Principal Component Analysis

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Introduction to Machine Learning



Dimensionality Reduction

Sometimes we would like to plot some high-dimensional data in two dimensions, e.g. to see if there are clusters in the data.

How should we represent the high-dimensional data in two dimensions?

Idea 1: Project the data to one of the planes spanned by the axes, i.e. plot coordinate *i* versus coordinate *j* of the data points.

Idea 2: Project the data onto the plane spanned by the directions along the largest variance of the data.



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- 6. Principal Component Regression



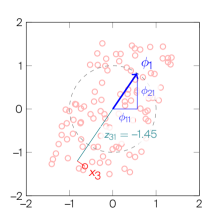
First Principal Component

Given
$$x_{ij}$$
 for $i=1,\ldots,n, j=1,\ldots,p$ column-wise zero mean $\bar{x}_j=\frac{1}{n}\sum_{i=1}^n x_{ij}=0$.

First Principal Component

Find the direction onto which the projection of the data has the highest variance.

(projection) **scores**
$$z_{i1} = \langle \phi_1, x_i \rangle = \sum_{j=1}^p x_{ij} \phi_{j1}$$
 Find **loadings** $\phi_{11}, \phi_{21}, \dots, \phi_{p1}$ that
$$\max_{\phi_{11}, \dots, \phi_{p1}} \frac{1}{n} \sum_{i=1}^n z_{i1}^2$$
 under the constraint $\sum_{i=1}^p \phi_{i1}^2 = 1$.



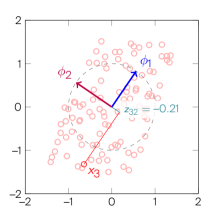


Principal Component Analysis

Second Principal Component

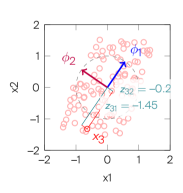
Find the direction onto which the projection of the data has the highest variance under the constraint that it is orthogonal to the first PC.

k-th Principal Component Find the direction onto which the projection of the data has the highest variance under the constraint that it is orthogonal to the first k-1 PCs.

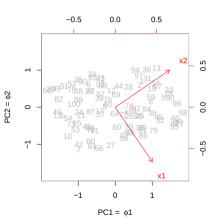


Biplots: Visualizing Scores and Loadings Simultaneously

Plot of raw data



Biplot



scores in gray: read bottom-left coordinates, loadings in red: read top-right coordinates. (read the coordinates of the text label, e.g. x1, not the arrow tip).

First Principal Component Rewritten in Matrix Notation

$$z_1 = X\phi_1$$

data
$$X$$
 $n \times p$ matrix row i contains observation i loadings ϕ_1 $p \times 1$ column vector first PC scores z_1 $n \times 1$ column vector first PC scores of all observation

Find
$$\phi_1$$
 that maximizes $\frac{1}{n}z_1^Tz_1=\max_{\|\phi_1\|=1}\phi_1^TX^TX\phi_1$ $\|\phi_1\|=1$

The solution of this optimization problem is the eigenvector of the largest eigenvalue of the covariance matrix X^TX .



PCA in Matrix Notation and Relation to SVD

$$Z = X\Phi$$

data $X = n \times p$ matrix row i contains observation i

loadings Φ $p \times p$ matrix column j contains PC j

scores $Z n \times p$ matrix column j = scores of PC j for all observations

The columns of the loading matrix Φ are eigenvectors of X^TX , i.e. $X^TX\phi_i = \lambda_i\phi_i$.

PCA is closely linked to **Singular Value Decomposition** $X = U\Sigma V^T$ where U and V are unitary matrices and Σ is a diagonal matrix. One can show that (see exercises)

$$Z = U\Sigma$$
 and $\Phi = V$

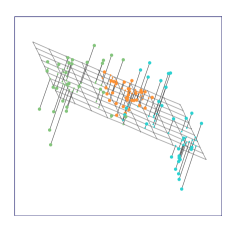


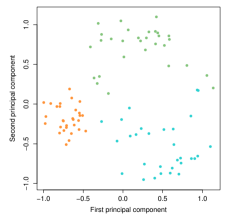
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PCA Provides Linear Subspaces Closest to the Data

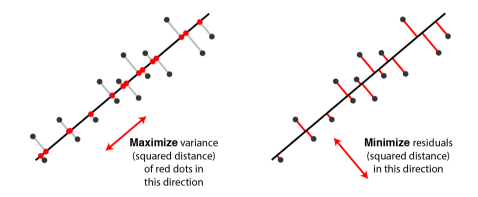




The data varies more within the plane than perpendicular to the plane.



