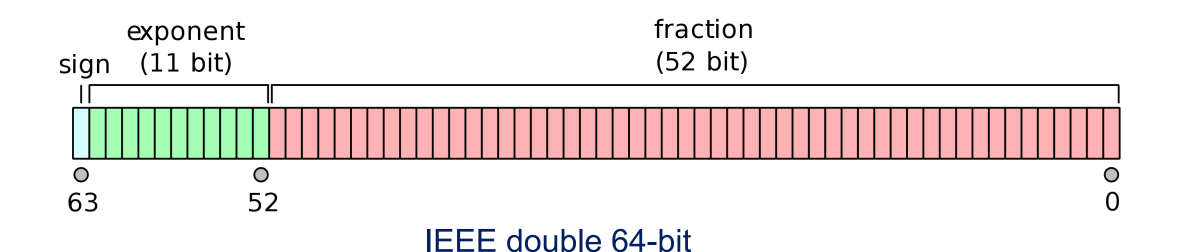
$$\approx 1.5707964 \times 2 = 3.1415928$$

- One bit for sign (unsigned number exisits)
- β is the base of the number system (2,10, 16, etc.)
- t is the significand precision
- $0 \le d_i \le \beta 1$ is the digits of the mantissa
- $L \le e \le U$ is the signed exponent

β	t	L	U
2	24	-126	127
2	53	-1022	1023
2	48	-16383	16384
16	6	-64	63
10	50	-999	999
	2 2 2 16	2 24 2 53 2 48 16 6	2 24 -126 2 53 -1022 2 48 -16383 16 6 -64



IEEE single 32-bit



- Normalized number require that the first digit in the mantissa to be nonzero.
- In binary system, the nonzero number is 1, thus we get one free bit for mantissa.

$$x = \pm \sum_{i=0}^{t} d_i \beta^{-i} \beta^e = (1 \times 2^{-0} + 1 \times 2^{-1} + 0 \times 2^{-2} + \dots \times 2^{-23}) \times 2^{1}$$

$$\approx 1.5707964 \times 2 = 3.1415928$$

- One bit for sign (unsigned number exisits)
- β is the base of the number system (2,10, 16, etc.)
- t is the significand precision
- $0 \le d_i \le \beta 1$ is the digits of the mantissa
- $L \le e \le U$ is the signed exponent

β	t	L	U
2	24	-126	127
2	53	-1022	1023
2	48	-16383	16384
16	6	-64	63
10	50	-999	999
	2 2 2 16	2 24 2 53 2 48 16 6	2 24 -126 2 53 -1022 2 48 -16383 16 6 -64

- Normalized number require that the first digit in the mantissa to be nonzero.
- In binary system, the nonzero number is 1, thus we get one free bit for mantissa.

$$x = \pm \sum_{i=0}^{t-1} d_i \beta^{-i} \beta^e = (1 \times 2^{-0} + 1 \times 2^{-1} + 0 \times 2^{-2} + \dots \times 2^{-23}) \times 2^1$$

$$\approx 1.5707964 \times 2 = 3.1415928$$

- Underflow level β^L
- Overflow level $(1 \beta^{-t})\beta^{U+1}$

	β	t	L	U
IEEE single (32 bit)	2	24	-126	127
IEEE double (64 bit)	2	53	-1022	1023
Old Cray 64bit	2	48	-16383	16384
IBM mainframe 32 bit	16	6	-64	63
packed decimal	10	50	-999	999

Representation error

- Error between a number x and its floating-point representation \tilde{x} :
 - \triangleright absolute $|x \tilde{x}|$
 - > relative $\frac{|x-\tilde{x}|}{|x|}$
- Equivalently, sometimes we say $\tilde{x} = x(1 \pm \varepsilon)$
- IEEE 754 standard gives five rounding rules, two of them are
 - truncation: $\varepsilon = \beta^{-t}$
 - rounding: $\varepsilon = 0.5 \, \beta^{-t}$
- The maximum relative error is called the machine precision. (Calculate it for a 64-bit double.)

