Lecture 8

Scalable PCA/SVD Dimensionality Reduction & Factor Analysis

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COM6012 Scalable Machine Learning Spring 2018

Week 8 Contents

• Unsupervised Learning

• PCA - Dimensionality Reduction

• SVD – Factor Analysis

• Scalable PCA in Spark

Unsupervised Learning

Supervised methods

$$y = f(X)$$

predict our data



as a function of other data

Unsupervised methods



find structure in the data on its own

Three Topics

- Principal component analysis (PCA) & SVD
 - Dimensionality reduction & factor analysis
- K-means
 - Clustering
- Matrix factorisation (with missing information)
 - Collaborative filtering → Recommender system
- Scale these algorithms for big data

Week 8 Contents

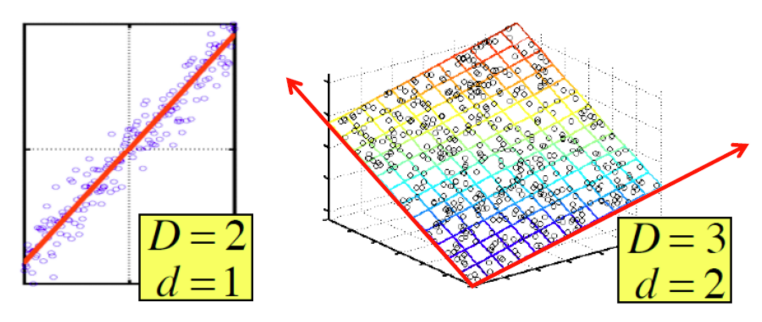
Unsupervised Learning

• PCA - Dimensionality Reduction

• SVD – Factor Analysis

• Scalable PCA in Spark

Dimensionality Reduction



- **Assumption:** Data lies on or near a low *d*-dimensional subspace
- Axes of this subspace are effective representation of the data

Why Reduce Dimensions?

Why reduce dimensions?

- Discover hidden correlations/topics
 - Words that occur commonly together
- Remove redundant and noisy features
 - Not all words are useful
- Interpretation and visualization
- Easier storage and processing of the data

Dimensionality Reduction

Raw data is complex and high-dimensional

• Dimensionality reduction describes the data using a simpler, more compact representation

• This representation may make interesting patterns in the data clearer or easier to see

Dimensionality Reduction

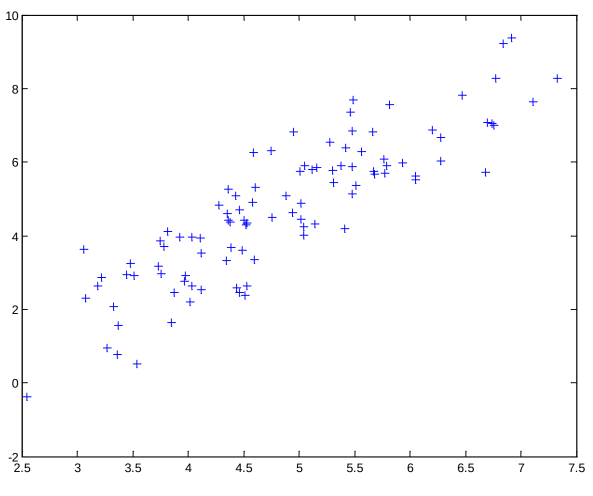
• Goal: Find a 'better' representation for data

- How do we define 'better'?
- For example
 - Minimise reconstruction error
 - Maximise variance
 - They give the same solution \rightarrow PCA!

PCA Algorithm

- Input: N data points, each \rightarrow D-dimensional vector
- PCA algorithm
 - 1. $\mathbf{X_0} \leftarrow \text{Form } N \times D \text{ data matrix, with one row vector } \mathbf{x_n}$ per data point
 - 2. **X**: subtract mean **x** from each row vector \mathbf{x}_n in \mathbf{X}_0
 - 3. $\Sigma \leftarrow \mathbf{X}^T\mathbf{X}$ Gramian (scatter) matrix for \mathbf{X}
 - ullet Find eigenvectors and eigenvalues of Σ
 - PCs U $(D \times d)$ \leftarrow the d eigenvectors with largest eigenvalues
- PCA feature for y D-dim: U^Ty (d-dimensional)
 - Zero correlations, ordered by variance

