CS434a/541a: Pattern Recognition Prof. Olga Veksler

Lecture 6

Today

- Introduction to nonparametric techniques
- Basic Issues in Density Estimation
- Two Density Estimation Methods
 - 1. Parzen Windows
 - 2. Nearest Neighbors

Non-Parametric Methods

- Neither probability distribution nor discriminant function is known
 - Happens quite often
- All we have is labeled data



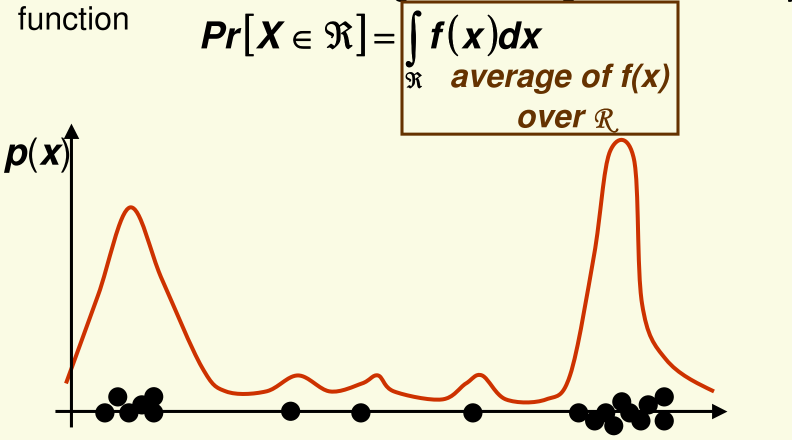
 Estimate the probability distribution from the labeled data a lot is known "easier"

> little is known "harder"

- In previous lectures we assumed that either
 - **1**. someone gives us the density p(x)
 - In pattern recognition applications this never happens
 - 2. someone gives us $p(x|\theta)$
 - Does happen sometimes, **but**
 - we are likely to suspect whether the given $p(x|\theta)$ models the data well
 - Most parametric densities are unimodal (have a single local maximum), whereas many practical problems involve multi-modal densities

- Nonparametric procedures can be used with arbitrary distributions and without any assumption about the forms of the underlying densities
- There are two types of nonparametric methods:
 - Parzen windows
 - Estimate likelihood $p(x | c_j)$
 - Nearest Neighbors
 - Bypass likelihood and go directly to posterior estimation $P(c_j | x)$

- Nonparametric techniques attempt to estimate the underlying density functions from the training data
 - Idea: the more data in a region, the larger is the density function $D_{\mu}[V = \infty]$



salmon length x

 $Pr[X \in \Re] = \int_{\Omega} f(x) dx$

How can we approximate \$\mathbf{Pr}_{r}^{\pi}[X \in \bar{x}_{1}]\$ and \$\mathbf{Pr}[X \in \bar{x}_{2}]\$?
\$\mathbf{Pr}[X \in \bar{x}_{1}] \approx \frac{6}{20}\$ and \$\mathbf{Pr}[X \in \bar{x}_{2}] \approx \frac{6}{20}\$

salmon length x

- Should the density curves above R₁ and R₂ be equally high?
 - No, since is \mathcal{R}_1 smaller than \mathcal{R}_2 $Pr[X \in \mathfrak{R}_1] = \int f(x) dx \approx \int f(x) dx = Pr[X \in \mathfrak{R}_2]$
 - To get density, normalize by region size

p(x)

• Assuming f(x) is basically flat inside \mathcal{R} ,

 $\frac{\# of \ samples \ in \Re}{total \ \# of \ samples} \approx \Pr[X \in \Re] = \int_{\Re} f(y) dy \approx f(x) * Volume(\Re)$

Thus, density at a point *x* inside *R*, can be approximated

 $f(x) \approx \frac{\#of \ samples \ in \Re}{total \ \#of \ samples} \ \frac{1}{Volume(\Re)}$

Now let's derive this formula more formally

Binomial Random Variable

- Let us flip a coin *n* times (each one is called "trial")
 - Probability of head ρ , probability of tail is 1- ρ
- Binomial random variable K counts the number of heads in n trials

$$P(K = k) = {\binom{n}{k}} \rho^{k} (1 - \rho)^{n-k}$$

where ${\binom{n}{k}} = \frac{n!}{k!(n-k)!}$

- Mean is $E(K) = n\rho$
- Variance is $var(K) = n\rho(1-\rho)$

Density Estimation: Basic Issues

From the definition of a density function, probability

 ρ that a vector x will fall in region R is:

$$\rho = \Pr[\mathbf{x} \in \Re] = \int_{\Re} p(\mathbf{x}') d\mathbf{x}'$$

Suppose we have samples x₁, x₂,..., x_n drawn from the distribution p(x). The probability that k points fall in R is then given by binomial distribution:

$$\Pr[K = k] = {\binom{n}{k}} \rho^{k} (1 - \rho)^{n-k}$$

• Suppose that **k** points fall in \mathcal{R} , we can use MLE to estimate the value of ρ . The likelihood function is

$$p(\mathbf{x}_1,...,\mathbf{x}_n \mid \rho) = {\binom{n}{k}} \rho^k (1-\rho)^{n-k}$$

Density Estimation: Basic Issues

$$p(\mathbf{x}_1,...,\mathbf{x}_n \mid \rho) = {\binom{n}{k}} \rho^k (1-\rho)^{n-k}$$

- This likelihood function is maximized at $\rho = \frac{\kappa}{n}$
- Thus the MLE is $\hat{\rho} = \frac{\kappa}{n}$
- Assume that *p*(*x*) is continuous and that the region *R* is so small that *p*(*x*) is approximately constant in *R*

$$\int_{\Re} p(x') dx' \cong p(x) V$$

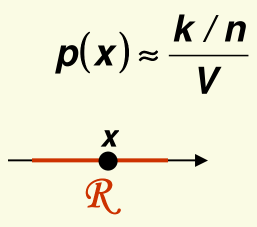
- \boldsymbol{x} is in $\boldsymbol{\mathcal{R}}$ and \boldsymbol{V} is the volume of $\boldsymbol{\mathcal{R}}$
- Recall from the previous slide: $\rho = \int p(x') dx'$
- Thus p(x) can be approximated:

$$\widehat{p}(x) \approx \frac{k/n}{V}$$

 $p(\mathbf{X})$

Density Estimation: Basic Issues

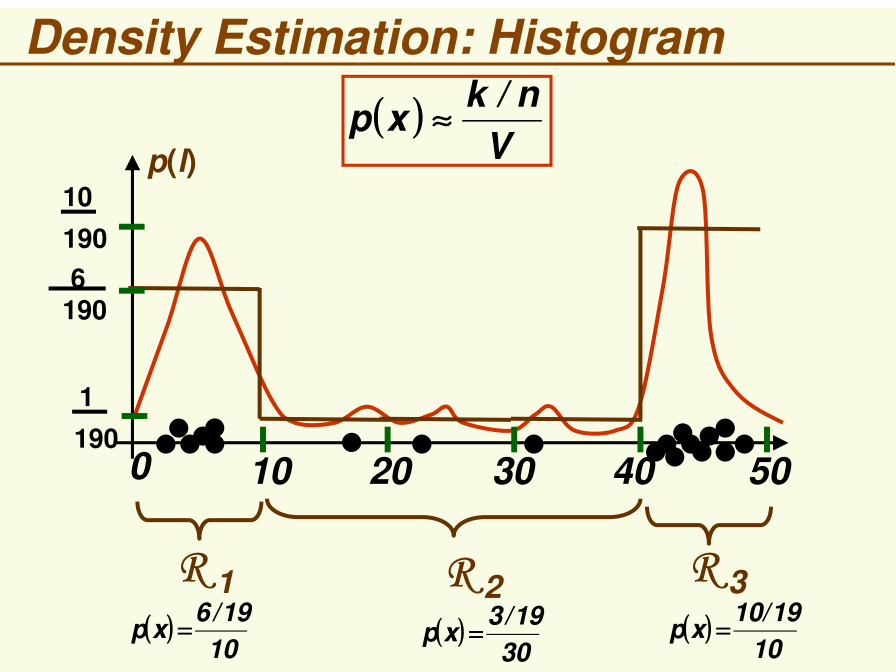
This is exactly what we had before:



- $p(x) \approx \frac{k/n}{V}$ x is inside some region \Re k = number of samples inside \Re **x** is inside some region \mathcal{R} *n*=total number of samples inside \mathcal{R} $V = volume of \mathcal{R}_{.}$
- Our estimate will always be the average of true density over \mathcal{R}_{i}

$$p(x) \approx \frac{k/n}{V} = \frac{\hat{\rho}}{V} \approx \frac{\int p(x')dx'}{V}$$

Ideally, $p(\mathbf{x})$ should be constant inside \mathcal{R}



• If regions \mathcal{R}_{i} 's do not overlap, we have a histogram

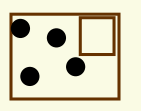
Density Estimation: Accuracy

- How accurate is density approximation $p(x) \approx \frac{k/n}{V}$?
- We have made two approximations
 - *n* as n increases, this estimate becomes more accurate

$$2. \int p(x')dx' \cong p(x)V$$

1. $\hat{\rho} = \frac{k}{k}$

as \mathcal{R} grows smaller, the estimate becomes more accurate



- As we shrink \mathcal{R} we have to make sure it contains samples, otherwise our estimated $\mathbf{p}(\mathbf{x}) = 0$ for all \mathbf{x} in \mathcal{R}
- Thus in theory, if we have an unlimited number of samples, to we get convergence as we simultaneously increase the number of samples *n*, and shrink region *R*, but not too much so that *R* still contains a lot of samples

Density Estimation: Accuracy $p(x) \approx \frac{k/n}{V}$

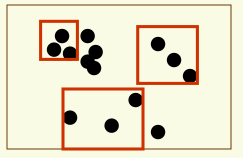
- In practice, the number of samples is always fixed
- Thus the only available option to increase the accuracy is by decreasing the size of *R* (*V* gets smaller)
 - If V is too small, p(x)=0 for most x, because most regions will have no samples
 - Thus have to find a compromise for **V**
 - not too small so that it has enough samples
 - but also not too large so that *p*(*x*) is approximately constant inside *V*

Density Estimation: Two Approaches

$$p(x) \approx \frac{k/n}{V}$$

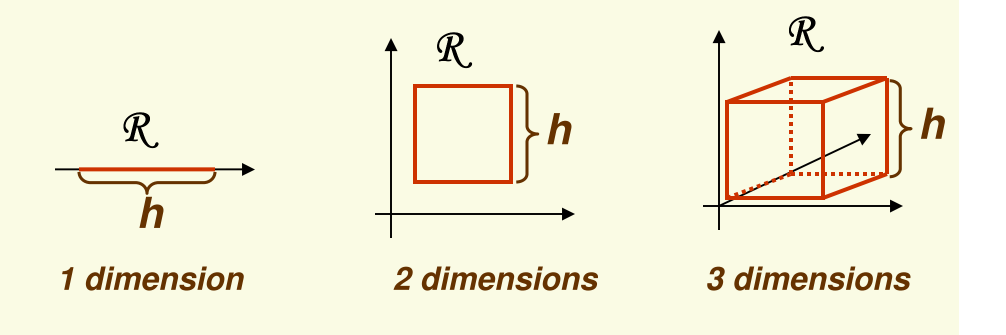
- 1. Parzen Windows:
 - Choose a fixed value for volume V and determine the corresponding k from the data

- 2. k-Nearest Neighbors
 - Choose a fixed value for *k* and determine the corresponding volume *V* from the data



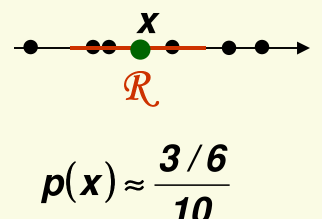
 Under appropriate conditions and as number of samples goes to infinity, both methods can be shown to converge to the true *p*(*x*)

- In Parzen-window approach to estimate densities we fix the size and shape of region *R*
- Let us assume that the region R is a *d*-dimensional hypercube with side length *h* thus it's volume is *h^d*



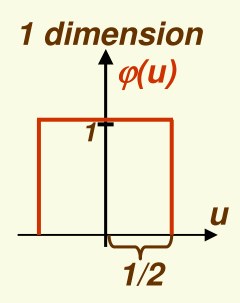
To estimate the density at point *x*, simply center the region *R* at *x*, count the number of samples in *R*, and substitute everything in our formula

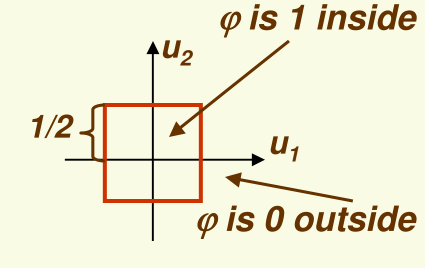
$$p(\mathbf{x}) \approx \frac{k/n}{V}$$



- We wish to have an analytic expression for our approximate density *R*
- Let us define a window function

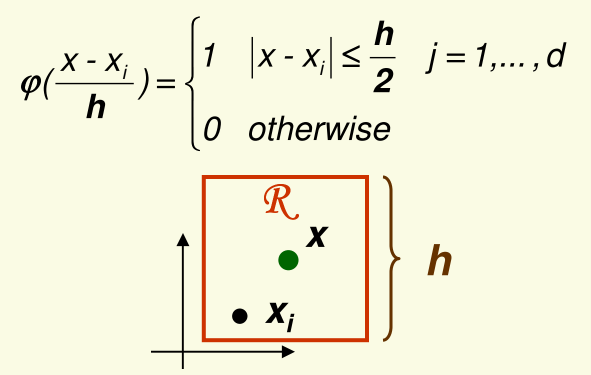
$$\varphi(u) = \begin{cases} 1 & |u_j| \le \frac{1}{2} & j = 1, \dots, d \\ 0 & otherwise \end{cases}$$