Connection Between the Two Interpretations of PCA





Reinterpretation of Loadings and Scores

Lossless transformation: change of basis

standard basis	${\mathbb I}$	$p \times p$	columns are standard basis vectors
data	X	$n \times p$	rows are coordinates of points in standard basis
loadings	Φ	$p \times p$	columns are new basis vectors
scores	$Z = X\Phi$	$n \times p$	rows are coordinates of points in new basis
reconstruction	$X = Z\Phi^T$	$n \times p$	rows are coordinates of points in standard basis

Lossy transformation: projection to lower-dimensional space

loadings	Φ_L	$p \times L$	$L=1,\ldots,p$ first columns of Φ are new basis vectors		
scores	$Z_L = X\Phi_L$	$n \times L$	rows are coordinates of points in new basis		
reconstruction	$X_L = Z_L \Phi_L^T$	$n \times p$	reconstructed coordinates in standard basis		
$\Phi_{\rm c}$ minimizes $\ \mathbf{V} - \mathbf{V}\Phi_{\rm c}\Phi_{\rm c}^T\ ^2$					





Quiz

Correct or wrong?

- ▶ The scores $z_{11} \approx -1.1$ and $z_{12} \approx 0.1$.
- ► The data has the highest variance in direction $\phi_1 \approx (0.65, -0.55, 0.5)$.
- The two-dimensional linear subspace that is closest to the data is parallel to the X1-X2-plane...

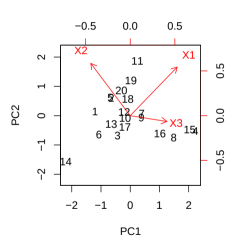




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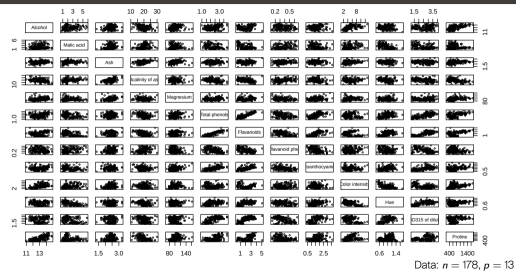


Exploring the Wine Dataset



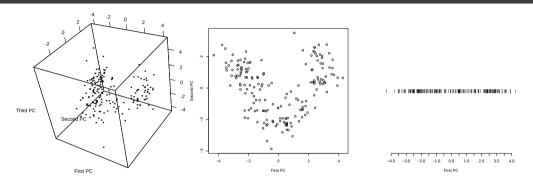
- ▶ 178 different wines from the same region in Italy were chemically analyzed.
- ▶ 13 different measurements per wine: 1) Alcohol 2) Malic acid 3) Ash 4) Alcalinity of ash 5) Magnesium 6) Total phenols 7) Flavanoids 8) Nonflavanoid phenols 9) Proanthocyanins 10) Color intensity 11) Hue 12) OD280/OD315 of diluted wines 13) Proline

Exploring the Wine Dataset





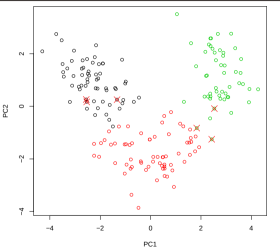
Linear Subspaces in the Wine Data



The 3D plot is obtained by projecting the 13D data onto the linear 3D subspace that is closest to the data. From there onwards we project onto the 2D and 1D subspaces closest to the data to obtain the 2D and 1D plots.



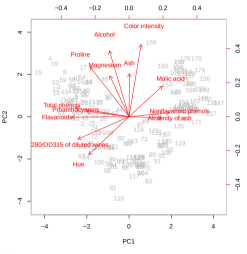
Clustering the Wine Dataset



K-means clustering

- The colors on in the image on the left indicate the classes found by 3-means clustering.
- ➤ The data actually came from wines of 3 different cultivars. The red crosses indicate data points that actually came from the red wine cultivar. All other observations were "correctly" clustered (without having seen a single class label in the training set!)

A Biplot of the Wine Data



PC1, PC2 scores of all observations in grey.

