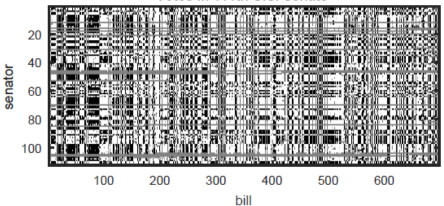
## Chapter 7 Matrix Analysis





# Section 7.1 From matrix to insight

### From matrix to insight

- Any two-dimensional array of numbers may be interpreted as a matrix
- These may come from widely disparate tasks
- Examples:
  - Text/document search
  - Voting patterns
  - o Preferences/ratings Netflix recommendations
  - o Graphs
  - Networks: social, political, co-authorship, casting in movies,...
  - Images

### Examples of matrices

- A term-document matrix may be used for analyzing a body of documents (or corpus)
- Each column may be a document; each row a term
- E.g, your textbook may have words like "numerical," "discretization," "matrix," "integration" and "function"
- An analysis textbook may have words like "integration," "function," "continuous" and so forth
- The occurrence of "function" may be often in both books, but the other terms are likely to be much different in frequency
- Meaning could be inferred from this kind of approach: latent semantic analysis

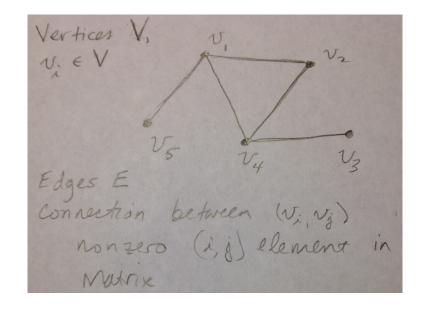
### Examples of matrices

• A term-document matrix example: matrix in green

Term	FNC by TAD+RJB	Analysis by Rudin	SM by Trefethen	NYT coffee table book
Numerical	251	2	179	0
Integration	37	275	33	18
Function	175	345	123	0
Matrix	151	11	87	0
Continuous	15	212	11	0
Spectral	15	0	124	0
Citizen	0	0	0	13

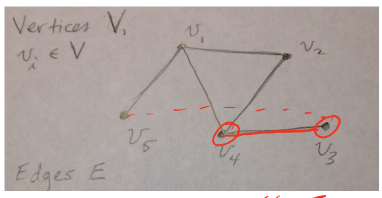
### Graphs as matrices

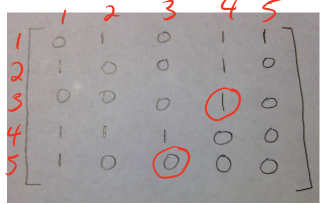
- Could be social network, internet (or subset of it), the web, etc
- Consider graphs first
- A graph is a set of nodes V connected by set of edges E
- Sometimes the graph is denoted G(V,E)



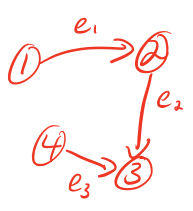
### Adjacency matrix

- We want adjacency matrix  $A = \{a_{ij}\}$  that represents this graph
- Edges are unweighted and undirected
- If an edge between  $(v_i, v_j)$ , then a one is placed in both  $a_{ij}$  and  $a_{ji}$  (symmetric)
- No self-connections
- Corresponding matrix at right





## Incidence matrix:

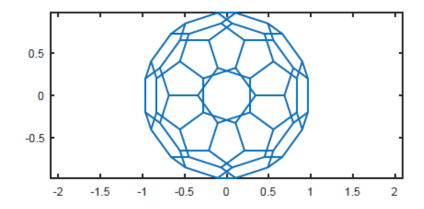


### Adjacency matrix: Buckyball

 Matlab has a built-in example of a graph representing the arrangements of carbon atoms in a C<sub>60</sub> molecule, a.k.a. the buckyball:

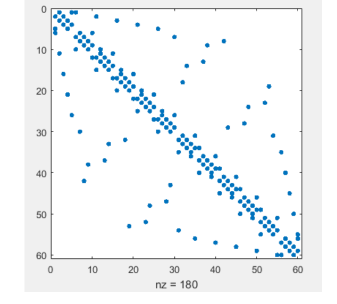
- The output has the adjacency matrix A and the vertex locations v
- Plotting the graph:

```
gplot(A,v), axis equal
```



### Adjacency matrix: Buckyball

- We know there are 60 nodes; how many edges?
- There is more than one way to compute this
- For undirected, you could use the spy command:
   |A,v|=bucky;
   spy(A)
- Is nnz it? Not quite
- You could use the triu and sum commands; how?





### **Images**

- Pictures and images are matrices in matlab
- Use imread and imshow to display them

```
A = imread('peppers.png');
size(A)
ans =
   384 512 3
```

- Three "layers" are RGB components
- To display: imshow (A)



### **Images**

- We can convert to grayscale to get only a 2D matrix
- Use rgb2gray and double:

```
A = rgb2gray(A); % collapse from 3 dimensions to 2
A = double(A); % convert to floating point
[m,n] = size(A)
```

```
m =
384
n =
512
```

- To display: imshow (A, [0, 255])
- What if different range spec'd?





