### Machine Learning Techniques

(機器學習技法)



Lecture 14: Radial Basis Function Network

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### Roadmap

- 1 Embedding Numerous Features: Kernel Models
- 2 Combining Predictive Features: Aggregation Models
- Oistilling Implicit Features: Extraction Models

#### Lecture 13: Deep Learning

pre-training with denoising autoencoder (non-linear PCA) and fine-tuning with backprop for NNet with many layers

#### Lecture 14: Radial Basis Function Network

- RBF Network Hypothesis
- RBF Network Learning
- k-Means Algorithm
- k-Means and RBF Network in Action

#### Gaussian SVM Revisited

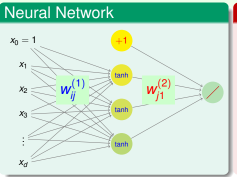
$$g_{\text{SVM}}(\mathbf{x}) = \text{sign}\left(\sum_{\text{SV}} \alpha_n y_n \exp\left(-\gamma \|\mathbf{x} - \mathbf{x}_n\|^2\right) + b\right)$$

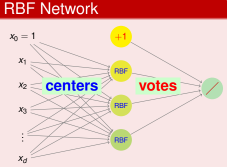
Gaussian SVM: find  $\alpha_n$  to combine Gaussians centered at  $\mathbf{x}_n$ ; achieve large margin in infinite-dimensional space, remember? :-)

- Gaussian kernel: also called Radial Basis Function (RBF) kernel
  - radial: only depends on distance between x and 'center' x<sub>n</sub>
  - · basis function: to be 'combined'
- let  $g_n(\mathbf{x}) = y_n \exp(-\gamma \|\mathbf{x} \mathbf{x}_n\|^2)$ :  $g_{\text{SVM}}(\mathbf{x}) = \text{sign}(\sum_{\text{SV}} \alpha_n g_n(\mathbf{x}) + b)$ 
  - —linear aggregation of selected radial hypotheses

Radial Basis Function (RBF) Network: linear aggregation of radial hypotheses

#### From Neural Network to RBF Network





- hidden layer different:
   (inner-product + tanh) versus (distance + Gaussian)
- output layer same: just linear aggregation

RBF Network: historically a type of NNet

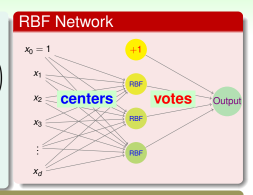
# **RBF Network Hypothesis**

$$h(\mathbf{x})$$

= Output 
$$\left(\sum_{m=1}^{M} \beta_m RBF(\mathbf{x}, \mu_m) + b\right)$$

key variables:

centers  $\mu_m$ ; (signed) votes  $\beta_m$ 



#### $g_{\text{SVM}}$ for Gaussian-SVM

- RBF: Gaussian; Output: sign (binary classification)
- M = #SV;  $\mu_m$ : SVM SVs  $\mathbf{x}_m$ ;  $\beta_m$ :  $\alpha_m y_m$  from SVM Dual

learning: given RBF and Output, decide  $\mu_m$  and  $\beta_m$ 

# **RBF** and Similarity

general similarity function between **x** and **x**':

Neuron(
$$\mathbf{x}, \mathbf{x}'$$
) = tanh( $\gamma \mathbf{x}^T \mathbf{x}' + 1$ )  
DNASim( $\mathbf{x}, \mathbf{x}'$ ) = EditDistance( $\mathbf{x}, \mathbf{x}'$ )

kernel: similarity via  $\mathbb{Z}$ -space inner product —governed by Mercer's condition, remember? :-) Poly( $\mathbf{x}, \mathbf{x}'$ ) =  $(1 + \mathbf{x}^T \mathbf{x}')^2$ 

$$Gaussian(\mathbf{x}, \mathbf{x}') = \exp(-\gamma ||\mathbf{x} - \mathbf{x}'||^2)$$

Truncated(
$$\mathbf{x}, \mathbf{x}'$$
) =  $[\|\mathbf{x} - \mathbf{x}'\| \le 1] (1 - \|\mathbf{x} - \mathbf{x}'\|)^2$ 

RBF: similarity via  $\mathcal{X}\text{-space distance}$ 

—often monotonically non-increasing to distance

RBF Network: distance similarity-to-centers as feature transform

#### Fun Time

Which of the following is not a radial basis function?

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Which of the following is not a radial basis function?

