CS5489 - Machine Learning

Lecture 6c - Non-Linear Dimensionality Reduction

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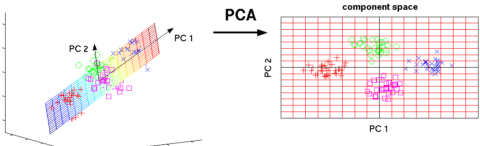
Outline

- 1. Linear Dimensionality Reduction for Vectors
- 2. Linear Dimensionality Reduction for Text
- 3. Non-linear Dimensionality Reduction
- 4. Manifold Embedding

Linear Dimensionality Reduction

- PCA, NMF, LSA are all linear dimensionality reduction methods
 - model the data as "living" on a linear manifold (line, plane, etc).

original data space



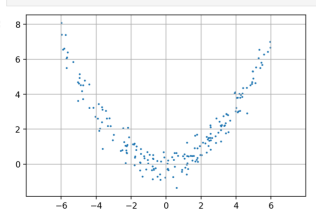
Non-linear surface

• What if the data "lives" on a non-flat surface?

In [4]: pfig



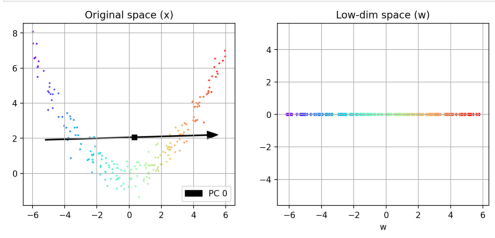
Out[4]:



- PCA can't capture the curvature of the data
 - purple points are close together

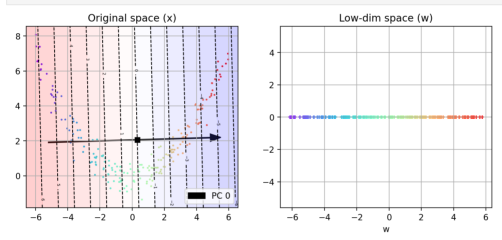
```
In [5]: pca = decomposition.PCA(n_components=1)
W = pca.fit_transform(X)

plt.figure(figsize=(10,4))
plot_basis(X, pca.components_, Y=Y, showlowarrow=False)
```



- · iso-contours of PCA projection
 - points on the same dashed line are projected to the same PCA coefficient.

```
In [6]: plt.figure(figsize=(10,4))
    plot_basis(X, pca.components_, Y=Y, showcontours=True, pca=pca, showlowarrow=False)
```



Kernel PCA

- How to project to a non-linear surface?
 - apply a high-dimensional feature transformation to the data

$$\circ \mathbf{x}_i \Rightarrow \phi(\mathbf{x}_i)$$

- project high-dim data to a linear surface
 - \circ i.e. run PCA on $\phi(\mathbf{x}_i)$
- in the original space, the projection will be non-linear

Kernel principal components

- ullet a principal component ${f v}$ is a linear combination of high-dim vectors
 - $\mathbf{v} = \sum_{i=1}^n a_i \phi(\mathbf{x}_i)$
 - where a_i are learned weights for component ${f v}$.
- For a new point \mathbf{x}_* , the KPCA coefficient for \mathbf{v} is

•
$$w = \phi(\mathbf{x}_*)^T \mathbf{v} = \sum_{i=1}^n a_i \phi(\mathbf{x}_*)^T \phi(\mathbf{x}_i) = \sum_{i=1}^n a_i k(\mathbf{x}_*, \mathbf{x}_i) = \mathbf{k}_*^T \mathbf{a}_i$$

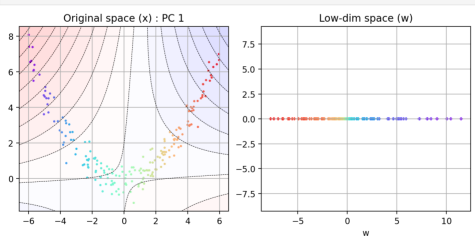
- coefficient is based on similarity to data points belonging to v.
- using the kernel trick saves computation.