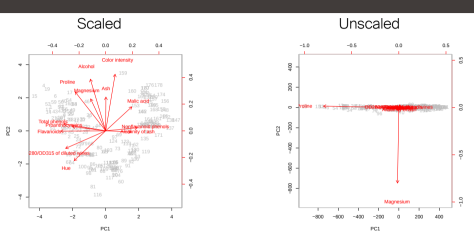
Loadings in red (axis and the top and right).

The first PC corresponds roughly to the total level of phenols (with subtypes Flavanoids, Non-Flavanoids and Proanthocyanins). The Alcalinity of the ash is strongly correlated.

The second PC corresponds roughly to the amount of ash, with e.g. the color intensity strongly correlated.



Scaling Matters



The direction of highest variance may depend on the choice of units (meters versus millimeters). It is common to preprocess each predictor to have mean 0 and variance 1.



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PCA Decorrelates

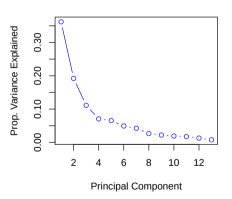
The covariance matrix of the scores is given by the diagonal matrix Λ with the eigenvalues $\lambda_1, \ldots, \lambda_p$ of the covariance matrix X^TX on the diagonal (sorted from largest to smallest).

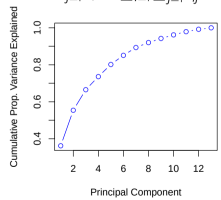
$$Z^TZ = \Phi^TX^TX\Phi = \Phi^T\Phi\Lambda = \Lambda$$
This means $\sum_{i=1}^n z_{im}z_{in} = \left\{ egin{array}{ll} 0 & m
eq n \\ \lambda_m & m = n \end{array}
ight.$



How Many Principal Components Matter?

Proportion of Variance Explained (PVE) of PC m: PVE $_{\rm m} =$



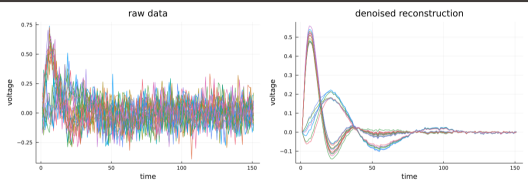


How Many Principal Components Matter?

- ▶ The Proportion of Variance Explained is decreasing, i.e. $PVE_m \ge PVE_{m+1}$.
- Sometimes the first *k* components reach a cumulative proportion of variance explained close to 1 and all the other components do not add much anymore.
- ▶ The question of how many principal components matter is not well defined.
- ▶ In practice, we tend to look at the first few components in order to find interesting patterns in the data.
- ▶ If the signal is along the first few principal components and the noise orthogonal to that, we can use this for compression and denoising.



Denoising



This is an artificial dataset with noise. For the reconstruction we assume that the variance along the first two principal components is the signal and the rest is noise, i.e. for the reconstruction we use L=2 and $X_L=Z_L\Phi_L^T$.



Lossy Image Compression with PCA



A dataset with n=400 grayscale images of faces with $p=64\times64$ pixels. This corresponds to $400\times64^2=1'638'400$ pixel values.

Idea: take each image as a 64 x 64 = 4096 dimensional vector, compute the first few principal components Φ_L and scores Z_L and store the data in this format.

If L=200 is sufficient, we store $4096 \times 200 + 400 \times 200 = 899'200$ values. This corresponds to a compression factor of $\frac{1'638'400}{899'200}$. If needed, we can reconstruct the images $X_L = Z_L \Phi_L^T$.

In general the compression ratio is $\frac{n \times p}{(p+n) \times L}.$

Lossy Image Compression with PCA

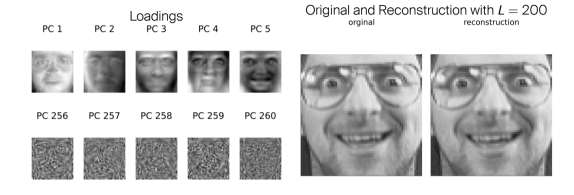




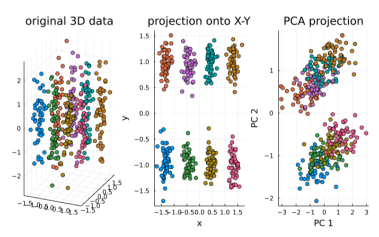
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Limitations of PCA

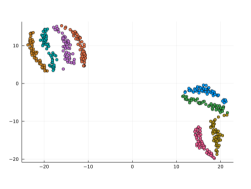
Neighbourhood-relationships can get lost in PCA due to projections.





