Dimension Reduction and Feature Selection

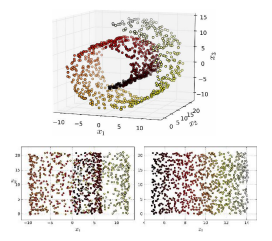
Elliott Ash

Text Data Course, Bocconi 2018

Dimensionality Reduction

- ► Especially in the case of text data, machine learning problems often involve thousands of features.
- Dimension reduction methods are needed, not just for computational tractability, but also to help find a good solution.
- Dimension reduction is also needed for data visualization for example, to plot data in two dimensions.

Swiss role dataset example

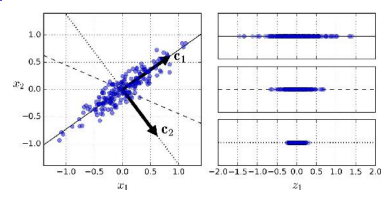


► The dimension reduction process matters: projecting down to two dimensions directly (left panel) might not isolate the variation we are interested in (as done in the right panel, which unrolls the Swiss Roll)

Manifold Learning

- ▶ The swiss roll is an example of a 2D manifold:
 - like many real-world data sets, the data are not uniformly distributed across the space.
 - can be modeled in a lower-dimensional subspace while retaining most of the information
- Dimension reduction methods in machine learning are motivated by this "manifold hypothesis."

PCA



- ▶ PCA (principal components analysis), a popular dimension reduction technique.
 - identifies the axis that accounts for the largest amount of variance in the training set.
 - finds a second axis, orthogonal to the first, that accounts for the largest amount of the remaining variance, and so on
- ▶ The unit vector defining the *i*th axis is called the *i*th principal component.

PCA Projection

▶ For principal components $c_1, c_2, ..., c_n$, define

$$V_n = \begin{bmatrix} \vdots & \vdots & & \vdots \\ c_1 & c_2 & \dots & c_n \\ \vdots & \vdots & & \vdots \end{bmatrix}$$

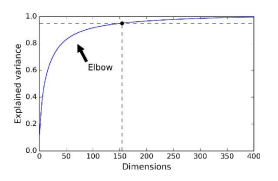
▶ Dimension reduction works by projecting the data set down to the hyperplane defined by the first *d* principal components.

$$\boldsymbol{X}_{PCA} = \boldsymbol{X} \cdot \boldsymbol{V}_d$$

where the d subscript specifies that we have used the first d columns of \boldsymbol{V}_n

```
from sklearn.decomposition import PCA
pca = PCA(n_components=2,svd_solver='randomized')
Xpca = pca.fit_transform(X)
pca.explained_variance_ratio_
plt.scatter(Xpca[:,0],Xpca[:,1], alpha=.1)
```

Choosing the number of dimensions



```
\begin{array}{lll} pca &=& PCA(\,n\_components\,=.95) \\ X95 &=& pca.\,fit\_transform\,(X) \\ pca.\,n\_components\_ \end{array}
```

PCA Inverse Transform

$$X_{\text{recovered}} = X_{\text{reduced}} \cdot V_d$$

Incremental PCA and numpy memmap

- Standard PCA requires you to load the whole data set into memory.
- numpy memmap loads big arrays from disk as needed

```
X_mm = np.memmap('X.pkl', shape = (32567, 525))
```

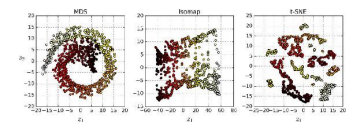
Incremental PCA splits the data into mini-batches and trains gradually.

```
\label{eq:from_sklearn.decomposition} \begin{array}{ll} \textbf{from} & \textbf{sklearn.decomposition} & \textbf{import} & \textbf{IncrementalPCA} \\ \textbf{inc\_pca} & = & \textbf{IncrementalPCA} \big( \begin{array}{ll} \textbf{n\_components} = 100, \\ \textbf{batch\_size} = 1000 \big) \\ \textbf{inc\_pca.fit} \big( \textbf{X\_mm} \big) \end{array}
```

Pros and Cons of PCA

- Advantages:
 - fast to compute
 - good performance on many tasks in practice
 - components are orthogonal by construction
- Disadvantages
 - ▶ lose (potentially a lot of) predictive information from X
 - Coefficients are not easily interpretable.