Example: decimal numerical system (i.e. β =10), t = 3, x = 0.1256, then \tilde{x}_r = 0.126 or \tilde{x}_t = 0.125.

Addition

- Steps for addition of floating-point numbers
 - align exponents
 - add mantissas
 - adjust exponent to normalize the result

```
Example: 123456.7 + 101.7654 = (1.234567 \times 10^5) + (1.017654 \times 10^2)
= (1.234567 \times 10^5) + (0.001018 \times 10^5)
= 1.235585 \times 10^5
```

• We consider x_i and their floating-point number $\tilde{x}_i = x_i(1 + \varepsilon_i)$ To compute $s = x_1 + x_2$, the sum is represented as $\tilde{s} = (\tilde{x}_1 + \tilde{x}_2) (1 + \varepsilon_3) = x_1(1 + \varepsilon_1) (1 + \varepsilon_3) + x_2(1 + \varepsilon_2) (1 + \varepsilon_3)$ $\approx x_1(1 + \varepsilon_1 + \varepsilon_3) + x_2(1 + \varepsilon_2 + \varepsilon_3) \approx s(1 + 2\varepsilon)$

Conclusion: Errors are added

Subtraction and associativity

Example: $123457.1467 - 123456.659 \approx (1.234571 \times 10^5) - (1.234567 \times 10^5) = 4.000000 \times 10^{-1}$

- The actual result is 4.877000×10^{-1}
- Relative error of the subtraction is about 20%.
- In extreme cases, all significant numbers can be lost due to cancellation.

Example: 7-digits decimal floating-point number to calculate (a+b)+c and a+(b+c), with a=1234.567, b=45.67834, c=0.0004.

a+b = 1280.24534, rounds to 1280.245; (a+b)+c = 1280.2454 rounds to 1280.245

(b+c) = 45.67874 rounds to 45.67874; a+(b+c) = 1280.24574 rounds to 1280.246

Because of this, compilers will not automatically reorder the operations for FP.

You need to enable "fast-math" options.

A toy problem

Evaluate $\sum_{n=1}^{10000} \frac{1}{n^2}$. The precise value is 1.644834 in a decimal numerical system (i.e. β =10) with t=7.

First term is 1.000000, so the partial sum will be greater than 1. So, for the terms that $\frac{1}{n^2} < 10^{-6}$, their contribution to the sum will be ignored.

Floating point sum is 1.644725: 4 correct digits.

Solution: sum in reverse order.

Unstable algorithm

Consider the recurrence

$$y_n = \int_0^1 \frac{x^n}{x+5} dx$$
. We may deduce that $y_n = \frac{1}{n} - 5y_{n-1}$.

The initial value of the recurrence is $y_0 = \log(6) - \log(5) = 1.82|322 \times 10^{-1}$. Consider a decimal numerical system (i.e. β =10) with t=3.

$$\begin{array}{lll} \text{FP arithmetic} & \text{Correct value} \\ \tilde{y}_0 = 1.82 \times 10^{-1} & y_0 = 1.82 \times 10^{-1} \\ \tilde{y}_1 = 9.00 \times 10^{-2} & y_1 = 8.84 \times 10^{-2} \\ \tilde{y}_2 = 5.00 \times 10^{-2} & y_2 = 5.80 \times 10^{-2} \\ \tilde{y}_3 = 8.30 \times 10^{-2} & y_3 = 4.31 \times 10^{-2} \\ \tilde{y}_4 = -1.65 \times 10^{-1} & y_4 = 3.43 \times 10^{-2} \end{array}$$

Let
$$\tilde{y}_n = y_n + \varepsilon_n$$
. Then $\tilde{y}_n = \frac{1}{n} - 5 \tilde{y}_{n-1} = \frac{1}{n} - 5 y_{n-1} + 5 \varepsilon_{n-1} = y_n + 5 \varepsilon_{n-1}$. $\varepsilon_n = 5 \varepsilon_{n-1}$. Error grows exponentially.