**3** 
$$\phi(\mathbf{x}, \mu) = [\![ \mathbf{x} = \mu ]\!]$$

## Reference Answer: (4)

Note that (3) is an extreme case of (1) (Gaussian) with  $\gamma \to \infty$ , and (2) contains an

 $\|\mathbf{x} - \boldsymbol{\mu}\|^2$  somewhere :-).

#### Full RBF Network

$$h(\mathbf{x}) = \text{Output}\left(\sum_{m=1}^{M} \frac{\beta_m}{\beta_m} \text{RBF}(\mathbf{x}, \mu_m)\right)$$

- full RBF Network: M = N and each  $\mu_m = \mathbf{x}_m$
- physical meaning: each  $x_m$  influences similar x by  $\beta_m$
- e.g. uniform influence with  $\beta_m = 1 \cdot y_m$  for binary classification

$$g_{\text{uniform}}(\mathbf{x}) = \text{sign}\left(\sum_{m=1}^{N} y_m \exp\left(-\gamma \|\mathbf{x} - \mathbf{x}_m\|^2\right)\right)$$

—aggregate each example's opinion subject to similarity

full RBF Network: lazy way to decide  $\mu_m$ 

### Nearest Neighbor

$$g_{\text{uniform}}(\mathbf{x}) = \operatorname{sign}\left(\sum_{m=1}^{N} y_{m} \operatorname{exp}\left(-\gamma \|\mathbf{x} - \mathbf{x}_{m}\|^{2}\right)\right)$$

- $\exp(-\gamma \|\mathbf{x} \mathbf{x}_m\|^2)$ : maximum when  $\mathbf{x}$  closest to  $\mathbf{x}_m$  —maximum one often dominates the  $\sum_{m=1}^{N}$  term
- take y<sub>m</sub> of maximum exp(...) instead of voting of all y<sub>m</sub>
   —selection instead of aggregation
- · physical meaning:

$$g_{\text{nbor}}(\mathbf{x}) = \mathbf{y}_{m}$$
 such that  $\mathbf{x}$  closest to  $\mathbf{x}_{m}$ 

- -called nearest neighbor model
- can uniformly aggregate k neighbors also: k nearest neighbor

k nearest neighbor:also lazy but very intuitive

### Interpolation by Full RBF Network

full RBF Network for squared error regression:

$$h(\mathbf{x}) = \text{Output}\left(\sum_{m=1}^{N} \beta_{m} \text{RBF}(\mathbf{x}, \mathbf{x}_{m})\right)$$

just linear regression on RBF-transformed data

$$\mathbf{z}_n = [\mathsf{RBF}(\mathbf{x}_n, \mathbf{x}_1), \mathsf{RBF}(\mathbf{x}_n, \mathbf{x}_2), \dots, \mathsf{RBF}(\mathbf{x}_n, \mathbf{x}_N)]$$

- optimal  $\beta$ ?  $\beta = (Z^TZ)^{-1}Z^Ty$ , if  $Z^TZ$  invertible, remember? :-)
- size of Z? N (examples) by N (centers)
   —symmetric square matrix
- theoretical fact: if  $\mathbf{x}_n$  all different, Z with Gaussian RBF invertible

optimal  $\beta$  with invertible Z:  $\beta = Z^{-1}y$ 

### Regularized Full RBF Network

full Gaussian RBF Network for regression:  $\beta = Z^{-1}y$ 

$$g_{\text{RBF}}(\mathbf{x}_1) = \boldsymbol{\beta}^T \mathbf{z}_1 = \mathbf{y}^T \mathbf{Z}^{-1} \text{ (first column of Z)} = \mathbf{y}^T \begin{bmatrix} 1 & 0 & \dots & 0 \end{bmatrix}^T = y_1$$
  
 $-g_{\text{RBF}}(\mathbf{x}_n) = y_n, \text{ i.e. } E_{\text{in}}(g_{\text{RBF}}) = 0, \text{ yeah!! :-})$ 

- called exact interpolation for function approximation
- but overfitting for learning? :-(
- how about regularization? e.g. ridge regression for β instead
   —optimal β = (Z<sup>T</sup>Z + λI)<sup>-1</sup>Z<sup>T</sup>y
- seen  $\mathbb{Z}$ ?  $\mathbb{Z} = [\mathsf{Gaussian}(\mathbf{x}_n, \mathbf{x}_m)] = \mathsf{Gaussian}$  kernel matrix  $\mathbb{K}$

effect of regularization in different spaces:

kernel ridge regression: 
$$\boldsymbol{\beta} = (K + \lambda I)^{-1} \mathbf{y}$$
; regularized full RBFNet:  $\boldsymbol{\beta} = (Z^T Z + \lambda I)^{-1} Z^T \mathbf{y}$ 