2D Data

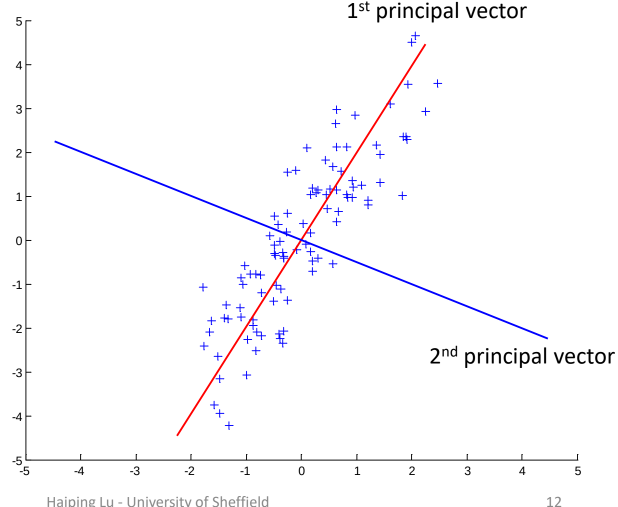


13/04/2018

Haiping Lu - University of Sheffield

Principal Components

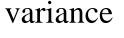
- The best axis to project
- Minimum RMS error
- Principal vectors are orthogonal

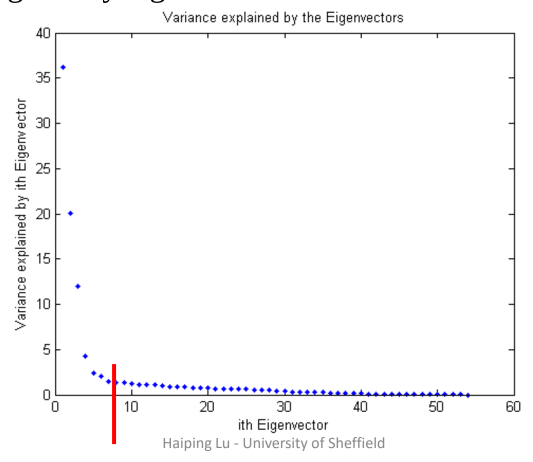


How Many Components?

• Check the distribution of eigen-values

• Take enough many eigen-vectors to cover 80-90% of the





13/04/2018

Other Practical Tips

- PCA assumptions (linearity, orthogonality) not always appropriate
- Various extensions to PCA with different underlying assumptions, e.g., manifold learning, Kernel PCA, ICA
- Centring is crucial, i.e., we must preprocess data so that all features have zero mean before applying PCA
- PCA results dependent on scaling of data
- Data is sometimes rescaled in practice before applying PCA

Problems and Limitations

- What if very large dimensional data?
 - e.g., Images (D $\geq 10^4 = 100 \times 100$)
- Problem:
 - Gramian matrix Σ is size (D²)
 - D= $10^4 \rightarrow |\Sigma| = 10^8$
- Singular Value Decomposition (SVD)!
 - Efficient algorithms available
 - Some implementations find just top d eigenvectors

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Unsupervised Learning

• PCA - Dimensionality Reduction

• SVD – Factor Analysis

• Scalable PCA in Spark

Singular Value Decomposition

- Factorization (decomposition) problem
 - #1: Find concepts/topics/genres → Factor Analysis
 - #2: Reduce dimensionality

\mathbf{term}	data	information	retrieval	brain	lung
document					
CS-TR1	1	1	1	0	0
CS-TR2	2	2	2	0	0
CS-TR3	1	1	1	0	0
CS-TR4	5	5	5	0	0
MED-TR1	0	0	0	2	2
${f MED-TR2}$	0	0	0	3	3
MED-TR3	0	0	0	1	1

The above matrix is actually "2-dimensional." All rows can be reconstructed by scaling [1 1 1 0 0] or [0 0 0 1 1]: D=5→d=2

