Solving Linear Equations

We have discussed general methods for solving
arbitrary equations. For the most part, we
restricted our attention to the one-dimensional case.
We also looked at a special class of equations, namely
those given by polynomials. Another special class is
given by linear equations. Of course, the one-dimensional case is absolutely trivial, so one naturally considers
systems of equations.
I assume that you are familiar with the bosic
results of linear algebra, such as the diagona lization
theorems and Gaysian Elimination. This section
provides a very quick review, then discusses SVD.
Applications: Applications for linear equations are numerous,
mose include of course, linear Programming
problems. Others are the local solution
of non-linear differential equations via
linearization, and the determination of
minima in least-squares problems

A quick review of important definitions and facts.

1. Given a linear function f: Rn - Rm, and given a pair of bases,

B₁ = {e,,..., en } Ar R^m,
B₂ = {d₁,..., d_m} Ar R^m,

We can represent f by an man matrix A.

Notice that the matrix A depends on the choice of bases B, *B2.

Often we just let B, *B2 be the obvious' bases for R* 1 Rm.

Moral: A matrix is a particular coordinate representation

Moral: A matrix is a particular coordinate representation of the linear function f.

2. Given an man matrix to we make the following definitions:

column space - linear combinations of the columns of A row space - linear combinations of the vous of A

If we think of A as defining the linear mapping

 $A: \mathbb{R}^{n} \to \mathbb{R}^{m}$ $= \begin{bmatrix} x_{1} \\ \vdots \\ y_{m} \end{bmatrix} \mapsto \begin{bmatrix} y_{1} \\ \vdots \\ y_{m} \end{bmatrix} = A \begin{bmatrix} x_{1} \\ \vdots \\ x_{m} \end{bmatrix}$

then the column space of A is a vector subspace of R".

consisting of all points in R" that are image vectors under A.

Note that the now space of A is just the column space of A, the transpose of A.

And we define:

null space — set of vectors x in Rn such that Ax =0.

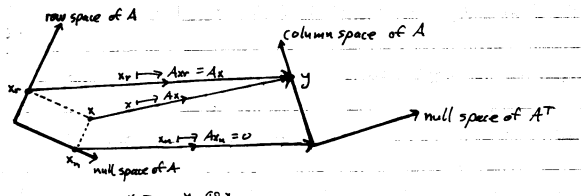
The following relationships are useful to remember:

- · dim (row space) = dim (col space), either of which is called the rank of A,
- · The row space and null space are complementary (perpendicular) subspaces of 18. In other words,

1Rn = row space @ null space

n = dim (row space) + dim (null space)

In picture form:



$$y = Ax$$

$$Ax = Ax_r + Ax_n$$

$$= Ax_r + Q$$

$$= Ax_n$$

- Suppose A is an man matrix, and consider the system of equetims Ax = b.
 - If b is not an element of the column space of A, then we say that the system is inconsistent (or ... overde termined).
- If bis in the column space of A and the null space of A is_ non-trivial, then we say that the system is underdetermined. In this case there is a whole family of solutions, given by the affine set

 $x_o + N$ Whene to is any particular solution $Ax_0 = b$, and

N is the nell space of A,

. If A is an new square matrix we say that A is singular ist det(A) = 0

iff rank (A) < niff the rows of A are not linearly independent iff the columns of A are not linearly independent iff the dimension of the null space of A is non-zero.

If A is not invertible.

Quick review of matrix decompositions: (we will look at SVD in more detail)

Factorization based on elimination

Twith min = rank(A)

Given an man matrix A, we can write A in the form

PA = LDU

where P is an man permutation matrix that specifies row interchanges, L is an mam square low-triangular matrix with Is on the diagonal, U is an min upper-triangular matrix with Is on the diagonal, and D is an mam square diagonal matrix.

L More generally: we may also need to perform column interchanges on A, in which case we get two permutation matrices, so P, AP2 = LDU. I Yet more generally: PA = LU with diagonal entries of Uno longer required to be 1.

- 1) The entries on the diagonal of D are sometimes called "pivots" (after the Ganssian Elimination algorithm).
- 2) The product of the pivots is equal to = det (A), whenever A is a square matrix. (-"if odd rowinerways)
- 3) If A is symmetric and P=I, then U=L.
- 4) If A is symmetric positive-definite, then $U = L^T$ and the diogonal entries of D are strictly positive.

[If mon, as in this example, then the last m-n rows of U are zero.]