### Section 7.2 Eigenvalue decomposition

### The eigenvalue decomposition

- We can decompose a matrix **A**, under some assumptions, into a useful set of other matrices using its eigenvalues and eigenvectors.
- Recall that the eigenvalue problem is  $Ax = \lambda x$ , for the eigenvalues  $\lambda$  and eigenvectors x
- (Sometimes the eigenvectors are referred to as the eigenspaces.)
- We need to review some results and terminology from linear algebra before proceeding
- In particular we need to recall the complex-valued case

### Complex vectors and matrices

- Recall complex numbers x = a + ib, where  $i^2 = -1$  and a, b are real-valued.
- The complex conjugate is  $\bar{x} = a ib$
- For a matrix A with complex-valued elements, the transpose needs to use complex conjugation at the same time:  $A^* = (\overline{A})^T = \overline{A^T}$
- The *hermitian* will keep certain properties analogous with the real-valued case
- e.g.,  $(A^TA)^T = A^TA$  so that  $A^TA$  is symmetric. So is  $(A^*A)^* = A^*A$
- Essentially, hermitian replaces transpose for complex-valued matrices and vectors

### Complex vectors and matrices

- Essentially, hermitian replaces transpose for complex-valued matrices and vectors
- For inner products with complex u and v:  $u^*v = \sum_{k=1}^n \overline{u}_k v_k$ ,
- This defines the two-norm for vectors,  $||u||_2 = \sqrt{u^*u}$ , and this will in turn define the two norm for matrices
- Definition of orthogonal
- And orthonormal set of vectors  $u_j$ , j=1,2,...,n:  $u_i^*u_j=1$  if i=j, otherwise  $u_i^*u_i=0$

### Complex vectors and matrices

- A square real-valued matrix with orthonormal columns is an orthogonal matrix
- A square complex matrix with orthonormal columns is unitary
- An  $n \times n$  unitary matrix U satisfies  $U^{-1} = U^*$  and  $||Ux||_2 = ||x||_2$  for any complex vector  $x \in \mathbb{C}^n$

### The eigenvalue problem

- The eigenvalue problem is  $Ax = \lambda x$ , for the eigenvalues  $\lambda$  and eigenvectors x
- It can also be written  $0 = (\lambda I A)x$
- We find eigenvalues  $\lambda_k$  such that  $\lambda_k I A$  is singular, and we find the corresponding eigenvectors  $x_k$  for k = 1, 2, ..., n
- When doing this by hand, one calculates the roots of the characteristic polynomial  $\det(\lambda I A)$
- There are thus n eigenvalues for an  $n \times n$  matrix, counting multiplicity
- Note: eigenvalues and eigenvectors not done that way in computer

### The eigenvalue decomposition

- Suppose we know the eigenvalues  $\lambda_k$  and eigenvectors  $\boldsymbol{v}_k$  so that  $A\boldsymbol{v}_k = \lambda_k \boldsymbol{v}_k$ , for k = 1, 2, ..., n
- For each k, we have a column vector on each side of the equation
- We can assemble each of those n column vectors into a matrix:

### The eigenvalue decomposition

• If we know that there is a complete set of linearly independent (LI) eigenvectors  $v_k$ , then  $V^{-1}$  exists, and we can postmultiply by it to get:

$$A = VDV^{-1}$$

- This is the eigenvalue decomposition, or EVD, of A
- If A is diagonalizable, it will have an EVD
- When does that happen?

### Theorem 7.2.1

If the  $n \times n$  matrix A has n distinct eigenvalues, then A is diagonalizable.

### Computing EVD in Matlab

- We can use the eig command to compute the EVD
- The eigenvalues arrive as the diagonal elements of **D**
- ullet The eigenvectors are the columns of  $oldsymbol{V}$
- A is singular, but has distinct eigenvalues, and because of that, it has a complete set of LI eigenvectors.
   Because of that, V is nonsingular

```
A = pi*ones(2,2);
lambda = eig(A)
lambda = 0
6.2832
```

```
V =
-0.7071  0.7071
0.7071  0.7071
D =
0  0
0  6.2832
```

[V,D] = eig(A)

### Computing EVD in Matlab

• We can easily check that the EVD is equivalent to A:

```
>> norm( A-V*D/V ) % /V is equivalent to *inv(V)

ans =

8.8818e-16

>>
```

What if A is not diagonalizable?