Summary

- Arithmetic on computer is done based on floating-point numbers, and thus simple calculations like addition may lead to error.
- Mathematically equivalent operations may not remain equivalent on computers, which could affect parallel computations.
- Algorithms need to be carefully designed to control the floating-point error
- Stability analysis is needed for ALL operations on computers.

Reference: Introudction to High Performance Scientific Computing by Victor Eijkhout, Chapter 2.

Linear Algebra

A known unusable algorithm

To solve Ax = b, one may think of the Cramer's rule:

$$x_{i} = \begin{vmatrix} a_{11} & a_{12} & \dots & a_{1i-1} & b_{1} & a_{1i+1} & \dots & a_{1n} \\ a_{21} & \dots & b_{2} & \dots & a_{2n} \\ \vdots & \vdots & & \vdots & & \vdots \\ a_{n1} & \dots & b_{n} & \dots & a_{nn} \end{vmatrix} / |A|$$

The time complexity is O(n!)

Recall that on a single CPU of TaiYi, we can achieve around 76.8 Gflops.

This algorithm is too expensive! Solving a 20-by-20 matrix problem may take 1 year!

Direct method

To solve Ax = b, one may do the old Gaussian elimination method:

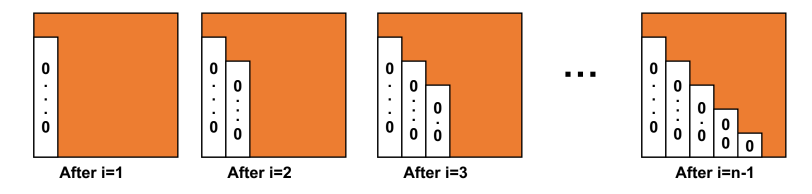
$$\begin{pmatrix} 6 & -2 & 2 \\ 12 & -8 & 6 \\ 3 & -13 & 3 \end{pmatrix} x = \begin{pmatrix} 16 \\ 26 \\ -19 \end{pmatrix}$$

$$\begin{bmatrix} 6 & -2 & 2 & | & 16 \\ 12 & -8 & 6 & | & 26 \\ 3 & -13 & 3 & | & -19 \end{bmatrix} \longrightarrow \begin{bmatrix} 6 & -2 & 2 & | & 16 \\ 0 & -4 & 2 & | & -6 \\ 0 & -12 & 2 & | & -27 \end{bmatrix} \longrightarrow \begin{bmatrix} 6 & -2 & 2 & | & 16 \\ 0 & -4 & 2 & | & -6 \\ 0 & 0 & -4 & | & -9 \end{bmatrix}$$

Algorithmic complexity is $O(n^3)$.

Gaussian Elimination algorithm

- Add multiples of each row to later rows to make A upper triangular
- Solve resulting triangular system Ux = c by substitution



Gaussian Elimination algorithm: math kernel

- LU factorization: if the above algorithm completes, we get A = LU. $\square = \triangle^{*}$
- Thus Ax=b becomes LUx=b.
 - ➤ We solve Ly=b first and solve Ux=y next.
- Solving Ax=b using GE:
 - Factorize A = L*U using GE $(\cos t = 2/3 \text{ n}^3 \text{ flops})$
 - \triangleright Solve L*y = b for y, using substitution (cost = n^2 flops)
 - \triangleright Solve U*x = y for x, using substitution (cost = n^2 flops)

Roundoff control

Consider a system

$$\begin{bmatrix} \varepsilon & 1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 + \varepsilon \\ 2 \end{bmatrix}.$$

The exact solution is

$$\begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 \\ 1 \end{bmatrix}.$$

Perform Gaussian elimination

$$\begin{bmatrix} \varepsilon & 1 \\ 0 & 1 - \frac{1}{\varepsilon} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 + \varepsilon \\ 1 - \frac{1}{\varepsilon} \end{bmatrix}.$$

We can do a "back-substitution" by solving x_2 first and solve x_1 next:

$$x_2 = 1 \Longrightarrow x_1 = 1$$
.

Roundoff control (cont.)

Suppose ε is smaller than the machine precision. $1 - \frac{1}{\varepsilon}$ becomes $-\frac{1}{\varepsilon}$, and $1 + \varepsilon$ becomes 1.