SVD - Definition

$$\mathbf{A}_{[\mathbf{n} \times \mathbf{m}]} = \mathbf{U}_{[\mathbf{n} \times \mathbf{r}]} \, \mathbf{\Lambda}_{[\mathbf{r} \times \mathbf{r}]} \, (\mathbf{V}_{[\mathbf{m} \times \mathbf{r}]})^{\mathrm{T}}$$

- A: $n \times m$ matrix (e.g., n documents, m terms)
- U: $n \times r$ matrix (n documents, r concepts)
- Λ : $r \times r$ diagonal matrix (strength of each 'concept') (r: rank of the matrix)
- V: $m \times r$ matrix (m terms, r concepts)

SVD - Properties

Always possible to decompose matrix \mathbf{A} into $\mathbf{A} = \mathbf{U} \mathbf{\Lambda} \mathbf{V}^{T}$, where

- **U**, **Λ**, **V**: unique (*)
- U, V: column orthonormal (i.e., columns are unit vectors, orthogonal to each other)
 - $U^TU = I$; $V^TV = I$ (I: identity matrix)
- A: singular value are positive, and sorted in decreasing order

SVD ←→Eigen-decomposition

- SVD gives us:
 - $\mathbf{A} = \mathbf{U} \boldsymbol{\Lambda} \mathbf{V}^{\mathrm{T}}$
- Eigen-decomposition:
 - $\mathbf{B} = \mathbf{W} \mathbf{\Sigma} \mathbf{W}^{\mathrm{T}}$
 - \mathbf{U} , \mathbf{V} , \mathbf{W} are orthonormal ($\mathbf{U}^{\mathrm{T}}\mathbf{U}=\mathbf{I}$),
 - Λ , Σ are diagonal
- Relationship:
 - $AA^T = U \Lambda V^T (U \Lambda V^T)^T = U \Lambda V^T (V \Lambda^T U^T) = U \Lambda \Lambda^T U^T$
 - $A^TA = V \Lambda^T U^T (U \Lambda V^T) = V \Lambda \Lambda^T V^T = V \Lambda^2 V^T$
 - B= $A^TA=W \Sigma W^T$

SVD for PCA

• PCA by SVD:

- 1. $\mathbf{X}_0 \leftarrow$ Form $N \times d$ data matrix, with one row vector \mathbf{x}_n per data point
- 2. **X** subtract mean **x** from each row vector \mathbf{x}_n in \mathbf{X}_0
- 3. U Λ V^T \leftarrow SVD of X
- The right singular vectors \mathbf{V} of \mathbf{X} are equivalent to the eigenvectors of $\mathbf{X}^T\mathbf{X} \rightarrow$ the PCs
- The singular values in Λ are equal to the square roots of the eigenvalues of $\mathbf{X}^T\mathbf{X}$

SVD - Properties

'spectral decomposition' of the matrix:

$$\begin{bmatrix} 1 & 1 & 1 & 0 & 0 \\ 2 & 2 & 2 & 0 & 0 \\ 1 & 1 & 1 & 0 & 0 \\ 5 & 5 & 5 & 0 & 0 \\ 0 & 0 & 0 & 2 & 2 \\ 0 & 0 & 0 & 3 & 3 \\ 0 & 0 & 0 & 1 & 1 \end{bmatrix} = \begin{bmatrix} \begin{vmatrix} 1 & 1 & 1 & 0 & 0 \\ 0 & 1 & 1 & 0 & 0 \\ 0 & 0 & 0 & 2 & 2 \\ 0 & 0 & 0 & 1 & 1 \end{bmatrix} \times \begin{bmatrix} \lambda_1 & \emptyset \\ 0 & \lambda_2 \end{bmatrix} \times \begin{bmatrix} \lambda_1 & 0 & 0 \\ 0 & \lambda_2 \end{bmatrix} \times \begin{bmatrix} \lambda_1 & 0 & 0 \\ 0 & \lambda_2 \end{bmatrix}$$

SVD - Interpretation

'documents', 'terms' and 'concepts':

- U: document-to-concept similarity matrix
- V: term-to-concept similarity matrix
- Λ : its diagonal elements: 'strength' of each concept

Projection:

• Best axis to project on: ('best' = min sum of squares of projection errors)

• $\mathbf{A} = \mathbf{U} \mathbf{\Lambda} \mathbf{V}^{\mathrm{T}}$ - example:

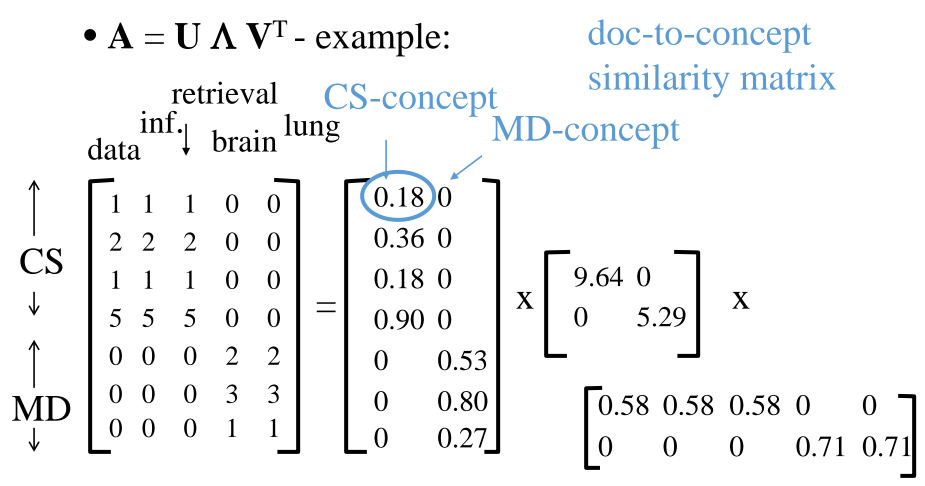
retrieval inf.↓ brain lung

\mathbf{term}	data	information	retrieval	brain	lung
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CS-TR1	1	1	1	0	0
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MED-TR3	0	0	0	1	1

	1	1	1	0	0
$\frac{1}{\text{CS}}$	2	2	2	0	0
	1	1	1	0	0
V	5	5	5	0	0
\uparrow	0	0	0	2	2
$\stackrel{ }{ m MD}$	0	0	0	3	3
↓	0	0	0	1	1_

$$\begin{bmatrix}
9.04 & 0 \\
0 & 5.29
\end{bmatrix}$$

$$\begin{bmatrix}
0.58 & 0.58 & 0.58 & 0 & 0 \\
0 & 0 & 0 & 0.71 & 0.71
\end{bmatrix}$$



• $\mathbf{A} = \mathbf{U} \mathbf{\Lambda} \mathbf{V}^{\mathrm{T}}$ - example:

retrieval
$$\inf_{\text{data}} \downarrow \text{ brain}$$
 brain lung 'strength' of CS-concept

$$\uparrow \quad \begin{bmatrix}
1 & 1 & 1 & 0 & 0 \\
2 & 2 & 2 & 0 & 0 \\
1 & 1 & 1 & 0 & 0 \\
5 & 5 & 5 & 0 & 0 \\
0 & 0 & 0 & 2 & 2 \\
0 & 0 & 0 & 3 & 3 \\
0 & 0 & 0 & 1 & 1
\end{bmatrix} = \begin{bmatrix}
0.18 & 0 \\
0.36 & 0 \\
0.18 & 0 \\
0.90 & 0 \\
0 & 0.53 \\
0 & 0.80 \\
0 & 0.27
\end{bmatrix} \times \begin{bmatrix}
9.64 & 0 \\
0 & 5.29
\end{bmatrix} \times \begin{bmatrix}
0.58 & 0.58 & 0.58 & 0 & 0 \\
0 & 0 & 0.71 & 0.71
\end{bmatrix}$$

• $\mathbf{A} = \mathbf{U} \mathbf{\Lambda} \mathbf{V}^{\mathrm{T}}$ - example:

retrieval similarity matrix data 0.18 0 **CS-concept** 0.36 0 0.18 0 \mathbf{X} 0.90 0 0.53 0.80 0.58 0.58 0.58 0