# Fewer Centers as Regularization

recall:

$$g_{\text{SVM}}(\mathbf{x}) = \text{sign}\left(\sum_{\text{SV}} \alpha_{m} y_{m} \exp\left(-\gamma \|\mathbf{x} - \mathbf{x}_{m}\|^{2}\right) + b\right)$$

—only '≪ N' SVs needed in 'network'

- next: M 

  N instead of M = N
- effect: regularization

by constraining number of centers and voting weights

• physical meaning of centers  $\mu_m$ : prototypes

remaining question: how to extract prototypes?

#### Fun Time

If  $\mathbf{x}_1 = \mathbf{x}_2$ , what happens in the Z matrix of full Gaussian RBF network?

- 1 the first two rows of the matrix are the same
- the first two columns of the matrix are different
- the matrix is invertible
- 4 the sub-matrix at the intersection of the first two rows and the first two columns contains a constant of 0

#### Fun Time

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- 3 the matrix is invertible
- 4 the sub-matrix at the intersection of the first two rows and the first two columns contains a constant of 0

# Reference Answer: (1)

It is easy to see that the first two rows must be the same; so must the first two columns. The two same rows makes the matrix singular; the sub-matrix in 4 contains a constant of  $1 = \exp(-0)$  instead of 0.

#### Good Prototypes: Clustering Problem

if  $\mathbf{x}_1 \approx \mathbf{x}_2$ ,

- $\implies$  no need both RBF( $\mathbf{x}, \mathbf{x}_1$ ) & RBF( $\mathbf{x}, \mathbf{x}_2$ ) in RBFNet,
- $\Longrightarrow$  cluster  $\mathbf{x}_1$  and  $\mathbf{x}_2$  by one prototype  $\mu \approx \mathbf{x}_1 pprox \mathbf{x}_2$ 
  - clustering with prototype:
    - partition  $\{x_n\}$  to disjoint sets  $S_1, S_2, \dots, S_M$
    - choose  $\mu_m$  for each  $S_m$
    - —hope:  $\mathbf{x}_1, \mathbf{x}_2$  both  $\in S_m \Leftrightarrow \mu_m \approx \mathbf{x}_1 \approx x_2$
  - cluster error with squared error measure:

$$E_{\text{in}}(S_1, \dots, S_M; \mu_1, \dots, \mu_M) = \frac{1}{N} \sum_{n=1}^{N} \sum_{m=1}^{M} [[\mathbf{x}_n \in S_m]] |[\mathbf{x}_n - \mu_m]|^2$$

goal: with 
$$S_1, \dots, S_M$$
 being a partition of  $\{x_n\}$ ,

$$\min_{\{S_1,\cdots,S_M;\mu_1,\cdots,\mu_M\}} E_{\text{in}}(S_1,\cdots,S_M;\mu_1,\cdots,\mu_M)$$

### Partition Optimization

with  $S_1, \dots, S_M$  being a partition of  $\{\mathbf{x}_n\}$ ,

$$\min_{\left\{\boldsymbol{S}_{1},\cdots,\boldsymbol{S}_{M}:\mu_{1},\cdots,\mu_{M}\right\}}\sum_{n=1}^{N}\sum_{m=1}^{M}\left[\!\left[\boldsymbol{\mathbf{x}}_{n}\in\boldsymbol{S}_{m}\right]\!\right]\left\|\boldsymbol{\mathbf{x}}_{n}-\boldsymbol{\mu}_{m}\right\|^{2}$$

- hard to optimize: joint combinatorial-numerical optimization
- two sets of variables: will optimize alternatingly

if  $\mu_1, \dots, \mu_M$  fixed, for each  $\mathbf{x}_n$ 

- $[\mathbf{x}_n \in S_m]$ : choose one and only one subset
- $\|\mathbf{x}_n \boldsymbol{\mu}_m\|^2$ : distance to each prototype

optimal chosen subset  $S_m$  = the one with minimum  $\|\mathbf{x}_n - \boldsymbol{\mu}_m\|^2$ 

for given  $\mu_1, \cdots, \mu_M$ , each  $\mathbf{x}_n$  'optimally partitioned' using its closest  $\mu_m$ 