2 dimensions

• Recall we have samples x_1, x_2, \ldots, x_n . Then



$$\varphi(\frac{X-X_i}{h}) = \begin{cases} 1\\ 0 \end{cases}$$

if x_i is inside the hypercube with width h and centered at x otherwise

How do we count the total number of sample points x₁, x₂,..., x_n which are inside the hypercube with side h and centered at x?

$$\boldsymbol{k} = \sum_{i=1}^{i=n} \varphi \left(\frac{\boldsymbol{x} - \boldsymbol{x}_i}{\boldsymbol{h}} \right)$$

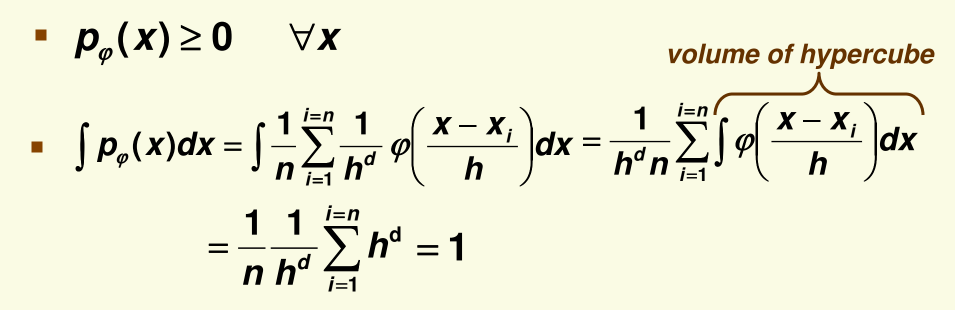
• Recall
$$p(x) \approx \frac{k/n}{V}$$

• Thus we get the desired analytical expression for the estimate of density $p_{\omega}(x)$

$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^{d}} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_{i}}{h}\right)$$

$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^{d}} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_{i}}{h}\right)$$

• Let's make sure $p_{\varphi}(\mathbf{x})$ is in fact a density



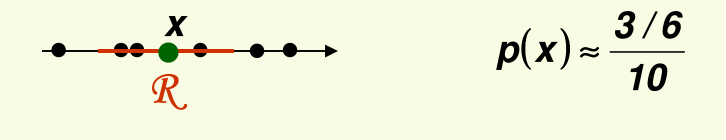
Today

- Continue nonparametric techniques
 - 1. Finish Parzen Windows
 - 2. Start Nearest Neighbors (hopefully)

$$p(\mathbf{x}) \approx \frac{k / n}{V}$$

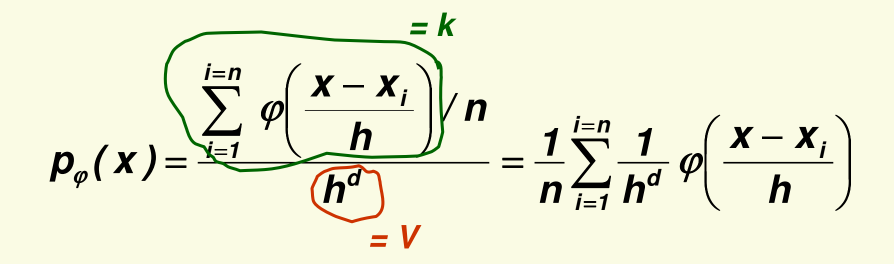
x is inside some region R k = number of samples inside R n=total number of samples inside R V = volume of R

To estimate the density at point *x*, simply center the region *R* at *x*, count the number of samples in *R*, and substitute everything in our formula





Formula for Parzen window estimation



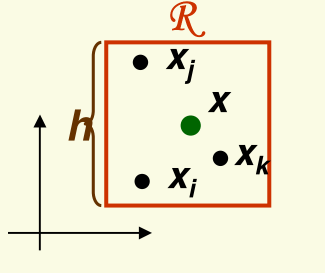
Parzen Windows: Example in 1D

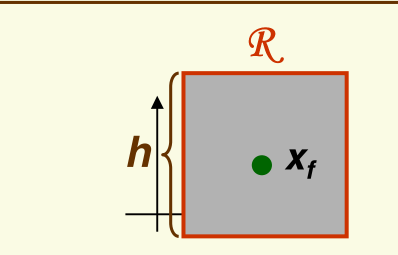
$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^d} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_i}{h}\right)$$

- Suppose we have 7 samples $D = \{2, 3, 4, 8, 10, 11, 12\}$
- Let window width h=3, estimate density at x=1 $p_{\varphi}(1) = \frac{1}{7} \sum_{i=1}^{1} \frac{1}{3} \varphi\left(\frac{1-x_i}{3}\right) = \frac{1}{21} \left[\varphi\left(\frac{1-2}{3}\right) + \varphi\left(\frac{1-3}{3}\right) + \varphi\left(\frac{1-4}{3}\right) + \dots + \varphi\left(\frac{1-12}{3}\right) \right]$ $\left| -\frac{1}{3} \right| \le 1/2$ $\left| -\frac{2}{3} \right| > 1/2$ $\left| -1 \right| > 1/2$ $\left| -\frac{11}{3} \right| > 1/2$ $p_{\varphi}(1) = \frac{1}{7} \sum_{i=1}^{1} \frac{1}{3} \varphi\left(\frac{1-x_i}{3}\right) = \frac{1}{21} \left[1+0+0+\dots+0 \right] = \frac{1}{21}$

Parzen Windows: Sum of Functions

• Fix **x**, let **i** vary and ask • For which samples \mathbf{x}_i is $\varphi\left(\frac{\mathbf{x} - \mathbf{x}_i}{h}\right) = 1$?





Now fix f and let x vary and ask
For which x is \$\varphi\bigg(\frac{x-x_f}{h}\bigg)=1\$? For all x in gray box
Thus \$\varphi\bigg(\frac{x-x_f}{h}\bigg)=1\$ is simply a function which is 1 inside square of width h centered at \$x_f\$ and 0 otherwise!