### Computing EVD in Matlab

- If A is not diagonalizable, it will not have all eigenvalues distinct, and it will not have a full set of eigenvectors
- eig still works, but  $V^{-1}$  doesn't exist:
- *V* is only rank 1, indicating a single LI column
- You should get in the habit of critically evaluating output to check whether it is consistent with theory

```
[Example 7.2.1]
```

### The EVD and similarity

- There is an important relationship between A and D
- For any nonsingular matrix S, then  $B = SAS^{-1}$  is said to be similar to A
- In many basic linear algebra texts, one can find a proof of this result:

Theorem 7.2.2 If X is an nonsingular matrix, then  $XAX^{-1}$  has the same eigenvalues as A.

• There is a really nice interpretation of the similarity transformation

Similarity transformations
 Consider the produce of a nonsingular matrix X with any vector:

$$y = Xz = z_1x_1 + \cdots + z_nx_n$$

- The  $x_i$  are the columns of X
- The columns of X are LI because it is invertible, so it is a basis for  $\mathbb{C}^n$
- The  $z_i$  are the coordinates of y using the columns of X as a basis
- But, we could left multiply by  $X^{-1}$  and then  $z = X^{-1}y$
- The elements of y are now coordinates for z using the columns of  $X^{-1}$  as a basis
- Thus: multiplication by the inverse of a matrix performs a change of **basis** into the coordinates associated with the matrix

### Similarity transformations

- Now consider the EVD
- Let u = Ax, or

- Premultiply by to get  $u = Ax = VDV^{-1}x$

$$V^{-1}u = D(V^{-1}x)$$

- ullet This equation says that using the columns of  $oldsymbol{V}$  for a basis, that there is a diagonal relation between the two vectors  $\boldsymbol{u}$  and  $\boldsymbol{x}$
- Said another way, the EVD finds a basis for  $\mathbb{C}^n$  so that the map is diagonal:

$$x \mapsto Ax$$

In that case, each coordinate is just multiplied by its own scalar

### Similarity transformations

- This can be really convenient for matrix powers; these will be an important operation for us
- Consider  $A^2$ , and use the EVD:

$$A^2 = (VDV^{-1})(VDV^{-1}) = VD(V^{-1}V)DV^{-1} = VD^2V^{-1},$$

- If we new the EVD, we could just square the diagonal elements of D, then reconstruct  $A^2$
- Higher powers? Then

$$A^k = VD^kV^{-1}$$

• Raise each eigenvalue to the k-th power to get  $D^{k}$ !

### Conditioning of EVD computation

- There are theorems around to tell us how much eigenvalues change in response to changes in the matrix
- One is the Bauer-Fike theorem:

```
Theorem 7.2.3 Let {\pmb A} \in {\mathbb C}^{n \times n} be diagonalizable, {\pmb A} = {\pmb V} {\pmb D} {\pmb V}^{-1}, with eigenvalues \lambda_1, \dots, \lambda_n. If \mu is an eigenvalue of {\pmb A} + {\pmb E} for a complex matrix {\pmb E}, then \min_{j=1,\dots,n} |\mu - \lambda_j| \le \kappa({\pmb V}) \|{\pmb E}\|, \tag{7.2.6} where \|\cdot\| and \kappa are in the 2-norm.
```

- Eigenvalues by as much as a factor of the condition number  $\kappa(V)$
- If V is unitary,  $\kappa(V) = 1$ , A is normal, eigenvalue calculation robust
- If V nearly singular,  $\kappa(V)\gg 1$ , eigenvalues can change significantly

## From Wikipedia "Bauer-Fite theorem":

Alternate formulation:

Let  $(\lambda^{\alpha}, \nu^{\alpha})$  be an approximate eigenvalue - eigenvector pair and

 $V = A V^a - \lambda^a V^a.$ 

Then there is an eigenvalue  $\lambda$  of A such that

 $|\lambda - \lambda^{\alpha}| \leq \kappa(V) \frac{\|r\|}{\|v^{\alpha}\|}.$ 

# Conditioning of EVD computation

- Example: triangular matrix with n = 15
- The bound on the change in the condition number is around 1e7
- Test the theorem with 1e-7 size perturbations
- The max change in these few cases is about 25% to 50% of the possible change

```
[Example 7.2.2]
```

```
lambda = (1:n)';
A = triu( ones(n,1)*lambda');
The Bauer-Fike theorem provides an upper bound
these eigenvalues.
[V,D] = eig(A);
kappa = cond(V)
kappa =
   7.1978e+07
for k = 1:3
   E = randn(n); E = 1e-7*E/norm(E);
   mu = eig(A+E):
   max_change = norm( sort(mu)-lambda, inf )
end
max_change =
   0.2407
max_change =
   0.4492
max_change =
   0.2737
```

### Method of EVD computation

- The practical methods for computing the EVD are beyond the scope of this class (and book)
- It is worth pointing out that the methods often use the idea of matrix powers as part of the computation.
- ullet If the eigenvalues are distinct, then raising them to a power separates them for large powers k
- There is an easy and elegant way to accomplish this separation

### Method of EVD computation

• Example: create a matrix with known eigenvalues:

Now do QR factorization and reverse it:

$$A=QR$$
 [Q,R] = qr(A);  
 $QTA=R$ 

We have same eigenvalues!

