Classical iterative methods for linear systems

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MATH 615 Numerical Analysis of Differential Equations

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Classical iterative methods for linear systems

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example linear systems

suppose we want to solve the linear system

$$A\mathbf{x} = \mathbf{b} \tag{1}$$

where $A \in \mathbb{R}^{m \times m}$ and $\mathbf{b} \in \mathbb{R}^{m}$, to find $\mathbf{x} \in \mathbb{R}^{m}$.

 throughout these notes we use just two examples: LS1

$$\begin{bmatrix} 2 & 1 & 0 \\ 0 & 2 & 1 \\ 1 & 0 & 3 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 2 \\ 1 \\ 4 \end{bmatrix}$$

LS2

$$\begin{bmatrix} 1 & 2 & 3 & 0 \\ 2 & 1 & -2 & -3 \\ -1 & 1 & 1 & 0 \\ 0 & 1 & 1 & -1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \\ x_4 \end{bmatrix} = \begin{bmatrix} 7 \\ 1 \\ 1 \\ 3 \end{bmatrix}$$

- on P17 (Assignment #5) you will check that these are well-conditioned linear systems
- it is trivial to find solutions of LS1, LS2 using a "x = A\b" black box, but these examples stand-in for the large linear systems we get from applying FD schemes to ODE and PDE problems

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residual

• the residual of a vector v in linear system (1) is the vector

$$\mathbf{r}(\mathbf{v}) = \mathbf{b} - A\mathbf{v} \tag{2}$$

• making the residual zero is the same as solving the system:

$$A\mathbf{x} = \mathbf{b} \iff \mathbf{r}(\mathbf{x}) = \mathbf{0}$$

- evaluating $\mathbf{r}(\mathbf{v})$ needs a matrix-vector product and a vector subtraction
 - requires $O(m^2)$ operations at worst
 - by comparison, applying Gauss elimination to solve linear system (1) is an $O(m^3)$ operation in general
- FD schemes for DEs generate matrices A for which the majority, often 99% or more, of the entries are zero
 - a matrix with enough zeros to allow exploitation of that fact is called *sparse*
 - evaluating the residual of a sparse matrix typically requires O(m) operations
 - even if A is sparse, A^{-1} is generally *dense*, i.e. most entries are nonzero

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Richardson iteration

- iterative methods for linear system (1) attempt to solve it based only on operations like computing the residual, or applying A to a vector
 - one wants the sequence of approximations, the iterates, to *converge* to the solution $\mathbf{x} = A^{-1}\mathbf{b}$
 - Iterative methods always require an initial iterate x₀
- *Richardson iteration* adds a multiple ω of the last residual at each step:

$$\mathbf{x}_{k+1} = \mathbf{x}_k + \omega(\mathbf{b} - A\mathbf{x}_k) \tag{3}$$

• for system LS1, using initial iterate $\mathbf{x}_0 = \mathbf{0}$ and $\omega = 1/5$, (3) gives:

$$\mathbf{x}_{0} = \begin{bmatrix} 0\\0\\0 \end{bmatrix}, \, \mathbf{x}_{1} = \begin{bmatrix} 0.4\\0.2\\0.8 \end{bmatrix}, \, \mathbf{x}_{2} = \begin{bmatrix} 0.6\\0.16\\1.04 \end{bmatrix}, \, \mathbf{x}_{3} = \begin{bmatrix} 0.728\\0.088\\1.096 \end{bmatrix}, \, \dots, \, \mathbf{x}_{10} = \begin{bmatrix} 0.998\\-0.017\\1.01 \end{bmatrix}, \dots$$

these iterates seem to be converging to $\mathbf{x} = \begin{bmatrix} 1 & 0 & 1 \end{bmatrix}^{\top}$, which is the solution to LS1

- a complex number λ ∈ C is an *eigenvalue* of a square matrix B ∈ ℝ^{m×m} if there is a nonzero vector v ∈ C^m so that Bv = λv
- the set of all eigenvalues of *B* is the *spectrum* $\sigma(B)$ of *B*
- the spectral radius $\rho(B)$ is the maximum absolute value of an eigenvalue:

$$ho({\it B}) = \max_{\lambda \in \sigma({\it B})} |\lambda|$$

even if B is real, λ may be complex—the roots of a polynomial with real coefficients may be complex—and if λ is complex and B is real then v must be complex

spectral properties and convergence of iterations

- properties of a matrix B described in terms of eigenvalues are generically called spectral properties
- some examples:
 - $\rho(B)$
 - $\|B\|_2 = \sqrt{\rho(B^\top B)}$
 - the 2-norm condition number $\kappa(B) = ||B||_2 ||B^{-1}||_2$
- a general idea:

whether an iterative method for solving $A\mathbf{x} = \mathbf{b}$ converges, or not, depends on spectral properties of A, or on matrices built from A