Parzen Windows: Sum of Functions

• Now let's look at our density estimate $p_{\varphi}(x)$ again:

$$p_{\varphi}(x) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^{d}} \varphi\left(\frac{x - x_{i}}{h}\right) = \sum_{i=1}^{i=n} \frac{1}{nh^{d}} \varphi\left(\frac{x - x_{i}}{h}\right)$$

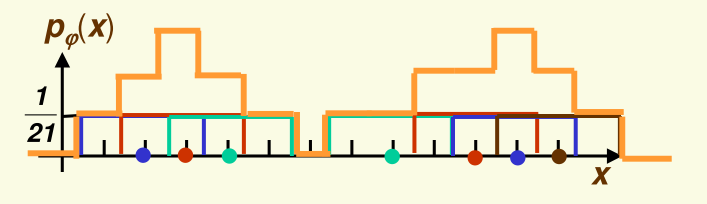
$$1 \text{ inside square centered at } x_{i}$$

$$0 \text{ otherwise}$$

• Thus $p_{\varphi}(\mathbf{x})$ is just a sum of \mathbf{n} "box like" functions each of height $\frac{1}{nh^{d}}$

Parzen Windows: Example in 1D

- Let's come back to our example
 - 7 samples **D**={2,3,4,8,10,11,12}, **h**=3

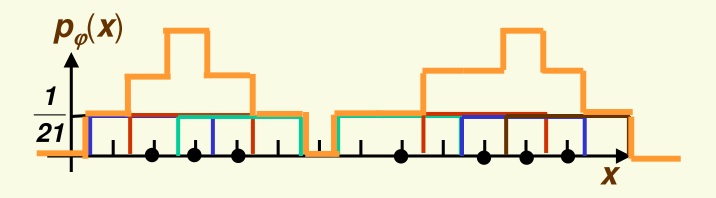


- To see what the function looks like, we need to generate 7 boxes and add them up
- The width is *h*=3 and the height, according to previous slide is

$$\frac{1}{nh^d} = \frac{1}{21}$$

Parzen Windows: Interpolation

 In essence, window function φ is used for interpolation: each sample x_i contributes to the resulting density at x if x is close enough to x_i

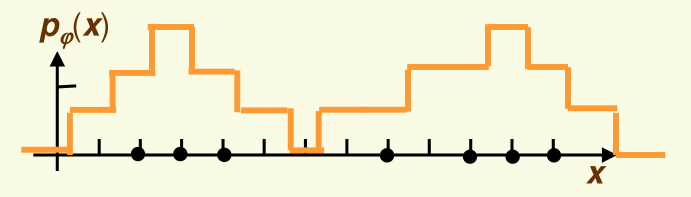


Parzen Windows: Drawbacks of Hypercube φ

As long as sample point x_i and x are in the same hypercube, the contribution of x_i to the density at x is constant, regardless of how close x_i is to x

$$\varphi\left(\frac{x-x_{1}}{h}\right) = \varphi\left(\frac{x-x_{2}}{h}\right) = 1$$

• The resulting density $p_{\varphi}(\mathbf{x})$ is not smooth, it has discontinuities



Parzen Windows: general φ

$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^d} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_i}{h}\right)$$

• We can use a general window φ as long as the resulting $p_{\varphi}(\mathbf{x})$ is a legitimate density, i.e. $\varphi_1(\mathbf{u}) = \varphi_2(\mathbf{u})$

1.
$$p_{\varphi}(u) \geq 0$$

• satisfied if $\varphi(u) \ge 0$

$$2. \int p_{\varphi}(x) dx = 1$$

• satisfied if
$$\int \varphi(u) du = 1$$

$$\mathbf{P}_{\varphi}(\mathbf{x}) d\mathbf{x} = \frac{1}{nh^{d}} \sum_{i=1}^{i=n} \int \varphi \left(\frac{\mathbf{x} - \mathbf{x}_{i}}{h} \right) d\mathbf{x} = \frac{1}{nh^{d}} \sum_{i=1}^{n} \int h^{n} \varphi(\mathbf{u}) d\mathbf{u} = 1$$

change coordinates to
$$u = \frac{x - x_i}{h}$$
, thus $du = \frac{dx}{h}$

Parzen Windows: general φ

$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^d} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_i}{h}\right)$$

- Notice that with the general window φ we are no longer counting the number of samples inside \mathcal{R} .
- We are counting the weighted average of potentially every single sample point (although only those within distance *h* have any significant weight)

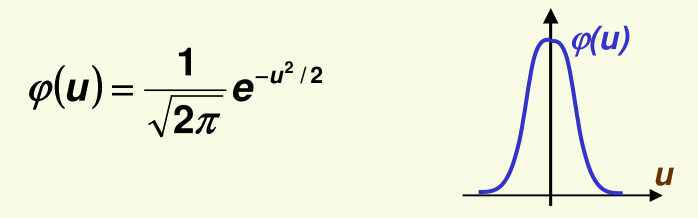
With infinite number of samples, and appropriate conditions, it can still be shown that

 $p_{\varphi}^{n}(x) \rightarrow p(x)$

Parzen Windows: Gaussian φ

$$\boldsymbol{p}_{\varphi}(\boldsymbol{x}) = \frac{1}{n} \sum_{i=1}^{i=n} \frac{1}{h^d} \varphi\left(\frac{\boldsymbol{x} - \boldsymbol{x}_i}{h}\right)$$

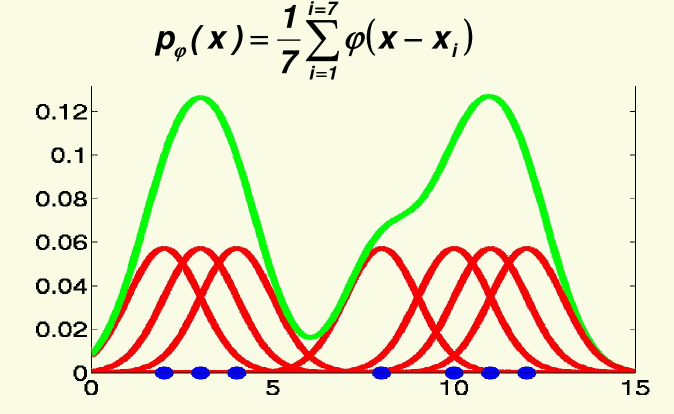
• A popular choice for φ is N(0, 1) density



- Solves both drawbacks of the "box" window
 - Points x which are close to the sample point x_i receive higher weight
 - Resulting density $p_{\varphi}(\mathbf{x})$ is smooth

Parzen Windows: Example with General φ

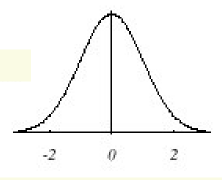
- Let's come back to our example
 - 7 samples **D**={2,3,4,8,10,11,12}, **h**=1



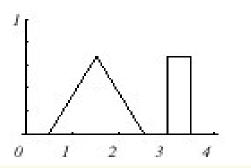
• $p_{\varphi}(\mathbf{x})$ is the sum of of 7 Gaussians, each centered at one of the sample points, and each scaled by 1/7

Parzen Windows: Did We Solve the Problem?