The Gaussian elimination

$$\begin{bmatrix} \varepsilon & 1 \\ 0 & 1 - \frac{1}{\varepsilon} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 + \varepsilon \\ 1 \\ 1 - \frac{\varepsilon}{\varepsilon} \end{bmatrix}$$

becomes

$$\begin{bmatrix} \varepsilon & 1 \\ 0 & -\frac{1}{\varepsilon} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 \\ -\frac{1}{\varepsilon} \end{bmatrix}$$

We can do a "back-substitution" by solving x_2 first and solve x_1 next: $x_2 = 1 \Rightarrow x_1 = 0$.

Machine round-off error has a dramatic impact on the results. We call this numerical instability.

Pivoting in Gaussian Elimination

- During the LU factorization, when the diagonal is small, swap the row (or column) to avoid division by small numbers.
 - This is referred to as pivoting in GE
 - We choose the largest possible pivot.
 - > In fact, we always choose the largest possible value as the pivot in GE.
 - > There is always a nonzero pivot if the matrix is non-singular.

Roundoff control with pivoting

Consider the system again,

$$\begin{bmatrix} \varepsilon & 1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 1 + \varepsilon \\ 2 \end{bmatrix}.$$

Pivot the row

$$\begin{bmatrix} 1 & 1 \\ \varepsilon & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 + \varepsilon \end{bmatrix}.$$

Perform Gaussian elimination

$$\begin{bmatrix} 1 & 1 \\ 0 & 1 - \varepsilon \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 - \varepsilon \end{bmatrix}.$$

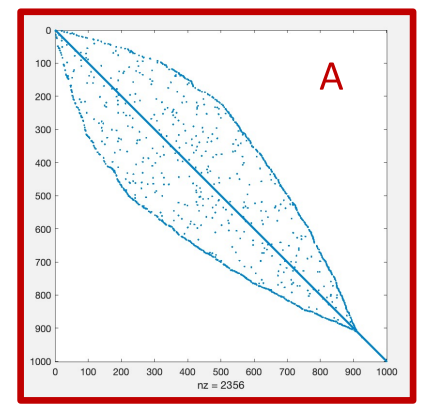
If ε is small,

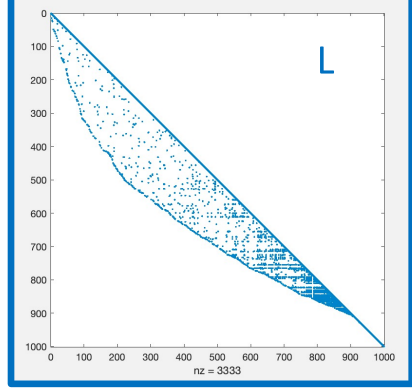
$$\begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 \end{bmatrix}.$$

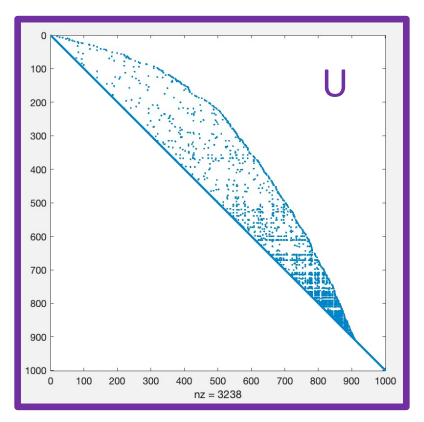
Back substitution: $x_2 = 1 \implies x_1 = 1$.