The qr function takes a random 5 by 5 matrix, then returns orthogonal matrix V and upper triangular matrix R

A and D are similar

eig(A)

### Method of EVD computation

- It turns out that we can repeat this and not change the eigenvalues!
- This is Francis QR iteration

```
• For this example:
```

 Look at diagonal elements, and off diagonal elements are getting small

```
[Example 7.2.3]
```

```
for k = 1:15
     [Q,R] = qr(A);
     A = R*Q;
end
A
```

```
-0.1336
-5.9984
                       0.0100
                                 -0.0000
                                              0.0000
-0.1336
            4.9960
                      -0.0491
                                  0.0000
                                             -0.0000
0.0100
           -0.0491
                       4.0024
                                 -0.0001
                                             -0.0000
-0.0000
            0.0000
                      -0.0001
                                  2.0000
                                             -0.0001
0.0000
           -0.0000
                      -0.0000
                                 -0.0001
                                             -1.0000
```

### Section 7.3 Singular value decomposition

### Singular value decomposition

Here is another matrix factorization; it is widely used in many fields:

Theorem 7.3.1

Let  $\mathbf{A} \in \mathbb{C}^{m \times n}$ . Then  $\mathbf{A}$  can be written as

$$A = USV^*, (7.3.1)$$

where  $U \in \mathbb{C}^{m \times m}$  and  $V \in \mathbb{C}^{n \times n}$  are unitary and  $S \in \mathbb{R}^{m \times n}$  is real and diagonal with nonnegative entries. If A is real, then so are U and V (which are then orthogonal matrices).

- This is the singular value decomposition, or SVD
- Cols of U: left singular vectors; cols of V: right singular vectors
- Diagonal elements of S are the singular values  $\sigma_1, ..., \sigma_r, r = \min\{m, n\}$

### Singular value decomposition

• The SVD is

$$A = USV^*$$

- $A \in \mathbb{C}^{m \times n}$  is  $m \times n$ , with complex entries
- Columns of (unitary)  $U \in \mathbb{C}^{m \times m}$ : left singular vectors
- Columns of (unitary)  $V \in \mathbb{C}^{n \times n}$ : right singular vectors
- Diagonal elements of  $S \in \mathbb{R}^{m \times n}$  are the singular values  $\sigma_1, \dots, \sigma_r, r = \min\{m, n\}$
- Usually the ordering of the singular values is

$$\sigma_1 \geq \sigma_2 \geq \cdots \geq \sigma_r \geq 0$$

- $\sigma_1$  is the principal singular value
- $oldsymbol{u}_1$  and  $oldsymbol{v}_1$  are the principal singular vectors

### Singular value decomposition

- What about real-valued A?
- Consider the case with  $A \in \mathbb{R}^{m \times n}$

$$A = USV^T$$

- Columns of orthogonal  $U \in \mathbb{R}^{m \times m}$ : left singular vectors
- Columns of unitary  $V \in \mathbb{R}^{n \times n}$ : right singular vectors
- Diagonal elements of  $S \in \mathbb{R}^{m \times n}$  are the singular values  $\sigma_1 \geq \sigma_2 \geq \cdots \geq \sigma_r \geq 0$ ,  $r = \min\{m, n\}$
- This is still an SVD because it satisfies the requirements:
  - o the first and last matrices are orthogonal,
  - o and the middle matrix is diagonal with non negative entries

### Interpreting the SVD

• Let's rewrite the SVD

$$A = USV^*$$

• As follows (postmultiply by V):

$$AV = US$$

• Considering one column at a time, we have  $Av_k = \sigma_k u_k$ , k = 1, 2, ..., r,  $r = \min\{m, n\}$ 

• This means that each right singular vector  $v_k$  is mapped by A to a scalar multiple  $(\sigma_k)$  of the corresponding left singular vector  $u_k$ 

## Interpreting the SVD: example

• Compute an SVD:

```
>> A = [2 1;3 4; 5 6]
A =
>> [U,S,V] = svd(A)
  -0.2164 0.9497 -0.2265
  -0.5258 -0.3088 -0.7926
  -0.8226 -0.0524 0.5661
S =
   9.4939 0
          0.9303
                0
V =
  -0.6450 0.7642
  -0.7642
         -0.6450
```

• Compare the first column of AV with the first column of U: this should be just  $\sigma_1$ 

```
>> AV = A*V
AV =

-2.0541     0.8834

-4.9917     -0.2872

-7.8101     -0.0488

>> AV(:,1)./U(:,1)

ans =

9.4939

9.4939

9.4939

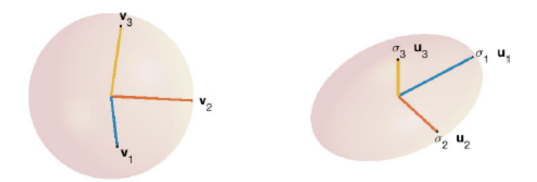
>> S(1,1)

ans =

9.4939
```

# Interpreting the SVD: visualizing it

- The SVD can be visualized for 3x3 real-valued matrices
- Unit vectors in the directions of the columns of V (left) are distorted by multiplication by A (right) to



# Contrasting the EVD and the SVD

Table 7.1. Differences between the EVD and the SVD.