(b)
$$\begin{bmatrix} 1 & 1 & 0 \\ 2 & 1 & -1 \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ 2 & 1 \end{bmatrix} \begin{bmatrix} 0 & -1 \end{bmatrix} \begin{bmatrix} 0 & 1 & 1 \\ 0 & 1 & 1 \end{bmatrix}$$

Algorithm: Gaussian Elimination directly yields this decomposition.

Application: As with most decompositions/factorizations, the

Ax = b.

Suppose A is square and non-singular.

Then solving Ax = b really means solving.

LDUx = Pb.

In turn this entails solving two simpler problems;

Each of these problems can be solved easily using forward backsubstitution (D-1 is easy to compate since D is diagonal with non-zero entires).

· Factorizations based on eigenvalues

These are the standard factorizations one learns in a linear algebra course. Two important ones are:

(i) If A is a square nxn matrix with a linearly independent eigenvectors, then

 $A = S \Lambda S^{-1},$

where A is a diagonal matrix whose entries are the eigenvalues of A,

and S is a matrix whose columns are the eigenvector

This factoritation is not always possible. (One case in which it is possible occurs when A has, distinct eigenvalues.)

(ii) one can always decompose A in Jordan form, is.

 $A = MJM^{-1}$

where $J = \begin{bmatrix} J_1 & 0 \\ 0 & J_5 \end{bmatrix}$ is a block matrix, st each block

Ji = [li!] with li an eigenvalue of A.

Here s is the number of independent eigenvectors of A. M. countsts of eigenvectors and "generalized" etaluvectors.

Side note

What is a "generalized" eigenvector?

well, suppose a Jordan block is of the form $J_{i} = \begin{bmatrix} 4 & 1 \\ 0 & 4 \end{bmatrix} \qquad (1i = 4)$

Then -xi = [1] is an elgenvector

and is a generalised eigenvector

 $\int_{\mathcal{C}} X_1 = \lambda_i X_1$

a Jix2 = lix2 +x

"not quite" an eigenvalue.

•	Factorizations Dased on AA
w 40 - 20°	Let's look at two such factorizations, QR and SVD.
(i)	<u>QR</u>
	Suppose A is an men matrix with independent columns. We can factor A as:
	A = QR Q is man
	R is nkn
	Q has the same column space as A, but it's columns are orthonormal vectors. In other words, $Q^TQ = I$ (QQT may not be I , if A is not square).
	R is invertible and upper triangular.
*	Here are two possible algorithms for computing this factorization:
	1) Use Gram-Schmidt to orthogonalize the
	vectors constitute the columns of Q.
	The matrix R is formed by keeping truck
·	of the Gram-Schwidt operations (specifically, R expresses the columns of A as linear combinations of the columns of Q).

(It is positive definite because the columns of A are independent.

Second, compute the LDU theterization of ATA.

One finds that ATA = LDL, with Llower-triangular and D diagonal strictly positive.

Finally, let $R = D^{\frac{1}{2}}L^{T}$ and $Q = AR^{-1}$

Unfortunately, both of these straightforward algorithms have poor numerical stability. In practice one uses a different algorithm. See \$11.3 of NRiC for a good algorithm.

Applications:

- (a) The QR algorithm. This is an iterative method

 that repeatedly produces QR fectorizations of matrices

 derival from A, building the eigenvalues of A in the process
- (b) Suppose we wish to solve an overconstrained system Ax = b in the least-squares sense. The last-squares solution \overline{x} is given by $\overline{x} = (A^TA)^TA^Tb$, assuming the columns of A are independent. But $Q^TQ = \overline{I}$, so $\overline{x} = R^{-1}Q^Tb$.

So, we can obtain \bar{x} by computing Q^Tb , then using backsubstitution to solve $R\bar{x} = Q^Tb$. — This is numerically more stable than solving the system $A^TA\bar{x} = A^Tb$ directly.

(ii) <u>5VD</u>

Any man matrix A can be factored as

 $A = U \leq V^{T}$

where U is an mxm orthogonal matrix whose cohumns are the eigenvectors of AAT

V is an nxn orthogonal metrix whose columns are the eigenvectors of ATA

E is a diagonal man matrix of the form

with 0, 20, 2 --- 20, 70

and k = rank (A)

The {Gi} are the square-roots of the eigenvalues of A'A.

They are called the singular value of A.