The fill-in phenomenon

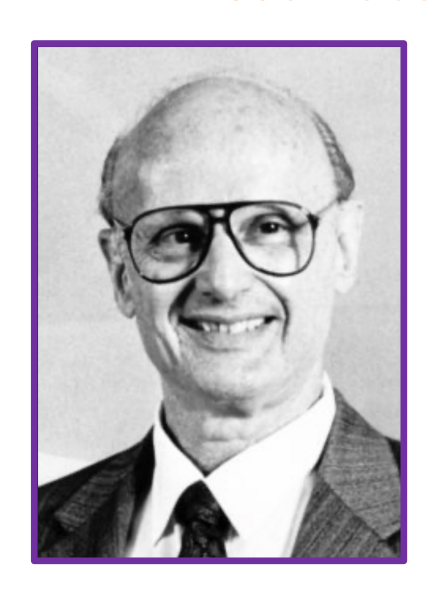
- LU factorization of a sparse matrix typically leads to additional nonzeros. This phenomenon is called **fill-in**.
- The memory requirement can thus become a bottleneck if one wants to use direct method to solve a sparse matrix.







1990 Nobel Prize in Economics



Our models strained the computer capabilities of the day [1950s]. I observed that most of the coefficients in our matrices were zero; i.e., the nonzeros were sparse in the matrix.

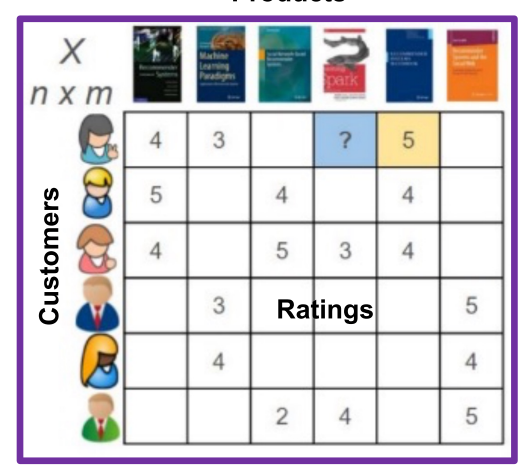
-- Harry Markowitz

Sparse matrices arise in many applications:

- simulating climate
- analyzing images
- web page ranking in search engines
- ·

Examples

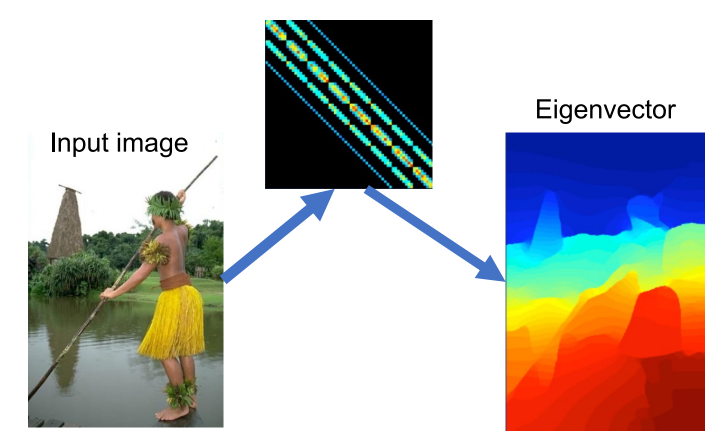
Products



Recommendation Matrix

Image segmentation – identify the object boundaries in an image.

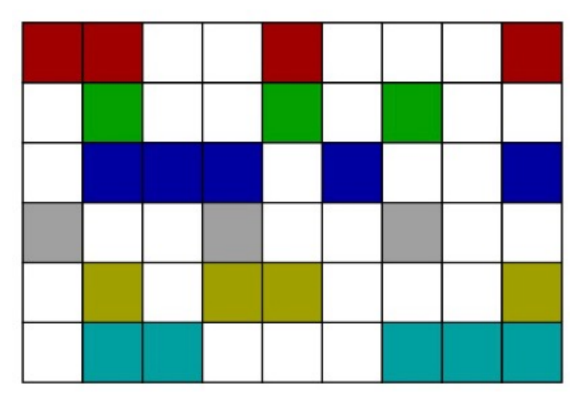
Find the eigenvalue of the affinity matrix. Efficient, High-quality image contour detection by Catanzaro, Su, et al. 2009.