#### EVD

most square matrices

 $Ax_k = \lambda_k x_k$ 

same basis for domain and range of A
may have poor conditioning

#### SVD

all rectangular and square matrices  $Av_k = \sigma_k u_k$ 

two orthogonal bases perfectly conditioned

### Connection between the SVD and 2-norm (Thrm 7.3.2)

- Let  $A \in \mathbb{C}^{m \times n}$  have an SVD with  $A = USV^*$  and  $\sigma_1 \geq \sigma_2 \geq \cdots \geq \sigma_r \geq 0, r = \min\{m, n\}$
- Then:
  - 1. The 2-norm satisfies  $|A|_2 = \sigma_1$ .
  - 2. The rank of A is the number of nonzero singular values.
  - 3. The condition number satisfies

$$\kappa(\mathbf{A}) = ||\mathbf{A}||_2 ||\mathbf{A}^+||_2 = \sigma_1/\sigma_r.$$

Division by zero here implies that A does not have full rank. Recall that  $A^+ = (A^T A)^{-1} A^T$  is the pseudoinverse (chapter 3).

## SVD example 7.3.2: try it

- ➤ A=vander(1:5); % Vandermonde matrix
- A = A(:,1:4) % 5 by 4 now
- $\triangleright$ [U,S,V] = svd(A);
- ➤norm(U'\*U-eye(5)) % check U is orthogonal
- ➤norm(V'\*V-eye(4)) % check V is orthogonal
- ➤ sigma = diag(S)
- $\triangleright$ [norm(A) sigma(1)] % Thrm 7.3.2, no. 1
- $\triangleright$ [cond(A) sigma(1)/sigma(end)] % Thrm 7.3.2, no. 3

#### Connections between SVD and EVD

- Let  $A = USV^*$
- Create the square hermitian matrix  $B = A^*A$
- Then

$$B = (VS^*U^*)(USV^*) = VS^*SV^* = V(S^TS)V^{-1}.$$

- $S^TS$  is a diagonal  $n \times n$  matrix
- There are two cases:

$$S^{T}S = \begin{bmatrix} \sigma_{1}^{2} & & & \\ & \ddots & \\ & & \sigma_{n}^{2} \end{bmatrix}, \qquad S^{T}S = \begin{bmatrix} \sigma_{1}^{2} & & \\ & \ddots & \\ & & \sigma_{m}^{2} & \\ & & & 0 \end{bmatrix},$$

• The lower right zero is  $n - m \times n - m$ 

#### Connections between SVD and EVD

• There are two cases:

$$\underline{m \geq n:} \qquad \underline{m < n:} \\ S^T S = \begin{bmatrix} \sigma_1^2 & & & \\ & \ddots & \\ & & \sigma_n^2 \end{bmatrix}, \qquad S^T S = \begin{bmatrix} \sigma_1^2 & & & \\ & \ddots & & \\ & & \sigma_m^2 & \\ & & & & 0 \end{bmatrix},$$

- The squares of the singular values of A are the eigenvalues of  $B = A^*A!$
- Conversely, the EVD of  ${\it B}$  yields the singular values of  ${\it A}$  and the right singular vectors of  ${\it A}$
- We could get the left singular vectors from AV = US by doing one column at a time

#### Connections between SVD and EVD

• We could also create

which is 
$$m + n \times m + n$$

$$C = \begin{bmatrix} 0 & A^* \\ A & 0 \end{bmatrix}.$$

for  $A m \times n$ 

- If  $\sigma$  is a singular value of B, the  $\sigma$  and  $-\sigma$  are eigenvalues of C
- ullet The associated eigenvector immediately gives a left and right singular vector of  ${\it A}$
- This connection is implicitly exploited by software to compute the SVD

#### Thin form of the SVD

- Like the QR factorization, we can have both full and thin versions of the factorization
- Let  $A = USV^*$  where A is  $m \times n$  with m > n
- Use svd(A,0) to get this in Matlab

# Section 7.4 Symmetry and definiteness

### Symmetry and Definiteness

- When we had real matrices A, we found in chapter 2 that there could be specializations of the factorizations there: e.g.,  $A = LDL^T$ , where D is diagonal and L is lower triangular
- We now study the analogous hermitian case for  $A^* = A$
- Let  $A = USV^*$  where A is  $n \times n$
- Since S is real and square,

$$A^* = VS^*U^* = VSU^*$$

• In this case A is diagonalizable with

$$A = VDV^{-1} = VDV^*$$

- ullet Here  $oldsymbol{V}$  is unitary, and  $oldsymbol{D}$  is diagonal and real
- This means that a hermitian matrix has a complete set of orthonormal eigenvectors (a unitary diagonalization), with real eigenvalues