Fact: If A is symmetric positive definite then C=V and Ξ is just the eigenvalue matrix J.

Motivation

- (i) The basic goal is to "solve" the system Ax = b for all matrices A and vectors b.
- (ii) A second goal is to solve Ax = b using a numerically stable algorithm.
- (iii) A third goal is to solve Ax=b in a reasonably efficient manner. For instance, we do not want to compute A-1 using determinants.

Comments:

Regarding the basic gool:

- If A is square and invertible we of course want the solution $x = A^{-1}b$
- set of solutions.
- But this case arises a lot in practice, so instead we will ask for the least-squires.

 Solution. In other words, we want that x which minimizes the error ||Ax-b||. Geometrically, Ax is the point in the column space of A closest to b.

Regarding stability and efficiency:

Gaussian Elimination is reasonably efficient, but it is not numerically very stable. In particular, elimination does not deal well with nearly singular metrices. The method is not designed for overconstrained systems. Even for underconstrained systems, the method requires extra work.

The poor numerical character of climination can be seen in a couple ways

The elimination process assumes a non-singular matrix. But singularity, and rank in general, is a slippery concept. After all, the matrix A contains continuous, possibly naisy, numbers. Yet, rank is a discrete integer. Technically both these sets are linearly independent vectors;

$$\left\{ \begin{bmatrix} 1 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}, \begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix} \right\} \left\{ \begin{bmatrix} 1,00 \\ 1,00 \\ 1,00 \end{bmatrix}, \begin{bmatrix} 1,00 \\ 1,00 \\ 1,01 \end{bmatrix} \right\}$$

Jet, the first set seems genuinely independent, while the second set seems "almost dependent."

2) Besed on elimination, one solves Ax = b by solving (via back substitution) Ly = b and $Ux = D^{-1}y$.

If A is nearly singular (eignitit is given by the second get of column vectors above), then D will contain near-zero entries on its diagonal, and thus D-1 will contain large numbers. That's all line and good. In principle one needs the large numbers to obtain a true solution. The problem is if A contains noisy entries. Then the large numbers may be pune noise — noise that dominates the true information. Furthermore, since Land U can be fairly arbitrary, they may distort or magnify that noise across other variables.

SVD in more detail

Rocall that it is sometimes possible to decompose an man matrix A as:

A = S.15-1

where A is diagonal, containing the eigenvalues of A, and S is a change-of-basis matrix containing the eigenvectors of A.

Why is this nice?

The answer lies in the change of coordinates $y = 5^{-1}x$.

Instead of working with the system Ax = b, we can work with the system Ay = c, where $c = 5^{-1}b$.

Since A is diagonal, we really have the trivial system.

Siyi = ci, i = 1, ..., n.

Esis are the eigenvalues of A.

If this system has a solution, then another change of coordinates gives us x (as x = Sy).

Note: Offhand, there is no reason to believe that computing 5-16 is any easier than computing A-16. However, in the ideal case the eigenvectors of A are orthogonal. This is true, for instance, if AAT = ATA. In that case the column of 5 are orthogonal, and so we can take 5 to be orthogonal. But then 5-1 = 5T, and the problem of solving Ax=6 becomes very simple.

worse, who	at do we	do if.	A is not	- square?	
The answ	r is to	work with	$A^{T}A$	al AAT	
Suppose A	is an mxv	matrix ,			
Suppose A The previous	discussion.	implies th	e following	decomposition	·s,·
ATA =	VDV^{T}	V nr		with the eigenvect	
			n diagonal, u	ith the (non-negatives of	ATA.
A A =	UD'UT	U mi	in orthogonal	, with the eigen	vectors of
		Di n.×	m diagonal,	with the (non-	AAT,
				4	1. 0
entries (except that I	he order might	nove the sav t be differ	ne non-zero d	Togonal

Recall $A = U \leq V^T$ $\leq = \begin{bmatrix} \sigma_i & \sigma_k \\ \sigma_i & \sigma_k \end{bmatrix}$
Observations
1) Rank (A) = Rank (E) = k
2) COLSPACE(A) = Span (first & columns of U) Note: these columns form an orthonormal basis for the oly
3) Nullspace(A) = Span (last n-k columns of V) Note: these columns form an orthonormal busis for the mall g
4) Each of U1 V is an orthornormal change of bosis matrix. We can therefore think of Z is a simple mapping between two different coordinate systems that amounts to dilation in each coordinate of the look at the effect on a unit circle, we get the picture
Rn In given In "U" In given coords coords coords coords