- the right-hand side b and the initial iterate x₀ generally do not determine whether an iteration converges
 - $\circ~$ a good choice of \boldsymbol{x}_{0} can speed up convergence

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convergence of the Richardson iteration

rewrite the Richardson iteration (3) as

$$\mathbf{x}_{k+1} = (I - \omega A)\mathbf{x}_k + \omega \mathbf{b}$$

 the lemma on the next slide shows that the Richardson iteration converges if and only if all the eigenvalues of the matrix *I* – ω*A* are inside the unit circle:

(3) converges if and only if
$$\rho(I - \omega A) < 1$$

• $\rho(I - \omega A) < 1$ means $(I - \omega A)\mathbf{x}_k$ is smaller in magnitude than \mathbf{x}_k • *if* $||I - \omega A|| < 1$ *then* (3) converges¹

¹recall $\rho(B) \leq ||B||$ in any induced matrix norm

convergence lemma

Lemma

$$\mathbf{y}_{k+1} = M\mathbf{y}_k + \mathbf{c}$$

converges to the solution of $\mathbf{y} = M\mathbf{y} + \mathbf{c}$ for all initial \mathbf{y}_0 if and only if

 $\rho(M) < 1.$

Proof.

Solve the iteration by writing out a few cases:

$$\begin{split} \mathbf{y}_2 &= M(M\mathbf{y}_0 + \mathbf{c}) + \mathbf{c} = M^2 \mathbf{y}_0 + (I + M) \mathbf{c}, \\ \mathbf{y}_3 &= M(M^2 \mathbf{y}_0 + (I + M) \mathbf{c}) + \mathbf{c} = M^3 \mathbf{y}_0 + (I + M + M^2) \mathbf{c}, \end{split}$$

By induction we get $\mathbf{y}_k = M^k \mathbf{y}_0 + p_k(M)\mathbf{c}$ where $p_k(x) = 1 + x + x^2 + \dots + x^{k-1}$. But $p_k(x) \to 1/(1-x)$ as $k \to \infty$ iff $x \in (-1, 1)$. Also, $\rho(M) < 1$ iff $M^k \to 0$. Thus $\mathbf{y}_k \to (I-M)^{-1}\mathbf{c}$ iff $\rho(M) < 1$.

convergence of the Richardson iteration 2

 since the Richardson iteration converges iff ρ(I – ωA) < 1, we choose ω based on the principle that

 ωA should be close to the identity I

o often not possible!

• in small cases we can graph $f(\omega) = \rho(I - \omega A)$:



matrix splitting

- unlike Richardson, most classical iteration methods "split" the matrix *A* before iterating
- the best known, and simplest, iteration based on splitting is *Jacobi iteration*, which extracts the diagonal of *A* (and inverts it)
- the splitting we consider is

$$A = D - L - U$$

where

- D is the diagonal of A
- *L* is strictly lower triangular ($\ell_{ij} = 0$ if $i \le j$)
- *U* is strictly upper triangular $(u_{ij} = 0 \text{ if } i \ge j)$
- you can split any matrix this way
- see section 4.2 of the textbook
- so that *D* is an invertible matrix, for the remaining slides we assume all diagonal entries of *A* are nonzero: $a_{ii} \neq 0$

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the Jacobi iteration is

$$D\mathbf{x}_{k+1} = \mathbf{b} + (L+U)\mathbf{x}_k \tag{4}$$

• if it converges then $D\mathbf{x} = \mathbf{b} + (L + U)\mathbf{x}$, which is the same as $A\mathbf{x} = \mathbf{b}$

• we could also write it as $\mathbf{x}_{k+1} = D^{-1} (\mathbf{b} + (L+U)\mathbf{x}_k)$ or as

$$x_{i}^{(k+1)} = \frac{1}{a_{ii}} \left(b_{i} - \sum_{j \neq i} a_{ij} x_{j}^{(k)} \right)$$
(5)

where $x_i^{(k)}$ denotes the *j*th entry of the *k*th iterate \mathbf{x}_k

• make sure you understand why (4) and (5) are the same!