# **Prototype Optimization**

with  $S_1, \dots, S_M$  being a partition of  $\{x_n\}$ ,

$$\min_{\{\boldsymbol{S}_1,\cdots,\boldsymbol{S}_M:\mu_1,\cdots,\mu_M\}} \sum_{n=1}^N \sum_{m=1}^M [\![\boldsymbol{x}_n \in \boldsymbol{S}_m]\!] \|\boldsymbol{x}_n - \boldsymbol{\mu}_m\|^2$$

- hard to optimize: joint combinatorial-numerical optimization
- two sets of variables: will optimize alternatingly

if  $\mathcal{S}_1,\cdots,\mathcal{S}_M$  fixed, just unconstrained optimization for each  $\mu_m$ 

$$\nabla_{\boldsymbol{\mu}_{m}} E_{\text{in}} = -2 \sum_{n=1}^{N} [\mathbf{x}_{n} \in \mathbf{S}_{m}] (\mathbf{x}_{n} - \boldsymbol{\mu}_{m}) = -2 \left( \left( \sum_{\mathbf{x}_{n} \in \mathbf{S}_{m}} \mathbf{x}_{n} \right) - |\mathbf{S}_{m}| \boldsymbol{\mu}_{m} \right)$$

optimal prototype  $\mu_m = \text{average of } \mathbf{x}_n \text{ within } \mathbf{S}_m$ 

for given  $S_1, \dots, S_M$ , each  $\mu_n$  'optimally computed' as **consensus** within  $S_m$ 

### k-Means Algorithm

use *k* **prototypes** instead of *M* historically (different from *k* nearest neighbor, though)

#### k-Means Algorithm

- 1 initialize  $\mu_1, \mu_2, \dots, \mu_k$ : say, as k randomly chosen  $\mathbf{x}_n$
- 2 alternating optimization of  $E_{in}$ : repeatedly
  - 1 optimize  $S_1, S_2, ..., S_k$ : each  $\mathbf{x}_n$  'optimally partitioned' using its closest  $\mu_i$
  - 2 optimize  $\mu_1, \mu_2, \dots, \mu_k$ : each  $\mu_n$  'optimally computed' as consensus within  $S_m$

until converge

**converge**: no change of  $S_1, S_2, ..., S_k$  anymore —guaranteed as  $E_{in}$  decreases during alternating minimization

*k*-Means: the most popular **clustering** algorithm through **alternating minimization** 

### RBF Network Using k-Means

#### RBF Network Using k-Means

- 1 run k-Means with k = M to get  $\{\mu_m\}$
- 2 construct transform  $\Phi(\mathbf{x})$  from RBF (say, Gaussian) at  $\mu_m$

$$\mathbf{\Phi}(\mathbf{x}) = [\mathsf{RBF}(\mathbf{x}, \boldsymbol{\mu}_1), \mathsf{RBF}(\mathbf{x}, \boldsymbol{\mu}_2), \dots, \mathsf{RBF}(\mathbf{x}, \boldsymbol{\mu}_M)]$$

- 3 run linear model on  $\{(\Phi(\mathbf{x}_n), y_n)\}$  to get  $\boldsymbol{\beta}$
- **4** return  $g_{RBFNET}(\mathbf{x}) = Linear Hypothesis (\beta, \Phi(\mathbf{x}))$ 
  - using unsupervised learning (k-Means) to assist feature transform—like autoencoder
  - parameters: M (prototypes), RBF (such as  $\gamma$  of Gaussian)

RBF Network: a simple (old-fashioned) model

#### Fun Time

For k-Means, consider examples  $\mathbf{x}_n \in \mathbb{R}^2$  such that all  $x_{n,1}$  and  $x_{n,2}$  are non-zero. When fixing two prototypes  $\mu_1 = [1,1]$  and  $\mu_2 = [-1,1]$ , which of the following set is the optimal  $S_1$ ?

- 1  $\{\mathbf{x}_n: x_{n,1} > 0\}$
- 2  $\{\mathbf{x}_n: x_{n,1} < 0\}$
- **3**  $\{\mathbf{x}_n: x_{n,2} > 0\}$
- 4  $\{\mathbf{x}_n: x_{n,2} < 0\}$

#### Fun Time

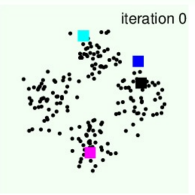
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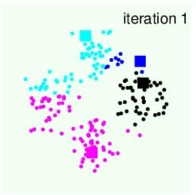
# Reference Answer: 1

Note that  $S_1$  contains examples that are closer to  $\mu_1$  than  $\mu_2$ .