Observation (1) revisited

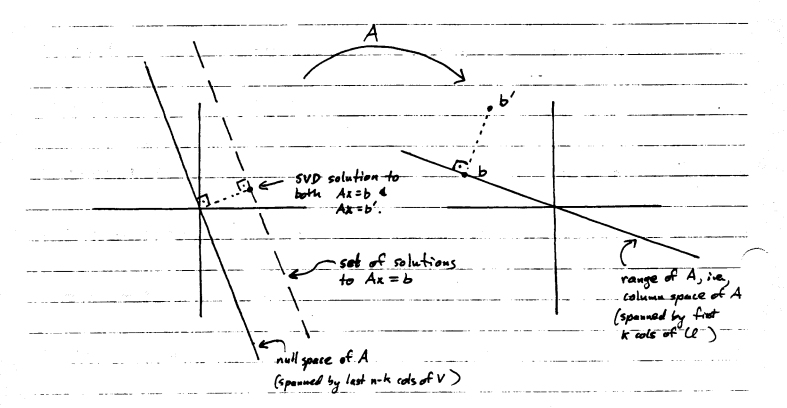
(1) tells us that we can determine the rank of A by counting the non-zero entries in E.
In fact, we can do better. Recall that one of our complaints about Gaussian elimination was that it did not handle noise or nearly singular matrices well. SVD remedies this situation.
For example; suppose that an new matrix A is nearly singular. Indeed, perhaps A should be singular, but due to revisy date it is not quite singular, this will show up in 2 as: 1) All of the a diagonal entries are almost zero,
More generally, an new matrix A may appear to have rank k, yet when we look at Z we may find that same of the singular values are very close to zero. If there are I such values, then the "true" rank of A is probably k-ly and we would do well to modify Z. Specifically, we should replace the I nearly zero singular values with zero.
Geometrically, the effect of this replacement is tex reduce the column space of A and increase its null space. The paint is that the column space is worped along directions for which $\sigma: \approx \mathcal{O}$. In effect, solutions to $Ax = b$ get pulled off to infinity. (since $\sigma: \propto \infty$) along vectors that are almost in the null space. So, it is better to ignore the its coordinate by serving $\sigma: \sim \infty$.
Ex A = (1.00 1.00 1.00) yields = (3.01 0 0)

For the moment, let us suppose A is a square motive.

Suppose A: IR" -> IR", __rank (A) = + < n.

The following: picture sketches the way in which SVD

solves the system Ax = b (we will see how to do this shortly).



Observe: i) The system Ax = b has an affine set of solutions,

given by $x_0 + N$, where x_0 is any solution and Nis the null space of A. It is easy to describe N, given

the SVD decomposition. N is just the span of the last n-t cols of V.

2) Ax = b' has no solution since b' is not in the colspace of A.

SVD will: 1) SVD solves Ax = b by determining that x which is closest to the origin (10, the x with minimum magnitude).

2) SVD solves Ax=b' by projecting b' onto the column
space of A, obtaining b, then solving Ax=b. In other
words, SVD obtains the least-squares solution.

So, how do we compute a solution to Ax=b for all these cases?

The cool thing is that the procedure is the same.

Def Given diagonal E, let us write E to mean the diagonal matrix whose diagonal entries are of the form:

$$\left(\frac{1}{2}\right)_{ii} = \begin{cases} \frac{1}{2ii} & \text{if } \sum_{ii} \neq 0 \\ 0 & \text{if } \sum_{ii} = 0 \end{cases}$$

And, as we mentioned sometimes it is useful to set Ziz = 0 19 Ziz 20.

then
$$\frac{1}{2} = \begin{pmatrix} \frac{1}{2} & 0 & 0 \\ 0 & \frac{1}{5} & 0 \\ 0 & 0 & 0 \end{pmatrix}$$

Observe a) If & is invertible, then \(\xi = \xi^{-1} \).

To solve Ax = b, we first compute the SVD decomposition $A = U \leq V^T$, then compute $\overline{x} = V \leq U^T b$

Comments: (i) If A is invertible, then \bar{x} is the unique solution to Ax = b.

(ii) If A is singular and b is in the range of A, then \bar{x} is the solution closest to the origin. $V \neq Cl^T$ is a <u>pseudo-inverse</u>,

The affine set of solutions is: $\bar{x} + span(last n + k cols of V)$, k = rank(A).