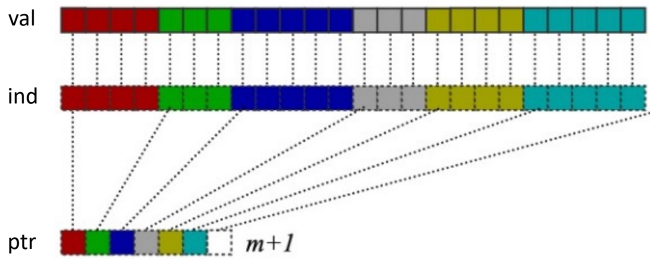


Data structure

- Use a proper data structure to represent the matrix by saving the nonzeros only.
- Compressed Row Storage (CRS) is the most widely used format.

Example:





```
y(i) ← y(i) + A(i,j) x(j)
for each row I
    for k = ptr[i] to ptr[i+1]
        y[i] += val[k] * x[ind[k]]
```

Data structure

- Use a proper data structure to represent the matrix by saving the nonzeros only.
- Compressed Row Storage (CRS) is the most widely used format.

Example:

$$A = \begin{pmatrix} 10 & 0 & 0 & 0 & -2 & 0 \\ 3 & 9 & 0 & 0 & 0 & 3 \\ 0 & 7 & 8 & 7 & 0 & 0 \\ 3 & 0 & 8 & 7 & 5 & 0 \\ 0 & 8 & 0 & 9 & 9 & 13 \\ 0 & 4 & 0 & 0 & 2 & -1 \end{pmatrix} \frac{\overline{\text{val}}}{\underline{\text{ind}}}$$

val	10	-2	3	9	3	7	8	7	3 9	13	4	2	-1
ind	0	4	0	1	5	1	2	3	$0 \cdots 4$	5	1	4	5
ptr	0	2	5	8	12	16	19						

Preallocation of the memory space can be critical! Otherwise, the initial use of the matrix can be extremely slow.

Stationary iterative method

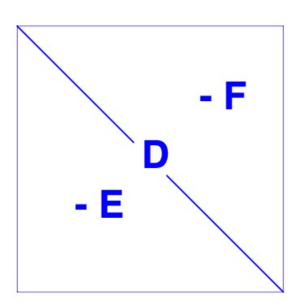
• To solve Ax = b with the matrix A being a sparse matrix, we want to devise an iterative method

$$Mx_{k+1} = Nx_k + f.$$
- h also satisfies $Mx - Nx + f$

such that the solution of Ax = b also satisfies Mx = Nx + f.

- A lot iterations can be devised based on the decomposition A = D E F.
 - Jacobi iteration: $Dx_{k+1} = (E + F)x_k + b$.
 - Gauss-Seidel iteration: $(D E)x_{k+1} = Fx_k + b$.
 - Successive Over-Relaxation (SOR):

$$(D - \omega E)x_{k+1} = (\omega F + (1 - \omega)D)x_k + \omega b$$



Stationary iterative method

- If the iteration converges, it converges to the linear system solution.
- In one iteration, in general, we only need to perform (1) matrix-vector multiplication; (2) perhaps solving a simpler matrix problem (usually can be achieved component-wisely). The cost of one iteration is $O(n^2)$.
- Stopping criterion:
 - monitor the residual $||b Ax_k||$
 - monitor the relative change $||x_{k+1} x_k||$
- It is often the case that we can achieve desirable accuracy in less than n iteration, meaning the iterative method is rather competitive, when compared against direct method.
- Is the method guaranteed to converge? How fast does it converge?
 There are theories to support this. Often is problem-specific.

Krylov iterative method

- We are considering solving Ax = b, in which A is large, sparse, and possibly with irregular structures.
- The idea is to solve a least-square problem: $min_{z \in K} ||b Az||$
- Krylov subspace: $K_m(A, v_1) = span\{v_1, Av_1, ..., A^{m-1}v_1\}$
- There are very fast algorithms that can locate the vector that minimizes ||b Az||, which are referred to as the Krylov method GMRES, Congujate gradient (CG), etc.