- Let's test if we solved the problem
 - 1. Draw samples from a known distribution
 - 2. Use our density approximation method and compare with the true density
- We will vary the number of samples *n* and the window size *h*
- We will play with 2 distributions

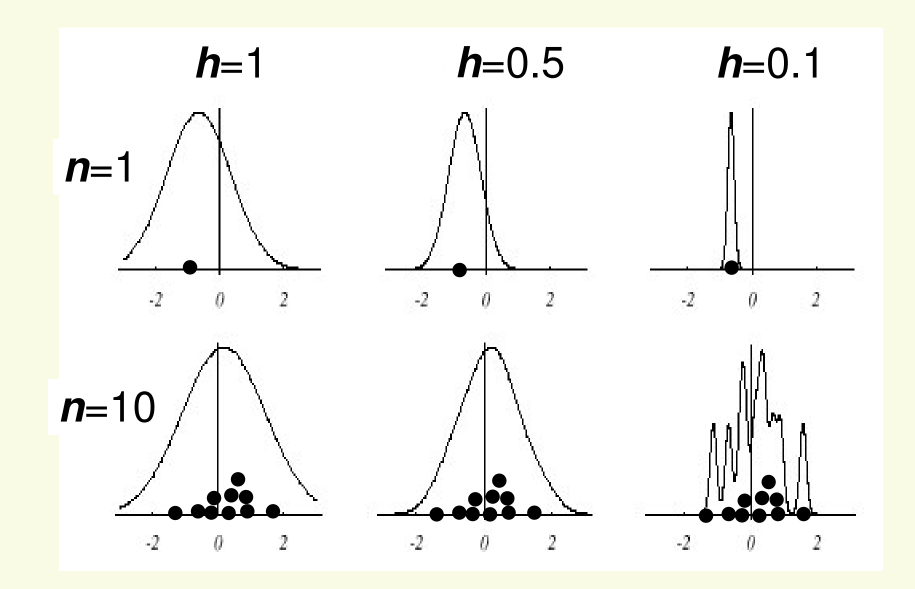


N(0,1)

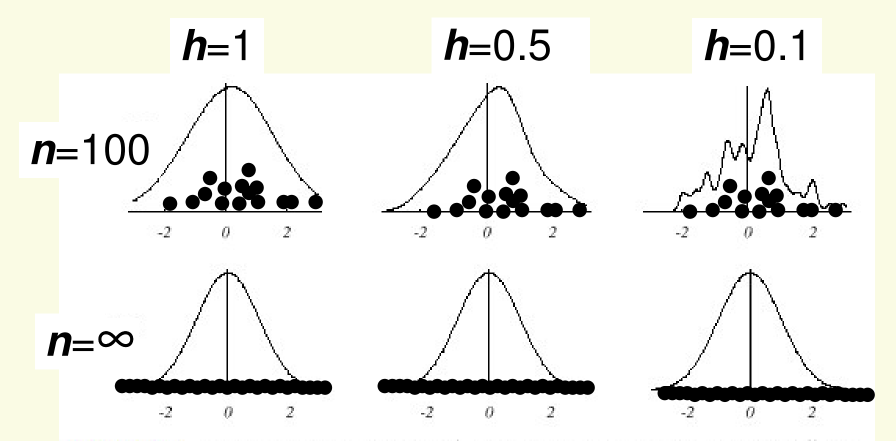


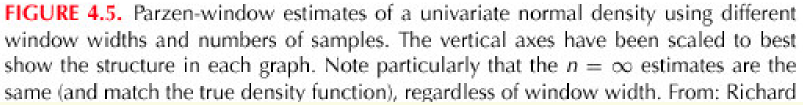
triangle and uniform mixture

Parzen Windows: True Density N(0,1)

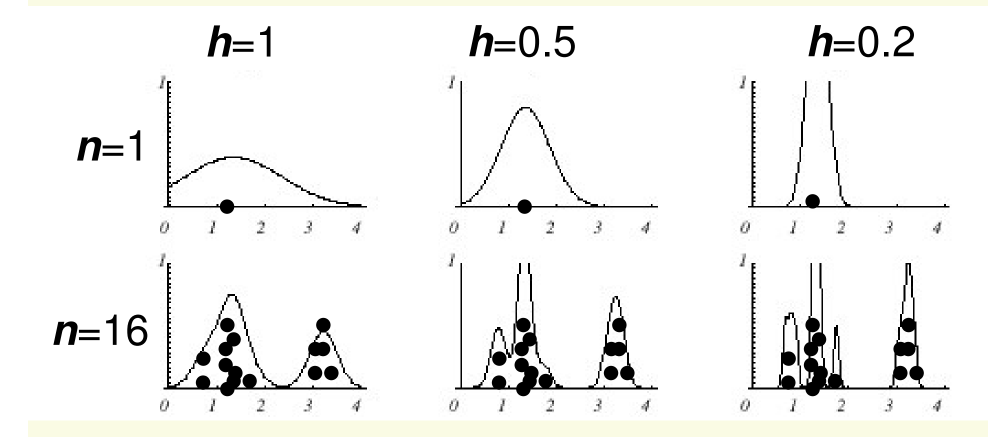


Parzen Windows: True Density N(0,1)





Parzen Windows: True density is Mixture of Uniform and Triangle



Parzen Windows: True density is Mixture of Uniform and Triangle

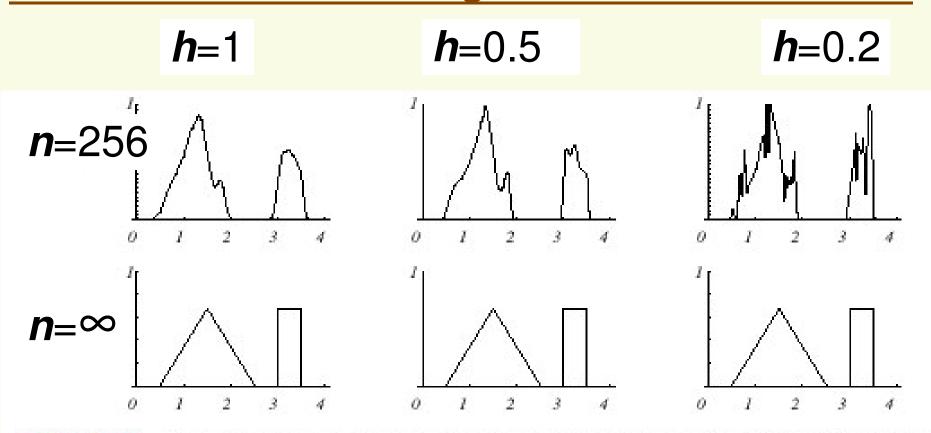
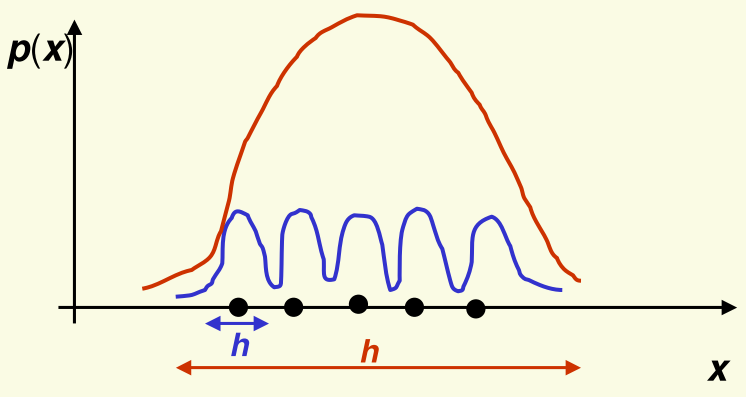


