INTRO TO DATA SCIENCE LECTURE 12: DIMENSIONALITY REDUCTION

RECAP 2

LAST TIME:

- LOGISTIC REGRESSION

AGENDA

I. DIMENSIONALITY REDUCTION
II. PRINCIPAL COMPONENTS ANALYSIS
III. SINGULAR VALUE DECOMPOSITION
IV. OTHER METHODS

EXERCISE:

IV. DIMENSIONALITY REDUCTION IN SCIKIT-LEARN

Q: What is dimensionality reduction?

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- A: A set of techniques for reducing the size (in terms of features, records, and/or bytes) of the dataset under examination.

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In general, the idea is to regard the dataset is a matrix and to decompose the matrix into simpler, meaningful pieces.

Dimensionality reduction is frequently performed as a pre-processing step before another learning algorithm is applied.

Q: What are the motivations for dimensionality reduction?

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The number of features in our dataset can be difficult to manage, or even misleading (eg, if the relationships are actually simpler than they appear).

For example, suppose we have a dataset with some features that are related to each other.

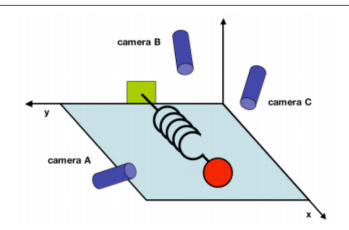
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If these relationships are *linear*, then we can use well-established techniques like PCA/SVD.



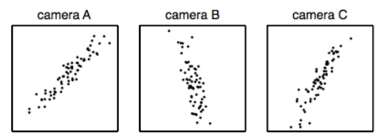


FIG. 1 A toy example. The position of a ball attached to an oscillating spring is recorded using three cameras A, B and C. The position of the ball tracked by each camera is depicted in each panel below.

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(More precisely, the sample size grows exponentially with $l \le d$, the dimension of the manifold *embedded* in the feature space).

Another way of characterizing this is to say that high-dimensional spaces are inherently **sparse**.

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ex: A high-dimensional orange contains most of its volume in the rind!

ex: A high-dimensional hypercube contains most of its volume in the corners!

In either case, most of the points in the space are "far" from the center.

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This illustrates the fact that local methods will break down in these circumstances (eg, in order to collect enough neighbors for a given point, you need to expand the radius of the neighborhood so far that locality is not preserved).

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This illustrates the fact that local methods will break down in these circumstances (eg, in order to collect enough neighbors for a given point, you need to expand the radius of the neighborhood so far that locality is not preserved).

The bottom line is that high-dimensional spaces can be problematic.

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More precisely: given an $n \times d$ matrix A (encoding n observations of a d-dimensional random variable), we want to find a k-dimensional representation of A (k < d) that captures the information in the original data, according to some criterion.

Q: What is the goal of dimensionality reduction?

- reduce computational expense
- reduce susceptibility to overfitting
- reduce noise in the dataset
- enhance our intuition

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feature selection — selecting a subset of features using an external criterion (*filter*) or the learning algo accuracy itself (*wrapper*)

feature extraction — mapping the features to a lower dimensional space

Feature selection is important, but typically when people say dimensionality reduction, they are referring to *feature extraction*.

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The goal of feature extraction is to create a new set of coordinates that *simplify the representation* of the data.

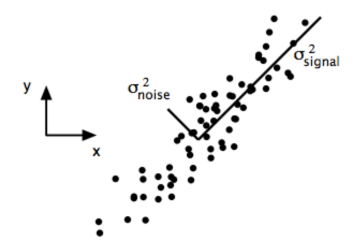


FIG. 2 Simulated data of (x,y) for camera A. The signal and noise variances σ_{signal}^2 and σ_{noise}^2 are graphically represented by the two lines subtending the cloud of data. Note that the largest direction of variance does not lie along the basis of the recording (x_A, y_A) but rather along the best-fit line.

Q: What are some applications of dimensionality reduction?

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- topic models (document clustering)
- image recognition/computer vision
- bioinformatics (microarray analysis)
- speech recognition
- astronomy (spectral data analysis)
- recommender systems

PCs # 0
PCs # 30



PCs # 10



PCs # 40



PCs # 20



PCs # 50



II. PRINCIPAL COMPONENT ANALYSIS

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The PCA of a matrix A boils down to the **eigenvalue decomposition** of the **covariance matrix** of A.