Learning KPCA weights

- Apply PCA to the transformed data $\{\phi(\mathbf{x}_i)\}$:
- After some derivation:
 - 1. Calculate the kernel matrix $\mathbf{K} = [k(\mathbf{x}_i, \mathbf{x}_i)]_{ij}$
 - 2. center the kernel (remove the mean in the high-dim space: $\tilde{\mathbf{K}} = (\mathbf{I} \frac{1}{N}\mathbf{1}\mathbf{1}^T)\mathbf{K}(\mathbf{I} \frac{1}{N}\mathbf{1}\mathbf{1}^T)$ $\mathbf{1}\mathbf{1}^T$ is a matrix of ones.
 - lacksquare 3. Find the top-K eigenvector/value pairs: $ilde{\mathbf{K}}\mathbf{a}_j=\lambda_j\mathbf{a}_j$
 - 4. Scale (normalize PC in high-dim space): $\mathbf{a}_j \leftarrow rac{1}{\sqrt{\lambda_j}} \mathbf{a}_j$
 - 5. Project new data \mathbf{x}_* : $w_i = \mathbf{k}_*^T \mathbf{a}_i$
- Example using polynomial kernel
 - purple points are further apart.
 - PC coefficient corresponds to location along the data curve.

```
In [8]: # run KPCA
kpca = decomposition.KernelPCA(n_components=1, kernel='poly', gamma=0.15, degree=2, coef0=0)
W = kpca.fit_transform(X)

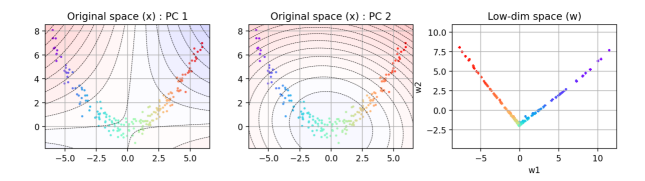
plt.figure(figsize=(8,4))
plot_kpca(X, W, kpca, showcontours=True, Y=Y)
```



- Example: 2 PCs
 - 2nd PC corresponds to the distance from the center

```
In [9]: # run KPCA
kpca = decomposition.KernelPCA(n_components=2, kernel='poly', gamma=0.15, degree=2, coef0=0)
W = kpca.fit_transform(X)

plt.figure(figsize=(10,3))
plot_kpca(X, W, kpca, showcontours=True, Y=Y)
```



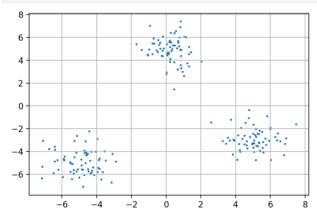
RBF kernel

- · principal components separate the data into clusters
- · coefficient is distance to clusters

Example

• data with 3 clusters

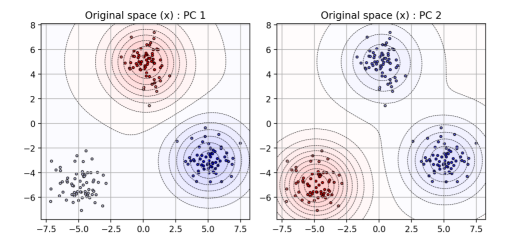
```
In [11]: plt.scatter(X[:,0], X[:,1], s=6, edgecolor="none")
  plt.grid(True);
```



- The first 2 PCs can split the data into 3 clusters
 - the color of the datapoint corresponds to the coefficient value.