FIGURE 4.7. Parzen-window estimates of a bimodal distribution using different window widths and numbers of samples. Note particularly that the $n = \infty$ estimates are the same (and match the true distribution), regardless of window width. From: Richard O. Duda, Peter E. Hart, and David G. Stork, *Pattern Classification*. Copyright © 2001 by John Wiley & Sons, Inc.

Parzen Windows: Effect of Window Width h

- By choosing *h* we are guessing the region where density is approximately constant
- Without knowing anything about the distribution, it is really hard to guess were the density is approximately constant



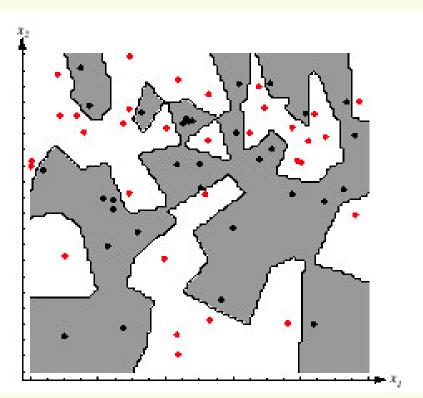
Parzen Windows: Effect of Window Width h

- If *h* is small, we superimpose *n* sharp pulses centered at the data
 - Each sample point x_i influences too small range of x
 - Smoothed too little: the result will look noisy and not smooth enough
- If *h* is large, we superimpose broad slowly changing functions,
 - Each sample point x_i influences too large range of x
 - Smoothed too much: the result looks oversmoothed or "outof-focus"
- Finding the best *h* is challenging, and indeed no single *h* may work well
 - May need to adapt *h* for different sample points
- However we can try to learn the best *h* to use from the test data

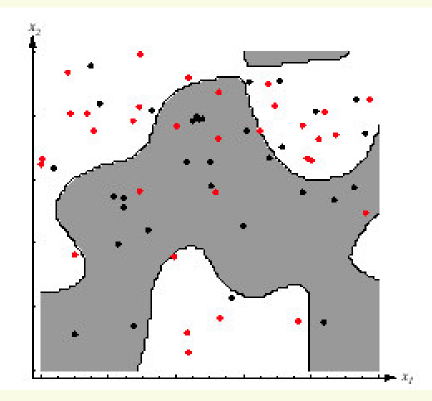
Parzen Windows: Classification Example

- In classifiers based on Parzen-window estimation:
 - We estimate the densities for each category and classify a test point by the label corresponding to the maximum posterior
 - The decision region for a Parzen-window classifier depends upon the choice of window function as illustrated in the following figure

Parzen Windows: Classification Example



- For small enough window size h is classification on training data is be perfect
- However decision boundaries are complex and this solution is not likely to generalize well to novel data



- For larger window size *h*, classification on training data is not perfect
- However decision boundaries are simpler and this solution is more likely to generalize well to novel data

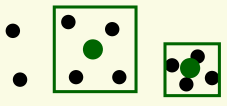
Parzen Windows: Summary

Advantages

- Can be applied to the data from any distribution
- In theory can be shown to converge as the number of samples goes to infinity
- Disadvantages
 - Number of training data is limited in practice, and so choosing the appropriate window size *h* is difficult
 - May need large number of samples for accurate estimates
 - Computationally heavy, to classify one point we have to compute a function which potentially depends on all samples
 - Window size *h* is not trivial to choose

- Recall the generic expression for density estimation $p(x) \approx \frac{k / n}{V}$
- In Parzen windows estimation, we fix V and that determines k, the number of points inside V
- In k-nearest neighbor approach we fix k, and find
 V that contains k points inside

- kNN approach seems a good solution for the problem of the "best" window size
 - Let the cell volume be a function of the training data
 - Center a cell about x and let it grows until it captures k samples
 - k are called the k nearest-neighbors of x



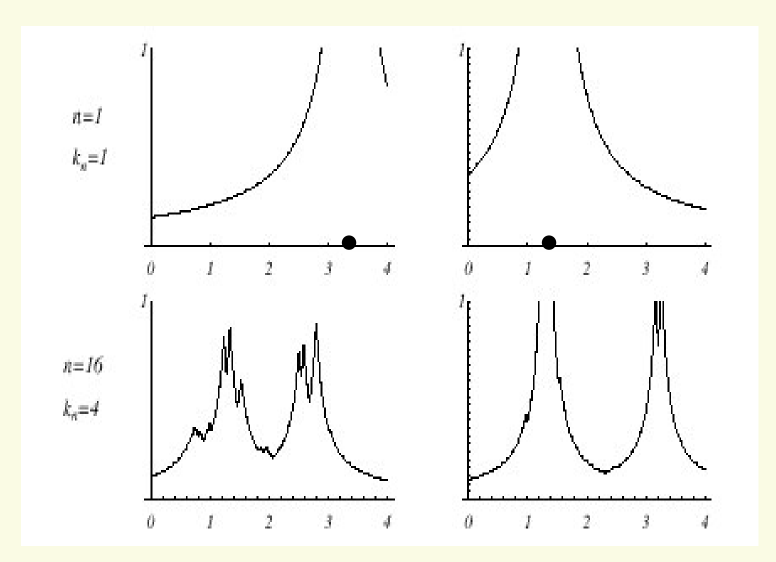
- 2 possibilities can occur:
 - Density is high near x; therefore the cell will be small which provides a good resolution
 - Density is low; therefore the cell will grow large and stop until higher density regions are reached

- Of course, now we have a new question
 - How to choose k?
- A good "rule of thumb" is $\mathbf{k} = \sqrt{\mathbf{n}}$
 - Can prove convergence if n goes to infinity
 - Not too useful in practice, however
- Let's look at 1-D example
 - we have one sample, i.e. n = 1