The covariance matrix C of a matrix A is always square:

$$C = \begin{bmatrix} E[(X_1 - \mu_1)(X_1 - \mu_1)] & E[(X_1 - \mu_1)(X_2 - \mu_2)] & \cdots & E[(X_1 - \mu_1)(X_n - \mu_n)] \\ E[(X_2 - \mu_2)(X_1 - \mu_1)] & E[(X_2 - \mu_2)(X_2 - \mu_2)] & \cdots & E[(X_2 - \mu_2)(X_n - \mu_n)] \\ \vdots & \vdots & \ddots & \vdots \\ E[(X_n - \mu_n)(X_1 - \mu_1)] & E[(X_n - \mu_n)(X_2 - \mu_2)] & \cdots & E[(X_n - \mu_n)(X_n - \mu_n)] \end{bmatrix}.$$

off-diagonal elements C_{ij} give the *covariance* between X_i , X_j ($i \neq j$) diagonal elements C_{ii} give the *variance* of X_i

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For an eigenvector ν of A and its eigenvalue λ , we have the important relation:

$$Av = \lambda v$$

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The columns of Q are the eigenvectors of A, and the values of A.

NOTE

This relationship defines what it means to be an eigenvector of

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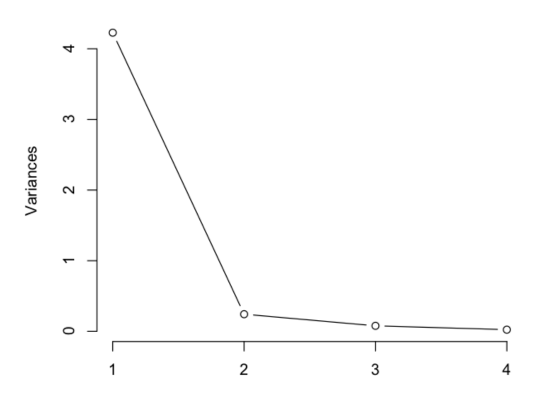
Furthermore the basis elements are ordered by their eigenvalues (from largest to smallest), and these eigenvalues represent the amount of variance explained by each basis element.

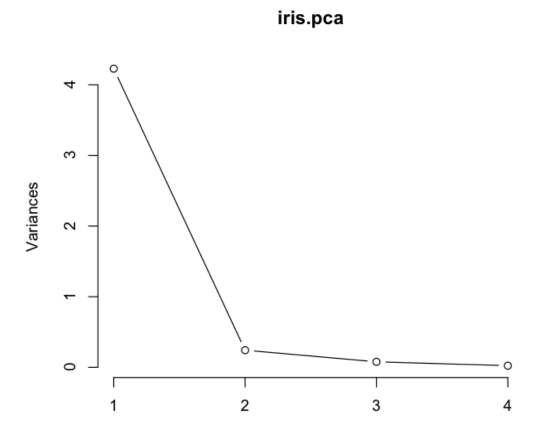
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This can be visualized in a **scree plot**, which shows the amount of variance explained by each basis vector.







NOTE

Looking at this plot also gives you an idea of how many principal components to keep.

Apply the *elbow test*: keep only those pc's that appear to the left of the elbow in the graph.

III. SINGULAR VALUE DECOMPOSITION

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Consider a matrix A with n rows and d features.

The **singular value decomposition** of *A* is given by:

$$A = U \Sigma V^T$$

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$$\rightarrow UU^T = I_n, \ VV^T = I_d \qquad \rightarrow \Sigma_{ij} = 0 \ (i \neq j)$$

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The columns of U & V are the (left- and right-) singular vectors of A.

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These singular vectors provide **orthonormal bases** for the spaces K_n & K_d (columns of U & V, respectively).

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The nonzero entries of Σ are the **singular values** of A. These are real, nonnegative, and rank-ordered (decreasing from left to right).

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NOTE

The number of singular values is equal to the rank of A.

The rank of a matrix measures its *non-degeneracy*.

The nonzero entries of Σ are the **singular values** of A. These are real, nonnegative, and rank-ordered (decreasing from left to right).

For a general SVD, the columns of U are the eigenvectors of AA^T , and the columns of V are the eigenvectors of A^TA .

Also, the singular values of A are the square roots of the eigenvalues of AA^T and A^TA .

SINGULAR VALUE DECOMPOSITION

Q: How do you interpret the SVD?

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Here "best" refers to the representation that minimizes the squared orthogonal distances from the points to the subspace. Q: How do you interpret the SVD?

A: Recall that given a set of n points in d-dimensional space (eg, a matrix A), we want to find the best k < d dimensional subspace to represent the data.

For k = 1, this subspace is a line passing through the origin.