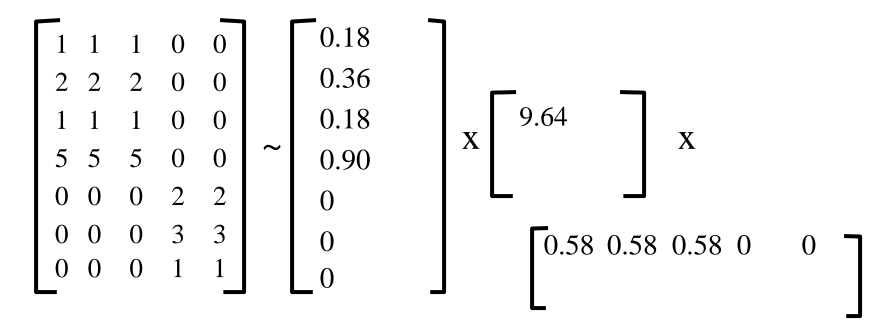
term-to-concept

SVD – Dimensionality Reduction

- Q: how exactly is (**further**) dim. reduction done?
- A: set the smallest singular values to zero:
- Note: 3 zero singular values already removed

$\begin{bmatrix} 1 & 1 & 1 & 0 & 0 \\ 2 & 2 & 2 & 0 & 0 \\ 1 & 1 & 1 & 0 & 0 \\ 5 & 5 & 5 & 0 & 0 \\ 0 & 0 & 0 & 2 & 2 \end{bmatrix} =$	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	
0 0 0 3 3 0 0 0 1 1	$\begin{bmatrix} 0 & 0.80 \\ 0 & 0.27 \end{bmatrix} \qquad \begin{bmatrix} 0.58 & 0.58 & 0.58 & 0 & 0 \\ 0 & 0 & 0 & 0.71 & 0 \end{bmatrix}$	71

SVD - Dimensionality Reduction



SVD - Dimensionality Reduction

• Best rank-1 approximation

Γ_1	1	1	0	0		1 1
	2	2	0	0		2 2
$\begin{array}{c c} 2 \\ 1 \end{array}$	1	1	0	0		1 1
	5	5	0	0	~	5 5
$\begin{bmatrix} 0 \\ 0 \end{bmatrix}$	0	0	2	2		0 0
$\begin{bmatrix} 0 \\ 0 \end{bmatrix}$	0	0	3	3		0 0
0	0	0	1	1		0 0
L	•	•	_		,	

```
      1
      1
      1
      0
      0

      2
      2
      2
      0
      0

      1
      1
      1
      0
      0

      5
      5
      5
      0
      0

      0
      0
      0
      0
      0

      0
      0
      0
      0
      0

      0
      0
      0
      0
      0

      0
      0
      0
      0
      0
```

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PCA - Dimensionality Reduction

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PCA & SVD in Spark MLlib

- Not scalable: computePrincipalComponents() from RowMatrix
- Scalable: computeSVD() from RowMatrix
- Code:

https://github.com/apache/spark/blob/v2.1.0/mllib/src/main/scala/org/apache/spark/mllib/linalg/distributed/RowMatrix.scala

• Documentation:

https://spark.apache.org/docs/2.1.0/api/scala/index.html#org.apache.spark.mllib.linalg.distributed.RowMatrix

PCA in Spark MLlib (RDD)

• https://spark.apache.org/docs/2.1.0/mllib-dimensionality-reduction.html

```
val mat: RowMatrix = new RowMatrix(dataRDD)

// Compute the top 4 principal components.

// Principal components are stored in a local dense matrix.

val pc: Matrix = mat.computePrincipalComponents(4)
```

• Not scalable, local computation

```
val brzSvd.SVD(u: BDM[Double], s: BDV[Double], _) = brzSvd(Cov)
```

Notebook 8

PCA in Spark ML (DF)

• Now in

https://spark.apache.org/docs/2.1.0/ml-features.html#pca

- Under features
- Scalable? Not likely

```
val pca = new PCA()
    .setInputCol("features")
    .setOutputCol("pcaFeatures")
    .setK(3)
    .fit(df)
```

SVD in Spark MLlib (RDD)

