Artificial Neural Networks Radial Basis Function Networks



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Pattern Recognition Pipeline





Classification

- Statistical classifiers
 - Bayesian classifier
 - Gaussian classifier
- Polynomial classifiers
- Non-Parametric classifiers
 - k-Nearest-Neighbor density estimation
 - Parzen windows
 - Artificial neural networks

Artificial Neural Network (ANN)



- There is no precise agreed definition among researchers as to what is an artificial neural network.
- Most would agree that it involves a network of simple processing elements (neurons), which can exhibit complex global behavior, determined by
 - the connections between the processing elements and
 - the element parameters.
- In a neural network model, simple nodes (*neurons*, or *processing elements* or *units*) are connected together to form a network of nodes.

ANN Operation



- In general an ANN operates as a function $f: x \rightarrow y$.
- The "network" arises because the function f(x) is defined as a composition of other functions g_i(x), which can further be defined as a composition of other functions, e.g. h_j(x).



General Form of ANN



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Sometimes the network is abstracted as an ANN black box.

There is great variation in ANNs, depending on:

- The number of layers
- Whether there are hidden layers or not
- The connectivity (We could have feedback loops.)
- The adaptability
- An ANN does not have to be adaptive. In practice, part of their strength comes from adapting: changing the weights of the connections in order to produce a desired signal flow.

Mathematical Description of an ANN



A widely used type of composition is the nonlinear weighted sum:

$$f(x) = \phi\left(\sum_{i} w_{i}g_{i}(x)\right)$$

- where ϕ is a predefined function that forces the output of a neuron to be in a certain range, typically [0,1] or [-1,1].
- ϕ is often referred to as an activation function.

Activation Function



- An activation function tries to mimic the firing of the neuron if the incoming signal is sufficiently strong.
- Mathematically, this is usually achieved with a sigmoid function,

$$\phi(t) = \frac{1}{1 + e^{-t}}$$



- Sigmoid functions have the following characteristic properties:
 - They are differentiable
 - They have 1 inflection point
 - They have a pair of horizontal asymptotes

Another typical sigmoid function employed in ANNs is the hyperbolic tangent, $\phi(t) = \tanh(t)$.

ANN and Classification



- The ANNs that we will examine are used in computing discriminant functions.
- Recall that, a discriminant function for class Ω_{κ} is a polynomial that evaluates to 1 if the feature vector belongs to that class. Otherwise it evaluates to zero.

$$d_{\kappa}(\vec{c}) = \begin{cases} 1 & \text{if } \vec{c} \in \Omega_{\kappa} \\ 0 & \text{otherwise} \end{cases}$$

The input of such an ANN is a feature vector \vec{c} and the output is a discriminant vector, $\vec{d} = (d_1, d_2, \dots, d_K)$.

$$\vec{c} \longrightarrow \text{ANN} \longrightarrow \vec{d}$$

Radial Basis Function ANNs



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- Radial Basis Function (RBF) networks use Radial Basis Functions in the nodes of the hidden layer.
- An RBF network is a feed-forward 3 layer network:
 - input layer, \vec{c} in our case
 - a hidden layer, where each node g_i is a separate RBF
 - an output layer, which is a weighted sum of the hidden layers.



Radial Basis Functions



- Radial basis functions were first used in 1987 by Powell.
- A radial basis function (RBF) is a real-valued function whose value depends only on the distance from the origin, so that

$$g(\vec{x}) = g(\|\vec{x}\|)$$

Alternatively, the RBF can be based on the distance from some other point \vec{q} , called a center:

$$g(\vec{x}, \vec{q}) = g(\left\| \vec{x} - \vec{q} \right\|)$$

Radial Basis Functions - continued

- So RBFs are a type of distance function.
- As a distance function, RBFs have the key characteristic that response decreases monotonically with distance from a central point.
- Its response decreases radially.



Different RBFs



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- Any distance function that decreases radially can be considered a radial basis function. Some commonly used RBFs are:
- Two different forms of Gaussians:

$$g(\|\vec{x} - \vec{q}\|) = e^{-\frac{(\|\vec{x} - \vec{q}\|)^2}{\sigma^2}} \qquad g(\|\vec{x} - \vec{q}\|) = \frac{1}{\sqrt{2\pi|\Sigma|}} e^{-\frac{1}{2}(\vec{x} - \vec{q})^T \Sigma^{-1}(\vec{x} - \vec{q})}$$

Multiquadric:

$$g(\|\vec{x} - \vec{q}\|) = \sqrt{r^2 + \|\vec{x} - \vec{q}\|^2} / r^2$$

Spline (a.k.a Logarithmic):

$$g(\|\vec{x} - \vec{q}\|) = \|\vec{x} - \vec{q}\|^2 \log(\|\vec{x} - \vec{q}\|)$$

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RBF and **Classification**



- Within the context of classification, RBFs work as follows.
- We are given a set of *N* training samples $\vec{c}_1, \vec{c}_2, \dots, \vec{c}_N$ and we want to find the best discriminant functions.
- One radial basis function (RBF) approach is to use a set of *N* basis functions, each centered around one of the training samples, i.e. $\vec{q}_i = \vec{c}_i$.
- Given a new feature vector \vec{c} we use RBFs to compute how far away it is from each of the training samples.

$$g(\vec{c} - \vec{c}_i) = g_i(\vec{c})$$

RBF and Classification – continued



The discriminant function is then treated as a linear combination of these radial basis functions.