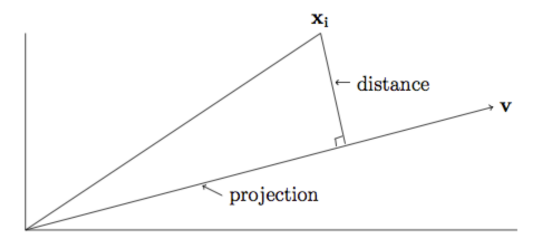


Figure 4.1: The projection of the point $\mathbf{x_i}$ onto the line through the origin in the direction of \mathbf{v}

SINGULAR VALUE DECOMPOSITION

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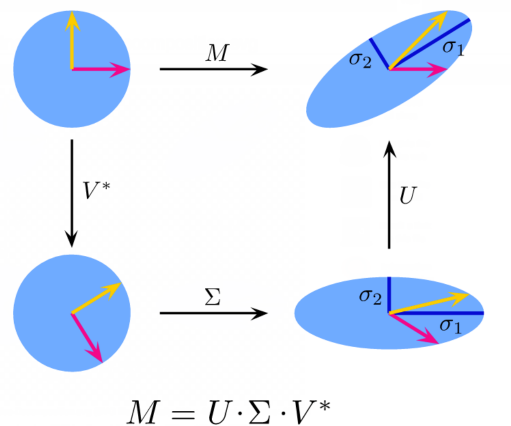
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The singular vectors of *T* correspond to the lengths of the axes of the *d*-dimensional ellipsoid.

The singular values give the magnitudes of the projection of each column of the original dataset on the elements of the new basis.

SINGULAR VALUE DECOMPOSITION



source: http://en.wikipedia.org/wiki/Singular_value_decomposition

III. OTHER METHODS

FACTOR ANALYSIS

Whereas PCA and SVD create new coordinates by transform the old coordinates, **factor analysis** requires new coordinates to be specified externally.

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The old coordinates are then modeled as linear combinations of the latent features.

FACTOR ANALYSIS

For example, consider a dataset that represents the results of a decathalon (rows = participants, columns = events, entries = times).

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Though this dataset contains 10 features X_i , we may be interested in modeling these features as functions of *latent variables* such as the speed and strength of the participants:

$$X_i = \lambda_1 f_1 + \lambda_2 f_2 + \varepsilon$$

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This would allow us to analyze the data in a more fundamental way.

NONLINEAR METHODS

SVD, PCA, and factor analysis are all linear techniques (eg, we use a linear transformation to embed the in a lower-dimensional space).

But sometimes linear techniques are not sufficient.

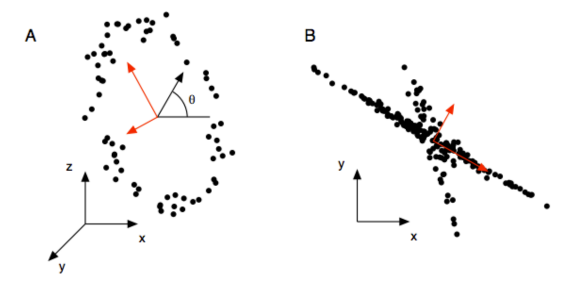
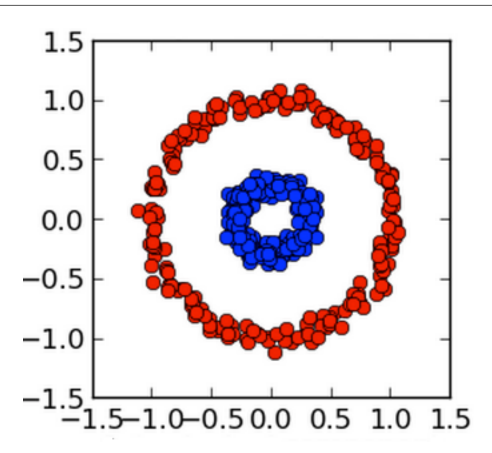


FIG. 6 Example of when PCA fails (red lines). (a) Tracking a person on a ferris wheel (black dots). All dynamics can be described by the phase of the wheel θ , a non-linear combination of the naive basis. (b) In this example data set, non-Gaussian distributed data and non-orthogonal axes causes PCA to fail. The axes with the largest variance do not correspond to the appropriate answer.



NONLINEAR METHODS

Some methods for nonlinear dimensional reduction (or *manifold learning*) include:

multidimensional scaling: low-dim embedding that preserves pairwise distances

locally linear embedding: approximates local structure of data (nbd preserving embedding)

Some methods for nonlinear dimensional reduction (or *manifold learning*) include:

kernel PCA: exploits PCA dependence on inner product (related to SVMs)

isomap: nonlinear dim reduction via MDS using geodesic (surfacebound) distances

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In any case, the key difficulties with dimensionality reduction are time/ space complexity, randomness (eg different results for different runs), and selecting the number of dimensions in the lower-dim subspace.

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Furthermore, there's an obvious (bias/variance) tradeoff between the number of subspace dimensions and the size of approximation error.