• But the estimated p(x) is not even close to a density function:

$$\int_{-\infty}^{1} \frac{1}{2|x-x_1|} dx = \infty \neq 1$$

k-Nearest Neighbor: Gaussian and Uniform plus Triangle Mixture Estimation



k-Nearest Neighbor: Gaussian and Uniform plus Triangle Mixture Estimation

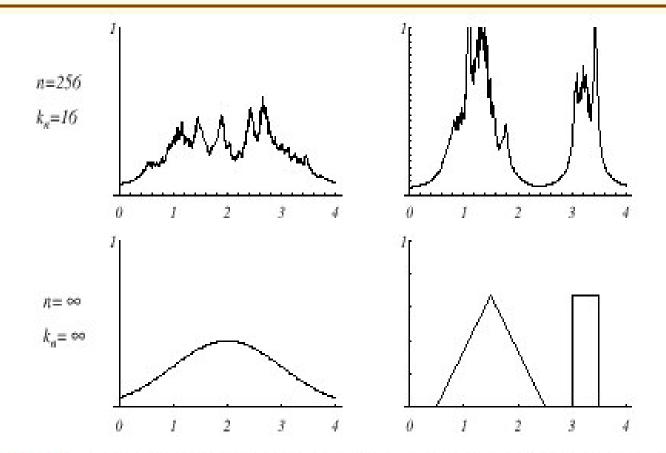


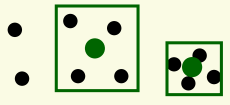
FIGURE 4.12. Several *k*-nearest-neighbor estimates of two unidimensional densities: a Gaussian and a bimodal distribution. Notice how the finite *n* estimates can be quite "spiky." From: Richard O. Duda, Peter E. Hart, and David G. Stork, *Pattern Classification*. Copyright © 2001 by John Wiley & Sons, Inc.

Today

- Continue with Nonparametric Density Estimation
 - Finish Nearest Neighbor

$$p(\mathbf{x}) \approx \frac{\mathbf{k} / \mathbf{n}}{\mathbf{V}}$$

- kNN approach seems a good solution for the problem of the "best" window size
 - Let the cell volume be a function of the training data
 - Center a cell about x and let it grows until it captures k samples
 - k are called the k nearest-neighbors of x



k-Nearest Neighbor: Gaussian and Uniform plus Triangle Mixture Estimation

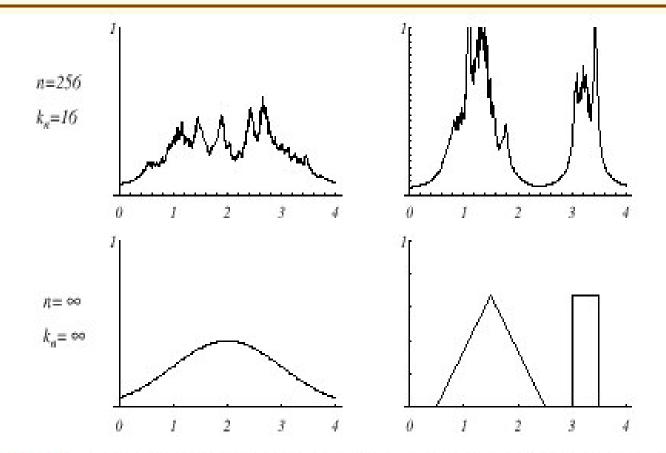


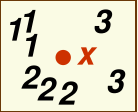
FIGURE 4.12. Several *k*-nearest-neighbor estimates of two unidimensional densities: a Gaussian and a bimodal distribution. Notice how the finite *n* estimates can be quite "spiky." From: Richard O. Duda, Peter E. Hart, and David G. Stork, *Pattern Classification*. Copyright © 2001 by John Wiley & Sons, Inc.

- Thus straightforward density estimation *p*(*x*) does not work very well with kNN approach because the resulting density estimate
 - 1. Is not even a density
 - 2. Has a lot of discontinuities (looks very spiky, not differentiable)
 - Even for large regions with no observed samples the estimated density is far from zero (tails are too heavy)
- Notice in the theory, if infinite number of samples is available, we could construct a series of estimates that converge to the true density using kNN estimation. However this theorem is not very useful in practice because the number of samples is always limited

- However we shouldn't give up the nearest neighbor approach yet
- Instead of approximating the density *p*(*x*), we can use kNN method to approximate the posterior distribution *P*(*c_i*|*x*)
 - We don't even need p(x) if we can get a good estimate on $P(c_i|x)$

- How would we estimate P(c_i / x) from a set of n labeled samples?
- Recall our estimate for density: $p(x) \approx \frac{k/n}{v}$
- Let's place a cell of volume V around x and capture k samples
 - *k_i* samples amongst *k* labeled *c_i* then:

$$p(c_i, x) \approx \frac{k_i / n}{V}$$



Using conditional probability, let's estimate posterior:

$$p(c_{i} | x) = \frac{p(x,c_{i})}{p(x)} = \frac{p(x,c_{i})}{\sum_{j=1}^{m} p(x,c_{j})} \approx \frac{k_{i} / n}{V \sum_{j=1}^{m} \frac{k_{j} / n}{V}} = \frac{k_{i}}{\sum_{j=1}^{m} k_{j}} = \frac{k_{i}}{k}$$

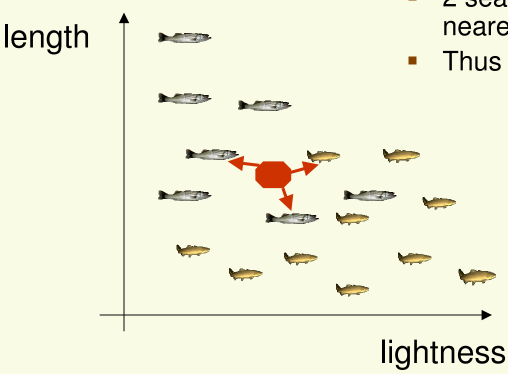
Thus our estimate of posterior is just the fraction of samples which belong to class c_i:

$$p(c_i \mid x) = \frac{k_i}{k}$$

- This is a very simple and intuitive estimate
- Under the zero-one loss function (MAP classifier) just choose the class which has the largest number of samples in the cell
- Interpretation is: given an unlabeled example (that is *x*), find *k* most similar labeled examples (closest neighbors among sample points) and assign the most frequent class among those neighbors to *x*

k-Nearest Neighbor: Example

- Back to fish sorting
 - Suppose we have 2 features, and collected sample points as in the picture
 - Let **k** = 3



- 2 sea bass, 1 salmon are the 3 nearest neighbors
- Thus classify as sea bass

kNN: How Well Does it Work?

- kNN rule is certainly simple and intuitive, but does it work?
- Pretend that we can get an unlimited number of samples
- By definition, the best possible error rate is the Bayes rate *E**
- Even for k =1, the nearest-neighbor rule leads to an error rate greater than E*
- But as $n \rightarrow \infty$, it can be shown that nearest neighbor rule error rate is smaller than **2E***
- If we have a lot of samples, the kNN rule will do very well !