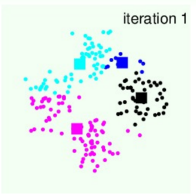
$$k = 4$$



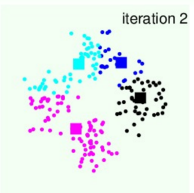
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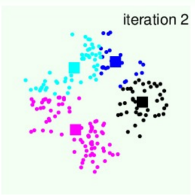
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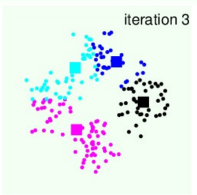
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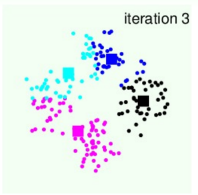
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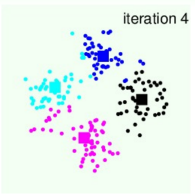
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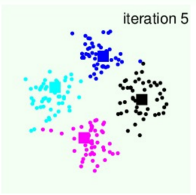
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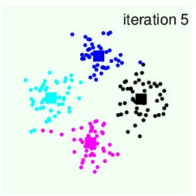
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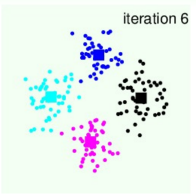
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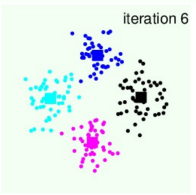
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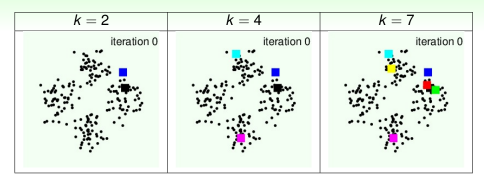
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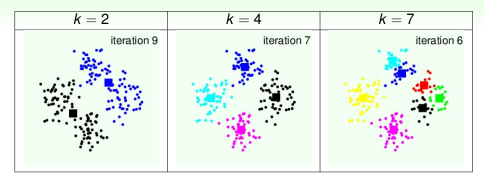


### Difficulty of k-Means



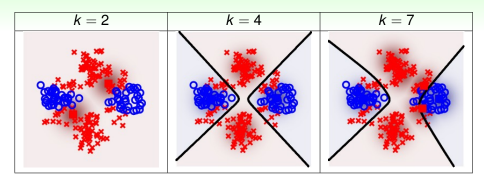
'sensitive' to k and initialization

### Difficulty of k-Means



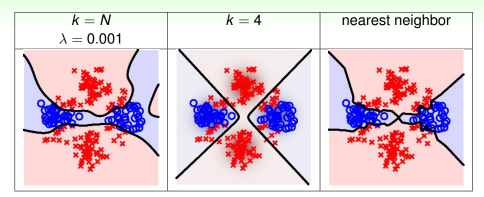
'sensitive' to k and initialization

### RBF Network Using *k*-Means



reasonable performance with proper centers

#### Full RBF Network



full RBF Network: generally less useful

#### Fun Time

When coupled with ridge linear regression, which of the following RBF Network is 'most regularized'?

- **1** small M and small  $\lambda$
- **2** small M and large  $\lambda$
- $\odot$  large M and small  $\lambda$
- 4 large M and large  $\lambda$

#### Fun Time

When coupled with ridge linear regression, which of the following RBF Network is 'most regularized'?

- $lue{1}$  small M and small  $\lambda$
- 2 small M and large  $\lambda$
- **3** large M and small  $\lambda$
- 4 large M and large  $\lambda$

# Reference Answer: 2

small M: fewer weights and more regularized; large  $\lambda$ : shorter  $\beta$  more and more regularized.

# Summary

- 1 Embedding Numerous Features: Kernel Models
- 2 Combining Predictive Features: Aggregation Models
- Oistilling Implicit Features: Extraction Models

#### Lecture 14: Radial Basis Function Network

- RBF Network Hypothesis prototypes instead of neurons as transform
- RBF Network Learning linear aggregation of prototype 'hypotheses'
- k-Means Algorithm clustering with alternating optimization
- k-Means and RBF Network in Action
   proper choice of # prototypes important
- next: extracting features from abstract data