- https://spark.apache.org/docs/2.1.0/mllib-dimensionality-reduction.html
- With distributed implementations

```
val mat: RowMatrix = new RowMatrix(dataRDD)

// Compute the top 5 singular values and corresponding singular vectors.
val svd: SingularValueDecomposition[RowMatrix, Matrix] = mat.computeSVD(5, computeU = true)
val U: RowMatrix = svd.U // The U factor is a RowMatrix.
val s: Vector = svd.s // The singular values are stored in a local dense vector.
val V: Matrix = svd.V // The V factor is a local dense matrix.
```

SVD in Spark MLlib (RDD)

- An $m \times n$ data matrix **A** with m > n (note different notations)
- For large matrices, usually we don't need the complete factorization but only the top *k* singular values and its associated singular vectors.
- Save storage, de-noise and recover the low-rank structure of the matrix (dimensionality reduction)

SVD in Spark MLlib (RDD)

- An $m \times n$ data matrix A
- Assume m>n, SVD $\mathbf{A} = \mathbf{U} \Lambda \mathbf{V}^{\mathrm{T}}$
- The singular values and the right singular vectors are derived from the eigenvalues and the eigenvectors of $\mathbf{A}^{T}\mathbf{A}$ (which is smaller than \mathbf{A})
- The left singular vectors are computed via matrix multiplication as $\mathbf{U}=\mathbf{A}\mathbf{V}\ \mathbf{\Lambda}^{-1}$, if requested by the user via the computeU parameter

- Auto
- If *n* is small (n<100) or *k* is large compared with *n* (k>n/2), compute $\mathbf{A}^{\mathrm{T}}\mathbf{A}$ first and then compute its top eigenvalues and eigenvectors **locally** on the driver
- Otherwise, compute A^TA v in a distributive way and send it to ARPACK to compute the top eigenvalues and eigenvectors on the driver node

• Auto (default)

```
if (n < 100 || (k > n / 2 && n <= 15000)) {
    // If n is small or k is large compared with n, we better compute the Gramian matrix first
    // and then compute its eigenvalues locally, instead of making multiple passes.
    if (k < n / 3) {
        SVDMode.LocalARPACK
    } else {
        SVDMode.LocalLAPACK
    }
} else {
        // If k is small compared with n, we use ARPACK with distributed multiplication.
        SVDMode.DistARPACK
}</pre>
```

Specify computeMode (private)

```
case "local-svd" => SVDMode.LocalLAPACK
case "local-eigs" => SVDMode.LocalARPACK
case "dist-eigs" => SVDMode.DistARPACK
```

computeMode (note brzSvd.SVD is local)

```
// Compute the eigen-decomposition of A' * A.
val (sigmaSquares: BDV[Double], u: BDM[Double]) = computeMode match {
  case SVDMode.LocalARPACK =>
    require(k < n, s"k must be smaller than n in local-eigs mode but got k=$k and n=$n.")
   val G = computeGramianMatrix().asBreeze.asInstanceOf[BDM[Double]]
    EigenValueDecomposition.symmetricEigs(v => G * v, n, k, tol, maxIter)
  case SVDMode.LocalLAPACK =>
   // breeze (v0.10) svd latent constraint, 7 * n * n + 4 * n < Int.MaxValue
    require(n < 17515, s"$n exceeds the breeze svd capability")</pre>
   val G = computeGramianMatrix().asBreeze.asInstanceOf[BDM[Double]]
   val brzSvd.SVD(uFull: BDM[Double], sigmaSquaresFull: BDV[Double], _) = brzSvd(G)
    (sigmaSquaresFull, uFull)
  case SVDMode.DistARPACK =>
    if (rows.getStorageLevel == StorageLevel.NONE) {
     logWarning("The input data is not directly cached, which may hurt performance if its"
        + " parent RDDs are also uncached.")
    require(k < n, s"k must be smaller than n in dist-eigs mode but got k=k = n.")
    EigenValueDecomposition.symmetricEigs(multiplyGramianMatrixBy, n, k, tol, maxIter)
```

Remark

- Acknowledgement
 - Some slides are adapted from slides by Jure Leskovec et al. http://www.mmds.org
- References
 - http://infolab.stanford.edu/~ullman/mmds/ch11.pdf
 - http://www.mmds.org
 - https://en.wikipedia.org/wiki/Principal_component_analysis
 - https://spark.apache.org/docs/2.1.0/mllib-dimensionality-reduction.html