## Symmetry and Definiteness: example

#### Example 7.4.1

The following matrix is not hermitian.

$$A = [0 \ 2; -2 \ 0]$$

so it is normal.

```
ans = 2.2204e-16
```

The eigenvalues are pure imaginary.

```
lambda = diag(D)
```

```
lambda =
0.0000 + 2.0000i
0.0000 - 2.0000i
```

The singular values are

```
ans = 2 2
```

### Symmetry and Definiteness

- Now we have a theorem that says that the condition number for the eigenvalues is bounded above by  $\kappa(V)$
- Here **V** is the eigenvector matrix
- But, it is unitary (or orthogonal), which means that  $\kappa = 1!$
- So, our last theorem implies that the condition number for a Hermitian or normal matrix is one, which is as good as it gets!
- In that case, that is for Hermitian or normal matrices, the eigenvalues can be changed by no more than the norm of the perturbation of the matrix!
- Can we verify this?

### Symmetry and Definiteness: example

#### Example 7.4.2

We construct a real symmetric matrix with known eigenvalues by using the QR factorization to produce a random orthogonal set of eigenvectors.

```
n = 30;
lambda = (1:n)';
D = diag(lambda);
[V,R] = qr(randn(n));
                         % get a random orthogonal V
A = V*D*V':
```

The orthonormal columns of V become the associated eigenvectors for the eigenvalues in matrix D, which are the same for A. Why?

Eigenvalues are 1,2,...,30 and put on diagonal of matrix D

> The qr function takes a random 30 by 30 matrix here, then returns an orthogonal matrix V and upper triangular matrix R

## Symmetry and Definiteness: example

Example 7.4.2

```
The condition number of these eigenvalues is one. Thus bounded by the norm of the perturbation to A.
```

```
for k = 1:3
    E = randn(n); E = 1e-4*E/norm(E);
    mu = sort(eig(A+E));
    max_change = norm(mu-lambda,inf)
end
```

Create perturbation matrix E that is random and has norm 1e-4

max\_change = 2.5564e-05 max\_change = 2.0501e-05 max\_change =

2.3712e-05

### Symmetry and Definiteness

- This is great, but it is not quite an SVD
- Why? The sign of the diagonal elements could be anything.
- Can we make it into an SVD? Yes!
- The trouble is with D; lets rewrite it as a product of two diagonal matrices: one is T with the sign $(d_{ii})$  on the diagonal; the other is |D|, which has  $|d_{ii}|$  on the diagonal. We still have

$$D = T|D|$$

Substitute to get

$$A = VDV^* = VT|D|V^* = (VT)|D|(V^*)$$

• This is an SVD, because the diagonal matrix has nonnegative entries and the left and right matrices are unitary

## Symmetry and definiteness

- Recall the quadratic form  $x^*Ax$  where A is  $n \times n$  and x is  $n \times 1$ .
- For real entries,  $x^T A x$  is the quadratic form of interest
- For n=2, we can easily write it out:

$$\begin{bmatrix} x_1 & x_2 \end{bmatrix} \begin{bmatrix} a_{11} & a_{12} \\ a_{21} & a_{22} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = a_{11}x_1^2 + (a_{12} + a_{21})x_1x_2 + a_{22}x_2^2$$

- $oldsymbol{\cdot}$  For some matrices, this quantity remains positive for any nonzero vector  $oldsymbol{x}$
- For example, if  $a_{11}>0$ ,  $a_{22}>0$ , and  $a_{21}=-a_{12}$ , then the quadratic form reduces to  $a_{11}x_1^2+a_{22}x_2^2>0$  for nonzero  $\boldsymbol{x}$
- This matrix is called positive definite
- If the matrix were symmetric and this were true, then it is symmetric positive definite or SPD

### Symmetry and definiteness

- The SPD matrix is important: it has all positive eigenvalues
- Complex case: a matrix is Hermitian positive definite (HPD) when

$$A = A^*$$
 and  $x^*Ax > 0$ 

for any nonzero compatible x

• It turns out that we can prove some equivalent statements:

#### Theorem 7.4.3

If  $A^* = A$ , then the following statements are equivalent.