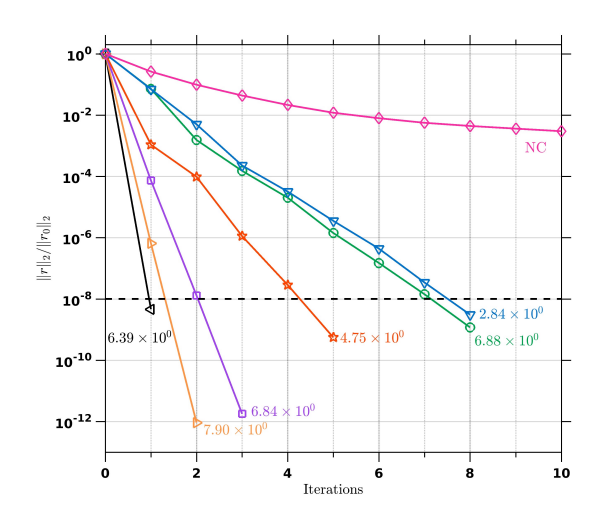
Krylov iterative method

 We can modify the linear system by considering solving

```
MAx = Mb,
or ANy = b, with x = Ny
or MANx = Mb, with x = Ny
```

The matrices M and N are left and right preconditioners, which may accelerate the Krylov iteration.

Examples: Jacobi, ILU, etc.



Summary

- Linear algebra problem is ubiquitous.
- The condition number measures the property of a matrix.
- Direct method such as LU factorization with pivoting can solve a problem in a very robust way.

Its computational cost is $O(n^3)$.

For sparse matrices, the factorization will lead to a fill-in phenomenon, which could cost a lot memory space.

 Iterative method strives to solve the linear problem by matrix-vector multiplications iteratively.

The cost per iteration is cheap $O(n^2)$ at most and O(n) for very sparse matrices.

Iteration drives the error down to certain prescribed tolerance.

Preconditioner may accelerate the convergence.

GMRES and CG are most commonly used method for solving linear systems.

Summary

Reference: Numerical Linear Algebra by L.N. Trefethen and D. Bau, III.

Differential equations

Initial value problem

Consider the problem

$$\frac{du}{dt}(t) = f(t, u(t)), t \in (0, T), \text{ with } u(0) = u_0.$$

- We may turn the continuous problem into a discrete one by looking at finite time steps: $t_0 = 0, t_1, t_2, \dots t_N = T$. Here we assume the time step size is uniform: Δt .
- Taylor series

$$u(t + \Delta t) = u(t) + u'(t)\Delta t + u''(t)\frac{\Delta t^2}{2} + \cdots$$

• Use the following: $u'(t) = \frac{u(t+\Delta t)-u(t)}{\Delta t} + O(\Delta t)$

We get a discrete problem:

$$\frac{u_{k+1} - u_k}{\Delta t} = f(t_k, u_k)$$

This is known as the Forward Euler or Explicit Euler method. First-order accurate.

Stability of the explicit Euler method

Consider the problem

$$\frac{du}{dt}(t) = f(t, u(t)), t \in (0, T), \text{ with } u(0) = u_0.$$

Let $f = -\lambda u$. The exact solution is $u(t) = u_0 e^{-\lambda t}$. It is monotonically decreasing for positive λ .

$$u_{k+1} = u_k + \Delta t f(t_k, u_k) = (1 - \lambda \Delta t) u_k = \dots = (1 - \lambda \Delta t)^{k+1} u_0$$

To have the discrete solution mimic its continuous counterpart,

$$|1 - \lambda \Delta t| < 1$$

which leads to $\Delta t < 2/\lambda$.

We call a method that has constraint on the selection of grid size to be conditionally stable.

Implicit Euler Method

Consider the problem

$$\frac{du}{dt}(t) = f(t, u(t)), t \in (0, T), \text{ with } u(0) = u_0.$$

Taylor series

$$u(t - \Delta t) = u(t) - u'(t)\Delta t + u''(t)\frac{\Delta t^2}{2} + \cdots$$

• Use the following: $u'(t) = \frac{u(t) - u(t - \Delta t)}{\Delta t} + O(\Delta t)$

We get a discrete problem:

$$\frac{u_{k+1}-u_k}{\Delta t} = f(t_{k+1}, u_{k+1})$$

This is known as the Backward Euler or Implicit Euler method. First-order accurate.