Alternative 1: t-SNE

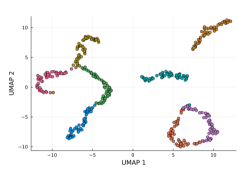
t-Distributed Stochastic Neighbor Embedding (t-SNE)



- t-SNE tries to preserve local neighbourhood-relationships, but the global structure disappears.
- t-SNE is a gradient descent procedure to find low-dimensional coordinates with the property that the probabilities of being neighbours in the low-dimensional space roughly match the probabilities of being neighbours in the high-dimensional space.
- Details can be found here: https://lvdmaaten.github.io/tsne and https://distill.pub/2016/misread-tsne

Alternative 2: UMAP

Uniform Manifold Approximation and Projection (UMAP)

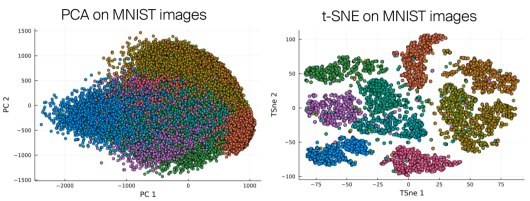


- ► UMAP also tries to preserve local neighbourhood-relationships. but the global structure disappears.
- It is usually faster than t-SNE.
- Details can be found here: https://github.com/lmcinnes/umap



Limitations of PCA

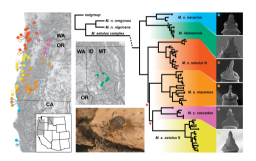
Also in real data there may be clusters that cannot be seen in the first few PCs.





Species Delimitation with Unsupervised Machine Learning

Species Delimitation based on SNP data, dimensionality reduction and clustering: if different methods find the same clusters, we can be somewhat confident that the individual clusters are meaningful.



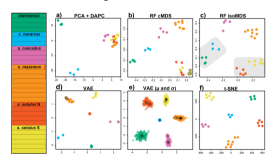




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Principal Component Regression

Unsupervised Feature Learning

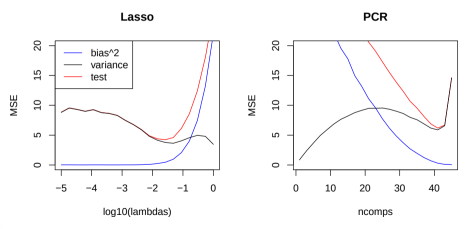
PCA can be used to reduce the dimensionality of the data before supervised learning.

Instead of fitting
$$y_i = \beta_0 + \beta_1 x_{i1} + \cdots + \beta_p x_{ip}$$
 fit
$$y_i = \beta_0 + \beta_1 z_{i1} + \cdots + \beta_m z_{im}$$
 with $m < p$.

In this case we can choose m with cross-validation.

Choosing the Number of PCs in PCR

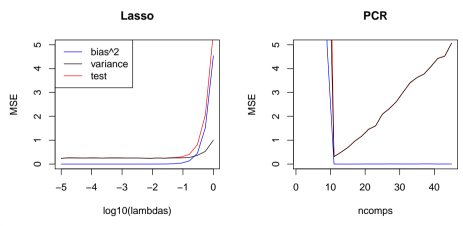
Choosing a small number of PCs in PCR acts as a regularizer that reduces the variance but may increase the bias





PCR versus Ridge Regression and the Lasso

Typically the Lasso (L1) or Ridge Regression (L2) are at least as good as PCR.





Summary

- Principal component analysis can be seen as
 - 1. finding the directions of largest variance in the data,
 - 2. finding linear subspaces that are closest to the data.
- ► The scores and loadings of PCA can be computed with Singular Value Decomposition or by finding the eigenvectors of the covariance matrix.
- Data with different units should be rescaled before applying PCA.
- ► The covariance matrix of the scores is diagonal.
- The (cumulative) proportion of variance explained can be used to see roughly how many PCs matter.
- ▶ PCA can fail to represent clusters faithfully; t-SNE may be more useful in this case.
- ▶ PCA can be used to reduce the dimensionality of a supervised learning problem, with the number of PCs acting as a regularization tuning parameter.