- 1. *A* is HPD.
- 2. The eigenvalues of A are positive numbers.
- 3. Any unitary EVD of A is also an SVD of A.
- This means, e.g., that all positive eigenvalues implies HPD (or SPD)

- In your linear algebra class, you may have discussed the Rayleigh quotient
- For A  $n \times n$  and x  $n \times 1$  with complex entries, the Rayleigh quotient is

$$R_A(x) = \frac{x^*Ax}{x^*x}$$

- In the special case that x is an eigenvector, say v, then  $Av = \lambda v$ , and substitution easily gives that  $R_A(v) = \lambda$
- We can conclude that: the Rayleigh quotient maps an eigenvector into its associated eigenvalue

• Consider the Hermitian case  $A = A^*$  and the Rayleigh quotient

$$R_A(x) = \frac{x^*Ax}{x^*x}$$

- Put in a vector that is close to an eigenvector:  $\mathbf{x} = \mathbf{v} + \epsilon \mathbf{z}$  for  $\epsilon \to 0$
- Using a multidimensional Taylor expansion gives that  $R_A(\boldsymbol{v} + \epsilon \boldsymbol{z}) = R_A(\boldsymbol{v}) + 0 + O(\epsilon^2) = \lambda + O(\epsilon^2)$
- This happens because, for an eigenvector,  $\nabla R_A(v) = \mathbf{0}$
- This means that if the input vector is within  $O(\epsilon)$  of an eigenvector then the Rayleigh quotient is within  $O(\epsilon^2)$  of the eigenvalue:  $R_A$  does a good job of approximating the eigenvalue! Let's explore this...

#### Example 7.4.3

```
We construct a symmetric matrix with a known EVD.
n = 20;
lambda = (1:n)'; D = diag(lambda);
[V,~] = qr(randn(n)); % get a random orthogonal V
A = V*D*V';
```

The Rayleigh quotient of an eigenvector is its eigenvalue.

```
R = @(x) (x'*A*x)/(x'*x);
format long, R(V(:,7))
```

```
ans = 7.000000000000001
```

#### Example 7.4.3

Now let's try different vectors that are closer and closer to an eigenvector

```
delta = 1./10.^(1:4)';
quotient = 0*delta;
for k = 1:4
    e = randn(n,1); e = delta(k)*e/norm(e);
    x = V(:,7)+e;
    quotient(k) = R(x);
end
table(delta, quotient)
```

ans =
delta quotient
----- 7.05738940427937
0.01 7.00066684894918
0.001 7.00000278235035
0.0001 7.00000005557751

Every time the input vector gets a factor of 10 closer to the eigenvector, there is a factor of 100 improvement in the eigenvalue approximation (another two zeros after the decimal point)

# Section 7.5 Dimension reduction

- We now return to the SVD
- We want to use it to approximate the information in a matrix with much less storage, or many less numbers
- We will see a few examples of this
- Consider matrix  $A \in \mathbb{R}^{m \times n}$  with m > n (for now)
- The thin SVD is then

• We truncated the n+1 to m columns of U and rows of S to get  $\widehat{U}$ ,  $\widehat{S}$ 

- Now let's rewrite the thin SVD carefully
- Note that  $\widehat{m{U}}$  has orthonormal columns  $m{u}_i$
- V has orthonormal rows  $\boldsymbol{v}_i^T$

$$A = \widehat{U}\widehat{S}V^T = \begin{bmatrix} u_1 & u_2 & \cdots & u_n \end{bmatrix} \begin{bmatrix} \sigma_1 & & & \\ & \ddots & & \\ & \vdots & \ddots & \\ & v_n^T \end{bmatrix}$$

$$m \times n \text{ here}$$

$$= \begin{bmatrix} \sigma_1 u_1 & \cdots & \sigma_n u_n \end{bmatrix} \begin{bmatrix} v_1^T \\ \vdots \\ v_n^T \end{bmatrix}$$

$$= \sigma_1 u_1 v_1^T + \cdots + \sigma_r u_r v_r^T = \sum_{i=1}^r \sigma_i u_i v_i^T,$$

$$u_i v_i^T \text{ is an outer product here, with each forming an } m \times n \text{ matrix}$$

• Each of those outer products is weighed with the singular value, then adding them *all* up recovers the original matrix

$$\mathbf{A} = \sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T,$$

- But, it so happens that in many matrices, there are only a few singular values that are sizable, and the rest may be quite small.
- By convention we ordered the singular values:

$$\sigma_1 \ge \sigma_2 \ge \cdots \ge \sigma_r \ge 0$$

- So, if the largest few singular values are say 1,2, ..., k, then maybe we need to keep only the first k terms to get a good approximation to the content of A!
- Let's make this more precise.

- We can think of the process of keeping more singular values as generating a sequence of matrices for increasing k, with  $1 \le k \le r$
- ullet Then, we only do a partial sum using the first k terms:

$$A_k = \sum_{i=1}^n \sigma_i u_i v_i^T = U_k S_k V_k^T.$$

- $\boldsymbol{U}_k$  and  $\boldsymbol{V}_k$  are the first k columns of  $\boldsymbol{U}$  and  $\boldsymbol{V}$
- $S_k$  is the upper left  $k \times k$  submatrix of S
- Because each  $u_iv_i^T$  is a matrix with unit norm, the overall size of each matrix added is given by the singular value
- Because of this, there are many cases where stopping at a small value of k relative to r will give a v

- ullet What do we know about the  $oldsymbol{A}_k$  ?
- The rank of a sum of matrices is less than or equal to the sum of the ranks of each: thus, the rank of  $A_k$  is at most k
- And, it turns out that  $A_k$  is the best rank k approximation to A!