Stability of the explicit Euler method

Consider the problem

$$\frac{du}{dt}(t) = f(t, u(t)), t \in (0, T), \text{ with } u(0) = u_0.$$

Let $f = -\lambda u$. The exact solution is $u(t) = u_0 e^{-\lambda t}$. It is monotonically decreasing for positive λ .

$$u_{k+1} = u_k + \Delta t f(t_{k+1}, u_{k+1}) \Rightarrow u_{k+1} = \frac{1}{1 + \lambda \Delta t} u_k = \dots = \left(\frac{1}{1 + \lambda \Delta t}\right)^{k+1} u_0$$

To have the discrete solution mimic its continuous counterpart, there is no limit on the choice of time step size.

We call a method that has NO constraint on the selection of grid size to be unconditionally stable.

Explicit vs Implicit

Consider the problem

$$\frac{du}{dt}(t) = f(t, u(t)), t \in (0, T), \text{ with } u(0) = u_0.$$

Let $f = u^2$.

Forward Euler: $u_{k+1} = u_k + \Delta t f(t_k, u_k) = u_k + u_k^2$.

Backward Euler: $u_{k+1} = u_k + \Delta t f(t_k, u_{k+1}) = u_k + u_{k+1}^2$

In implicit method, we need to solve equations to determine the solution.

There are otheer options: Trapezoidal/Crank-Nicholson rule, Runge-Kutta rule, etc.

Boundary value problem

Consider the problem

$$\frac{d^2u}{dx^2}(x) = f\left(x, u, \frac{du}{dx}\right), \ x \in (a, b), \text{ with } u(a) = u_a \text{ and } u(b) = u_b.$$

Taylor series

$$u(x + \Delta x) = u(x) + u'(x)\Delta x + u''(x)\frac{\Delta x^2}{2} + u'''(x)\frac{\Delta x^3}{6} + \cdots$$
$$u(x - \Delta x) = u(x) - u'(x)\Delta x + u''(x)\frac{\Delta x^2}{2} - u'''(x)\frac{\Delta x^3}{6} + \cdots$$

• We can get: $u''(x) = \frac{u(x+\Delta x)-2u(x)+u(x-\Delta x)}{\Delta x^2} + O(\Delta x^2)$

We get a discrete problem:

$$\frac{u_{k+1}-2u_k+u_{k-1}}{\Delta x^2} = f(x_k, u_k, u_k')$$

Boundary value problem

We get a discrete problem:

$$\frac{u_{k+1} - 2u_k + u_{k-1}}{\Delta x^2} = f(x_k, u_k, u_k')$$

Written as a matrix problem:

$$\begin{bmatrix} 2 & -1 & \dots & 0 \\ -1 & 2 & & & 0 \\ \vdots & & \ddots & \vdots \\ 0 & 0 & \dots & 2 \end{bmatrix} \begin{bmatrix} u_1 \\ u_2 \\ \vdots \\ \vdots \end{bmatrix} = -\Delta x^2 \begin{bmatrix} f_1 + u_a \\ f_2 \\ \vdots \\ \vdots \end{bmatrix}$$

Matrix properties: Very sparse (tri-diagonal), symmetric, positive definite.

Summary

- We can address the differential equations by replacing the differential operators by their difference counterpart.
- There are explict and implict method when marching in time.
- Typically, explicit method is conditionally stable, which poses a constraint on the choice of time step size.
- Implicit method can be either conditionally stable or unconditionally stable.
- Unconditionally stable algorithms is useful for delivering steady-state solutions.
- Accuracy analysis can be achieved by standard calculus techniques, such as the Taylor expansion.
- Accuracy can be verified by numerical results.

Summary

- Manufactured solution can be an effective way in desinging code verifications.
- One may plot the error in a log-log plot against the mesh size, and the error shall be in a straight line with the slope being the accuracy.

Reference:《微分方程数值解法》戴嘉尊,邱建贤。