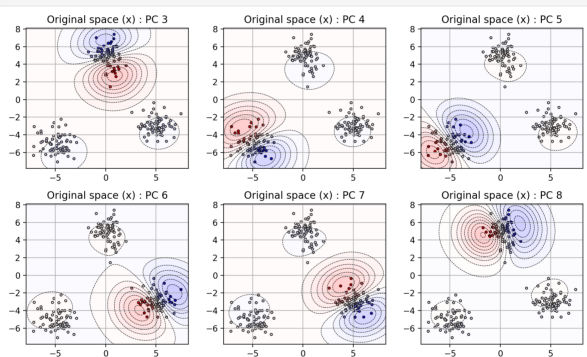
```
In [12]: # run KPCA
   kpca = decomposition.KernelPCA(n_components=8, kernel='rbf', gamma=0.15)
   W = kpca.fit_transform(X)

plt.figure(figsize=(8,4))
   plot_kpca(X, W, kpca, showcontours=True, showpcs=[0,1], colorcoefs=True)
```



- The remaining 6 PCs split each cluster into halves
 - multiple splits in orthogonal directions

```
In [13]: plt.figure(figsize=(10,6))
   plot_kpca(X, W, kpca, showcontours=True, showpcs=range(2,8), colorcoefs=True)
```

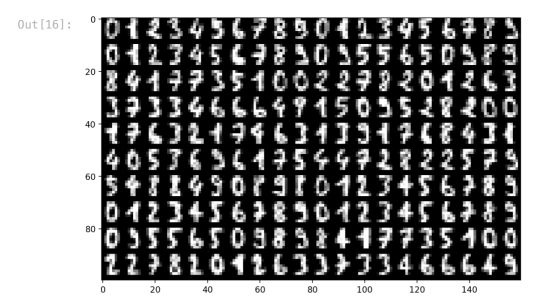


Example on digit images

• 8 x 8 images -> 64D vector

```
In [14]: digits = datasets.load_digits(n_class=10)
X = digits.data
Y = digits.target

# randomly split data into training and testing
trainX, testX, trainY, testY = \
model_selection.train_test_split(X, Y,
train_size=0.8, test_size=0.2, random_state=4487)
In [16]: digitfig
```



- · Apply KPCA with RBF kernel
 - (parallelize with n_jobs)

```
In [17]: kpca = decomposition.KernelPCA(n_components=10, kernel='rbf', gamma=0.001, n_jobs=-1)
    trainW = kpca.fit_transform(trainX)
```

- Top-5 positive and negative prototypes for each PC
 - the number is the a_i value for that image.
 - from the prototypes, the PCs are modeling the differences in appearance between digits

```
In [21]: plt.figure(figsize=(8,10))
  plot_kbasis(kpca, (8,8), trainX)
```

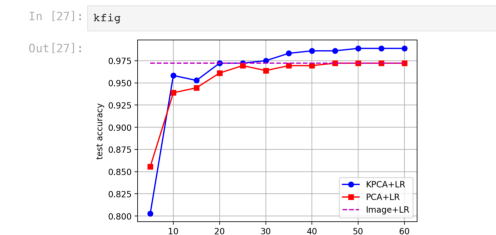


Classification experiment

- use KPCA coefficients as the new representation
 - train a logistic regression classifier
 - try different numbers of components
 - Note: can do this efficiently by selecting a subset of KPCA components.

```
In [22]: # apply kernel PCA
         kpca = decomposition.KernelPCA(n_components=60, kernel='rbf', gamma=0.001, n_jobs=-1)
         trainW = kpca.fit transform(trainX)
         testW = kpca.transform(testX)
In [23]: ncs = [5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60]
         for nc in ncs:
             # extract the first nc PCs
             trainWnew = trainW[:,0:nc]
             testWnew = testW[:,0:nc]
             # train classifier
             logreg = linear_model.LogisticRegressionCV(Cs=logspace(-4,4,10), cv=5, n_jobs=-1, max_it
             logreg.fit(trainWnew, trainY)
             # test classifier
             predYtest = logreg.predict(testWnew)
                        = metrics.accuracy_score(testY, predYtest)
             accs.append(acc)
```

- · Classification results on test set
 - KPCA can improve the performance, compared with PCA and raw image.



KPCA Summary

• Use kernel trick to perform PCA in high-dimensional space.

number of components

- Coefficients are based on a non-linear projection of the data.
- The type of projection is based on the kernel function selected.
- Using RBF kernel, KPCA can split the data into clusters.