Other Dimension Reduction Techniques



- Multidimensional Scaling (MDS) reduces dimensionality while trying to preserve the distances between the instances.
- ▶ Isomap creates a graph by connecting each instance to its nearest neighbors, then reduces dimensionality while trying to preserve the geodesic distances9 between the instances.
- t-Distributed Stochastic Neighbor Embedding (t-SNE) reduces dimensionality while trying to keep similar instances close and dissimilar instances apart. Useful for visualizing clusters of instances in high-dimensional space

Dimension Reduction for Visualization

```
from sklearn.manifold import MDS, Isomap, TSNE
mds = MDS()
iso = Isomap()
tsne = TSNE()

# these will be slow
Xmds = mds.fit_transform(X)
Xiso = iso.fit_transform(X)
Xtsne = tsne.fit_transform(X)
```

Principal Components Regression

- ▶ The "classic" way to deal with high-dimensionality.
 - ► Take the first few principal components of *X* and use those as predictors
 - Popular in macroeconomics and finance.

Partial Least Squares

- Partial Least Squares is another technique to determine linear combinations of the predictive variables.
- Unlike PCA, the PLS technique works by successively extracting factors from both predictive and target variables such that covariance between the extracted factors is maximized.
- Outline:
 - Assume X is a $n \times p$ matrix and Y is an $n \times q$ matrix (Y can include a multivariate response variable when q > 1).
 - ▶ PLS successively extracts factors from both *X* and *Y* such that covariance between the extracted factors is maximized.

PLS Goal

▶ Find a linear decomposition of X and Y such that

►
$$X = TPT + E$$

► T , X-scores $(n \times r)$
► P , X-loadings $(p \times r)$
► E , X-residuals $(n \times p)$
► $Y = UQT + F$
► U , Y-scores $(n \times r)$
► Q , Y-loadings $(q \times r)$
► F , Y-residuals $(n \times q)$

Covariance between T and U is maximized.

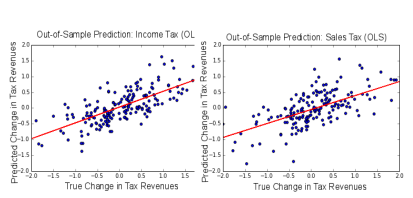
PLS Algorithm

- First step:
 - first extracted x-score is t = Xw, where w is the eigen vector corresponding to the first eigen value of X'YY'X.
 - first y-score is u = Yc, where c is eigen vector corresponding to the first eigen value of Y'XX'Y.
 - ▶ note that X'Y = Y'X = Cov[X, Y]
- Next step:
 - Residualize original values of X and Y as $X_1 = X tt'X$ and $Y_1 = Y tt'Y$
 - ▶ Repeat first step on X_1 and Y_1 to get second components.
 - etc...

PLS with Scikit-learn

```
\begin{array}{lll} \textbf{from} & \textbf{sklearn.cross\_decomposition} & \textbf{import} & \textbf{PLSRegression} \\ \textbf{pls} & = & \textbf{PLSRegression} \big( \textbf{n\_components} {=} 10 \big) \\ \textbf{Xpls} & , & \textbf{Ypls} & = & \textbf{pls.fit\_transform} \big( \textbf{X}, \textbf{Y} \big) \end{array}
```

Ash 2016: PLS predictions of tax revenue changes using tax code text

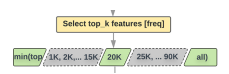


Sales Tax

Weak predictors filtered out; 80% training, 20% testing sample.

Income Tax

- ▶ Predicted change in revenue (vertical axis), plotted against true change in revenue (horizontal axis).
- Correlations between truth and prediction: 0.89 and 0.84.



► The Google text classification guide recommends a maximum 20,000 items in your feature set.

Feature selection using L1 Penalty

A popular way to reduce dimensionality is to run lasso or elastic net, and exclude any predictors whose weights are regularized to zero.

```
\begin{array}{lll} \textbf{from} & \textbf{sklearn.linear\_model import} & \textbf{ElasticNet} \\ \textbf{enet\_reg} & = & \textbf{ElasticNet} ( \textbf{alpha} = .1, \ \textbf{l1\_ratio} = .5) \\ \textbf{enet\_reg.fit} ( \textbf{X}, \textbf{Y}) \\ \textbf{nonzero} & = & \textbf{enet\_reg.coef\_} & != \ 0 \\ \textbf{X\_enet} & = & \textbf{X}[:, \texttt{nonzero}] \end{array}
```

Feature selection using univariate comparisions

- $\sim \chi^2$ is a very fast feature selection routine for classification tasks
 - features must be non-negative
 - works on sparse matrices

- With negative predictors, use f_classif.
- For regression tasks, use f_regression.
- There are also mutual_info_regression and mutual_info_classif, which will recover non-linear relations between predictor and outcome
 - they are much slower, can run on a sample of the data.

Graph-based feature selection

- Graph-based feature selection works by clustering collinear features and then iteratively removing features.
 - much slower than univariate comparisons, but probably gives better performance in text classification tasks.