#### Theorem 7.5.1

Suppose A has rank r and let  $A = USV^T$  be an SVD. Let  $A_k$  be as in (7.5.2) for  $1 \le k < r$ . Then  $A_k = \sum_{i=1}^k \sigma_i u_i v_i^T = U_k S_k V_k^T.$ 

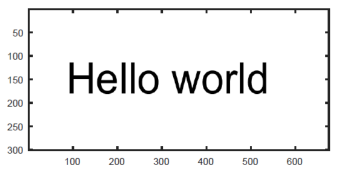
1. 
$$||A - A_k||_2 = \sigma_{k+1}, \qquad k = 1, \dots, r-1.$$

2. If the rank of B is k or less, then  $||A - B||_2 \ge \sigma_{k+1}$ .

- ullet For many matrices, k need not be very large to get a good approximation to the original matrix
- Example 7.5.1 gives an example with text
- Demo of that example and/or others...

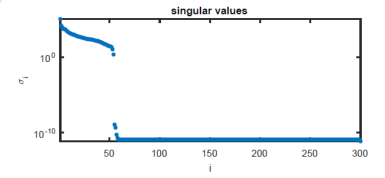
```
tobj = text(0,0,'Hello world','fontsize',44);
saveas(gcf,'hello.png')
A = imread('hello.png');
A = double(rgb2gray(A));
imagesc(A), colormap gray
[m,n] = size(A)
```

```
m = 300
n = 675
```



```
[U,S,V] = svd(A);
sigma = diag(S);
semilogy(sigma,'.')
r = find(sigma/sigma(1) > 10*eps,1,'last')
```

```
r = 56
```



#### Example 7.5.1

rank = 6 rank = 8

Hello world Hello world

Look how few singular values are needed to get a decent looking image! Less than 5% of the original storage to get the rank 8 approximation!

- These low rank approximations can be used to get at the essence of data
- One measure of how much content is contained in each added rank is the fractional "energy" given by

$$\tau_k = \frac{\sum_{i=1}^k \sigma_i^2}{\sum_{i=1}^r \sigma_i^2}, \quad k = 1, \dots, r.$$

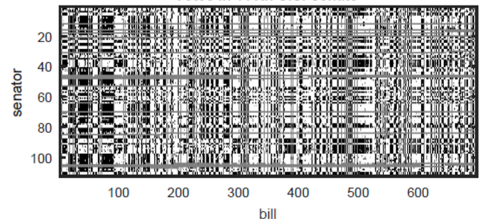
- Consider a different matrix of information now: Example 7.5.2
- Here we look at the voting pattern for the Senate in the 111<sup>th</sup> session of the US Congress

#### Example 7.5.2

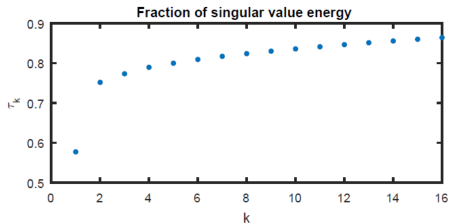
If we visualize the votes (white is "yea," black is "nay," and gray is anything else), we can see great similarity between many rows, reflecting party unity.

```
imagesc(A)
colormap gray
```

#### Votes in 111th U.S. Senate

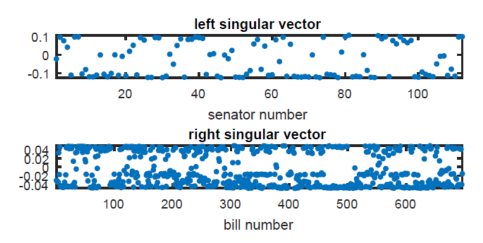


```
[U,S,V] = svd(A);
sigma = diag(S);
tau = cumsum(sigma.^2) / sum(sigma.^2);
plot(tau(1:16),'.')
```

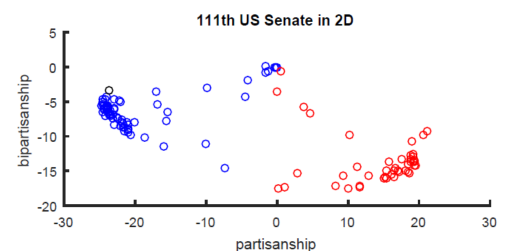


- The first two singular values account for 75% of the energy
- The remaining ones each account for relatively little of the energy
- Maybe just those first two are enough to capture the essence of the data?

```
subplot(211), plot(U(:,1),'.')
subplot(212), plot(V(:,1),'.')
```



- Note the different sizes of the two vectors
- Most of the values in these vectors are at ± C: there is not much in the middle



- Projecting each senator's votes in first two V coordinates, the right singular vectors: (1) is partisanship; (2) is bipartisanship
- Those coordinates are then plotted against each other
- Red: Republican
- Blue: Democrat
- · Black: Independent
- The scatter plot suggests that there is a clear separation between parties!