Gauss-Seidel iteration

- Gauss-Seidel iteration extracts the non-strict lower-triangular part of A (and inverts it)
- again if A = D L U then it is

$$(D-L)\mathbf{x}_{k+1} = b + U\mathbf{x}_k \tag{6}$$

- we could also write it " $\mathbf{x}_{k+1} = (D L)^{-1} (b + U\mathbf{x}_k)$ " but that would miss the point!
- instead we write it as $D\mathbf{x}_{k+1} = b + U\mathbf{x}_k + L\mathbf{x}_{k+1}$ or equivalently:

$$x_{i}^{(k+1)} = \frac{1}{a_{ii}} \left(b_{i} - \sum_{j > i} a_{ij} x_{j}^{(k)} - \sum_{j < i} a_{ij} x_{j}^{(k+1)} \right)$$
(7)

- the lower-triangular entries of A apply to those entries of x_{k+1} which have already been computed
- form (7) is actually *easier* to implement than Jacobi (5) (why?)

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- the convergence lemma says that
 - Jacobi iteration converges if and only if $\rho(D^{-1}(L+U)) < 1$
 - Gauss-Seidel iteration converges if and only if $\rho((D-L)^{-1}U) < 1$
- these conditions are hard to use in practice because computing a spectral radius can be just as hard as solving the original system

diagonally-dominant matrices

- definition. A is strictly diagonally-dominant if $|a_{ii}| > \sum_{i \neq i} |a_{ij}|$
 - LS1 is strictly diagonally-dominant
 - LS2 is not
- two relatively-famous theorems² are these:
 - *theorem.* if *A* is strictly diagonally-dominant then both the Jacobi and Gauss-Seidel iterations converge
 - *Theorem.* if A is symmetric positive definite then Gauss-Seidel iteration converges
- unlike the " $\rho(...) < 1$ " conditions on the last slide:
 - it is easy to check diagonal-dominance, and it is a common property of the matrices coming from FD schemes on ODEs and PDEs
 - these are only *sufficient* conditions, e.g. there are nonsymmetric *A*, which are *not* diagonally-dominant, but for which the iterations converge

see problems P19 and P20

²section 11.2 of Golub and van Loan, *Matrix Computations*, 4th edition 2013 • (= • • = • • • •

past

- the Jacobi and Gauss-Seidel iterations are from the 19th century
 - Richardson iteration first appears in a 1910 publication
- the early history of numerical partial differential equations, e.g. in the 1920 to 1970 period, heavily used these classical iterations
 - a generalization of Gauss-Seidel iteration called *successive over-relaxation*, was a particular favorite around 1970; see section 4.2 of the textbook
- none of these iterations work on system LS2

recent past and future

- there are better iterative ideas, and they flourished starting in the 1980-90s ... and far into the future
 - among the best known are CG = conjugate gradients (actually from 1950-60s) and GMRES = generalized minimum residuals (from a 1986 paper by Saad and Schultz)
 - GMRES works (i.e. converges at some rate) on LS2
 - but there can be no "good iteration" with a universally-fast convergence rate³
- iteration to solve linear systems is the future:
 - o it is obligatory on sufficiently-big systems
 - o it works better in parallel than direct methods like Gauss elimination
 - o it can exploit partial knowledge of the underlying model

³remarkably, there is a 1992 theorem by Nachtigal, Reddy, and Trefethen that says this 📱 🗠 🔍

biographies

- Gauss (1777–1855) did big stuff, not just the little Gauss-Seidel thing: en.wikipedia.org/wiki/Carl_Friedrich_Gauss
- Jacobi (1804–1851) also has his name on the "Jacobian", the matrix of derivatives appearing in Newton's method for systems of equations: en.wikipedia.org/wiki/Carl_Gustav_Jacob_Jacobi
- Seidel (1821–1896) is relatively little known: en.wikipedia.org/wiki/Philipp_Ludwig_von_Seidel
- Richardson (1881–1953) is the most interesting. He invented numerical weather forecasting, doing it by-hand for fun during WWI. Later, as a pacifist and quaker, he quit the subject entirely when he found his meteorological work was of most value to chemical weapons engineers and the British Air Force:

en.wikipedia.org/wiki/Lewis_Fry_Richardson