$$d_{\kappa}(\vec{c}) = \sum_{i=1}^{N} w_{\kappa i} g(\|\vec{c} - \vec{c}_i\|) = \sum_{i=1}^{N} w_{\kappa i} g_i(\vec{c})$$

- In this type of RBFs training corresponds to the estimation of the weights w_i from the training data.
- In more detail, recall that each $d_{\kappa}(\vec{c})$ is a binary function. Thus the training set has the form:

$$T = \left\{ \left(\vec{c}_{l}, d_{\kappa(l)}(\vec{c}_{l}) \right), l = 1, 2, \dots, N \right\}$$

where $d_{\kappa(l)}(\vec{c})$ is the discriminant function of the class $\Omega_{\kappa(l)}$ to which the sample \vec{c}_l belongs.

RBFN Training



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RBFN Training - continued



Since there are N samples in my training set and K classes, I have KN such equations.

$$\begin{aligned} d_{\kappa(1)}(\vec{c}_1) &= \left(g(\|\vec{c}_1 - \vec{c}_1\|), g(\|\vec{c}_1 - \vec{c}_2\|), \dots, g(\|\vec{c}_1 - \vec{c}_N\|)\right) \vec{w}_{\kappa(1)} \\ d_{\kappa(2)}(\vec{c}_2) &= \left(g(\|\vec{c}_2 - \vec{c}_1\|), g(\|\vec{c}_2 - \vec{c}_2\|), \dots, g(\|\vec{c}_2 - \vec{c}_N\|)\right) \vec{w}_{\kappa(2)} \\ & \vdots \\ d_{\kappa(N)}(\vec{c}_N) &= \left(g(\|\vec{c}_N - \vec{c}_1\|), g(\|\vec{c}_N - \vec{c}_2\|), \dots, g(\|\vec{c}_N - \vec{c}_N\|)\right) \vec{w}_{k(N)} \end{aligned}$$

which can be written more compactly as:

$$\vec{d}' = G \vec{w}$$
$$\Rightarrow \vec{w} = G^+ \vec{d}'$$

Important Comment on RBFN Training



- If we have many feature vectors in our training data and we have an RBF estimate for each individual training sample we end up with too many RBFs, too many nodes => Slow training and **Overfitting!!**
- Solution: Use centers of clusters of feature vectors for the RBFs, instead of the individual feature vectors.
- Each RBF is now centered around $\vec{\mu}_j, j = 1, 2, ..., s$ instead of $\vec{c}_i, i = 1, 2, ..., N$: $g(\vec{c} - \vec{\mu}_j) = g_j(\vec{c})$

Updated Training of RBFNs

- 2-stage process:
- **1.** Unsupervised selection of RBF centers $\vec{\mu}_j$
 - K-means:
 - pick $s \quad \vec{\mu}_i$ values at random.
 - Assign each training sample to its nearest $\vec{\mu}_j$. Recompute $\vec{\mu}_j$ as the mean value of the samples of cluster *j*.
 - Repeat this process until the $\vec{\mu}_j$ s are stabilized. If using a Gaussian RBF, use MLE to compute Σ_i
- 2. The estimation of \vec{w} can be done as before via linear algebra methods (e.g. SVD)



Weaknesses of the 2-stage Approach

- The estimation of $\vec{\mu}_j$ and Σ_j is not guided by the discriminant function that is used to compute \vec{w} .
- Hence we have a non-symmetric approach.
- Stage 2 relies on the results of Stage 1.
- Thus, we have a propagation of estimation errors which often means an amplification of errors.
- Better solution: use an integrated, fully supervised approach like the Orthogonal Least Squares approach.



RBFN Training via Orthogonal Least Squares

- Main idea of OLS: Do not cluster as a preprocessing step.
- Rather do a sequential selection of the centers $\vec{\mu}_j$ which leads to the largest reduction in the sum of squared errors.
- Which sum of squared error (SSE)?
- The difference between the computed and the expected result (value) of the discriminant functions:

$$SSE = \sum_{i=1}^{N} \left(\hat{\vec{d}}_{i} - \vec{d}_{i}\right)^{2}$$

Orthogonal Least Squares Algorithm

- 1. Start with *N* pairs $(\vec{c}_l, d_{\kappa(l)}(\vec{c}_l))$ and s=0
- 2. For each training pair *i* of the n=N-s features vectors
 - 2a. Add the current feature \vec{c}_i to the *s* centers

The new vector becomes an additional $\vec{\mu}_i$

2b. Compute the weights \vec{w}

Use linear algebra as previously described.

2c. Compute the sum of squared error, SSE.

- 3. Out of the *n* candidate cluster centers, select the one with the smallest SSE as the next cluster center.
- 4. s++;

Repeat until the desired # of clusters is reached.



References



- 1. The sigmoid function plot is courtesy of Wikipedia <u>http://en.wikipedia.org/wiki/File:Logistic-curve.svg</u>
- 2. The RBF graph is courtesy of P. Sherrod <u>http://www.dtreg.com/rbf.htm</u>
- 3. Additional information on Orthogonal Least Squares can be found at S. Chen, C.F.N. Cowan and P.M. Grant, "Orthogonal Least Squares Learning Algorithm for Radial Basis Function Networks," *IEEE Transactions on Neural Networks*, Vol. 2, No. 2, March 1991.