(iii) If A is singular and b is not in the range of A, then \bar{x} is.

the lent-source solution.

So far we have assumed that A is square.
So far we have assumed that A is square. Q: How do we solve $Ax = b$ for general $m \times n$ matrix A ?
A.: Just as we did before, namely x = V \(\frac{1}{2} \lambda \). The only issue is an implementation issue.
It is easiest to implement SVD when m?n.
So, let's take a quick look at the non-square cases.
1) Suppose m &n. So, there are fewer equations than unknowns.
we thus do not expect a unique solution.
First we turn A into an non matrix by adding rows of Os, and we turn b into an not vector by adding extra O's. Then we apply SVD Note that we are in a situation similar to case (is) - A is singular and b is in the range of A.
2) Suppose m7n. So, there are more equations than unknowns. This is a clossic least squares problem.
We can apply SUD directly, and get
$ (x) = (V) (\frac{\overline{\sigma_1}}{\overline{\sigma_n}} \sigma \cdots \sigma) (U^T) (b) $ $ (x) = (V) (x) (x) (x) (x) (x) (x) (x) (x) (x) (x$

In general there won't be any zero singular values.

However, if A has column degeneracies there may be near-zero.

singular values. It may be useful to zero these out, to remove noise.

(The implication is that the overdetermined set is not quite

as averdetermined as it seemed — i.e. the null space is not

trivial

Implementation note:

U i	on look at NRiC you will see that is not square mxm, but rather mxn. E is not mxn, but rather nxn.
What	is going on?
If n	this is an efficiency hack. 12 n , as NR: C assumes, then ost 12 singular values in E are non-zero.
—Thus —and	we can ignore the last m-n columns of Ul the last m-n rows of 5,
naga da na - 1 da sa - 2 da	

SVD Internals

You should have some idea of how the SVD algorithm actually works, so let's take a brief look.

(See _Ch. 9 of Forsythe et al. for more details.)

It is useful to establish a controst with Gaussian Elimination reduces

Elimination. Recall that Gaussian Elimination reduces

a matrix. A by a series of row operations

that zero out portions of columns of A.

Row operations imply pre-multiplying the matrix A.

The your aperations are all collected together in the matrix L-1 (recall A = LDLe).

In contrast, SVD zeros out portions of both

Nows and columns. Thus, whoreas Gaussian Elimination
only reduces A using pre-multiplication, SVD uses both

pre- and post-multiplication. As a result, SVD

can at each stage rely on orthogonal metrices

to perform its reductions on A. By using

orthogonal matrices, SVD reduces the risk of

magnifying noise and errors. (The pre-multiplication

metrices are gathered together in the matrix UT,

while the post-multiplication metrices are gathered

together in the matrix V.)

There are two phases to the SVD decomposition algorithm!
1) First SVD reduces A to bidiagonal form using a series of orthogonal transformations.
This phase is deterministic, with a running time that depends only on the size of the matrix A.
2) Next SVD removes the superclingonal elements from the matrix produced by phase (1), again using orthogonal transform
This phase is iterative (numerical), but converges quickly,
Let's take a slightly closer look at phose (1). The first step of this phase creates two oxthogonal matrices U, and V, such that
That $ \begin{array}{ccccccccccccccccccccccccccccccccccc$
and $U_1 A V_1 = \begin{pmatrix} q_1 & q_1 & 0 & \cdots & 0 \\ & & & & & \\ & & & & & \\ & & & & &$
If A is man than B" is (m-1) * (n-1). The next step of phase (1) recursively works on B", and so forth, until phase (1) produces orthogonal matrices (l,, llng, Vi,, ke, such that - (ln-10. U, A V Vn2 is bioliegemal

	structed from Householder matrices. from linear algebra:	
Def	If a is a unit vector in TR, then an nxn matrix P of the form	
	$P = I - 2\alpha\alpha^{T}$	
	is said to be a Householder Reflection.	
Note:	1) P is symmetric orthogonal	
	2) When a vector x is pre-multiplied by P it is reflected in the hyperplane span $\xi \alpha \xi^{\perp}$	-
,	D.L.	
	> span \$ x \$ \\	
	₹ Px	-
why	are Householder reflections so useful?	
	I leave that as an exercise for someone to write (See for instance, Ch. 9 of Forsythe et al. and 33.3 of Golub	-up,
	(Coe for include the of tarsythe et al. and	