1NN: Voronoi Cells

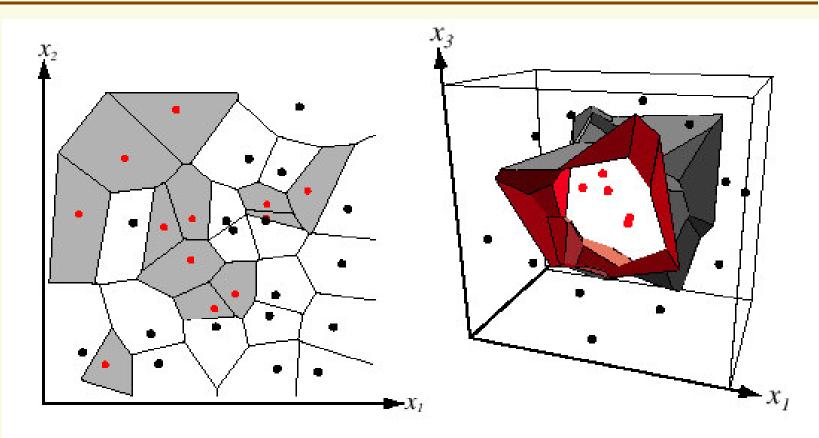
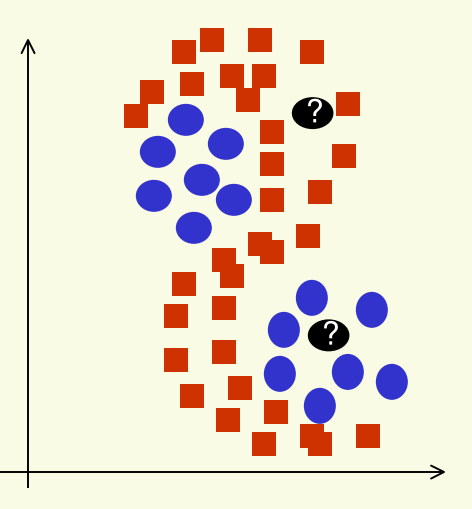


FIGURE 4.13. In two dimensions, the nearest-neighbor algorithm leads to a partitioning of the input space into Voronoi cells, each labeled by the category of the training point it contains. In three dimensions, the cells are three-dimensional, and the decision boundary resembles the surface of a crystal. From: Richard O. Duda, Peter E. Hart, and David G. Stork, *Pattern Classification*. Copyright © 2001 by John Wiley & Sons, Inc.

kNN: Multi-Modal Distributions

- Most parametric distributions would not work for this 2 class classification problem:
- Nearest neighbors will do reasonably well, provided we have a lot of samples



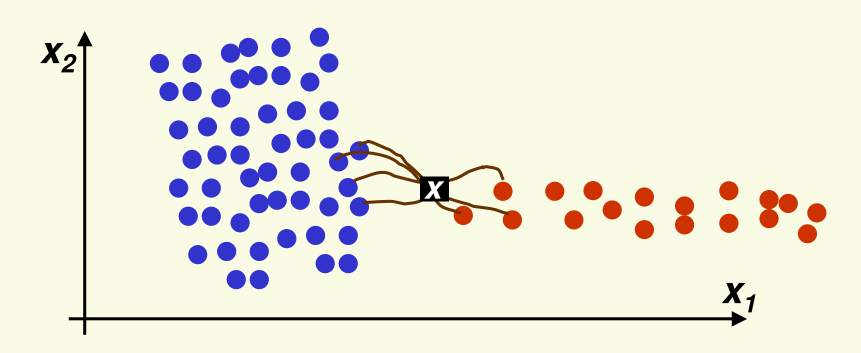
kNN: How to Choose k?

- In theory, when the infinite number of samples is available, the larger the *k*, the better is classification (error rate gets closer to the optimal Bayes error rate)
- But the caveat is that all k neighbors have to be close to x
 - Possible when infinite # samples available
 - Impossible in practice since # samples is finite

kNN: How to Choose k?

- In practice
 - k should be large so that error rate is minimized
 - k too small will lead to noisy decision boundaries
 - k should be small enough so that only nearby samples are included
 - k too large will lead to over-smoothed boundaries
- Balancing 1 and 2 is not trivial
 - This is a recurrent issue, need to smooth data, but not too much

kNN: How to Choose k?



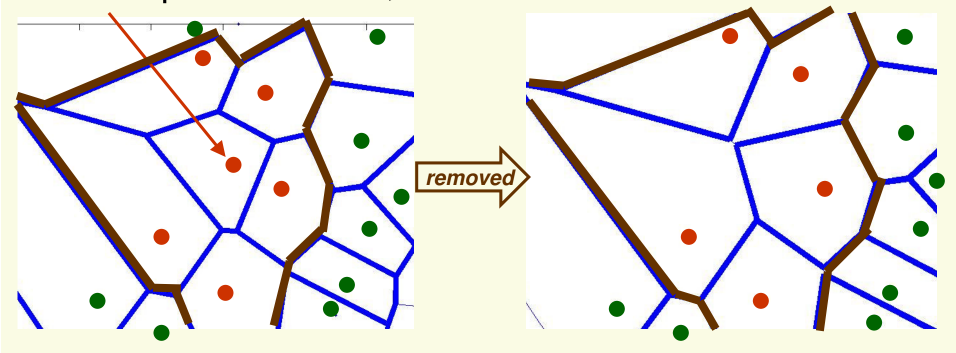
- For k = 1, ..., 7 point x gets classified correctly
 - red class
- For larger *k* classification of *x* is wrong
 - blue class

kNN: Computational Complexity

- Basic *kNN* algorithm stores all examples. Suppose we have *n* examples each of dimension *k*
 - **O**(**d**) to compute distance to one example
 - O(nd) to find one nearest neighbor
 - **O**(*knd*) to find *k* closest examples examples
 - Thus complexity is O(knd)
- This is prohibitively expensive for large number of samples
- But we need large number of samples for *kNN* to work well!

Reducing Complexity: Editing 1NN

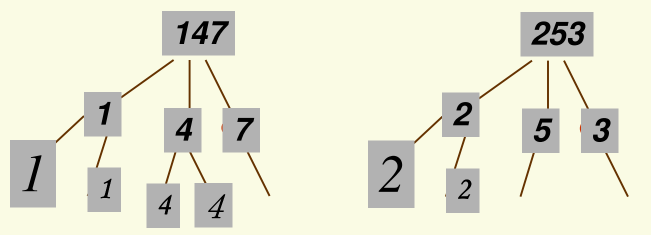
If all voronoi neighbors have the same class, a sample is useless, we can remove it:



- Number of samples decreases
- We are guaranteed that the decision boundaries stay the same

Reducing Complexity: kNN prototypes

 Explore similarities between samples to represent data as search trees of *prototypes*



- Advantages: Complexity decreases
- Disadvantages:
 - finding good search tree is not trivial
 - will not necessarily find the closest neighbor, and thus **not** guaranteed that the decision boundaries stay the same

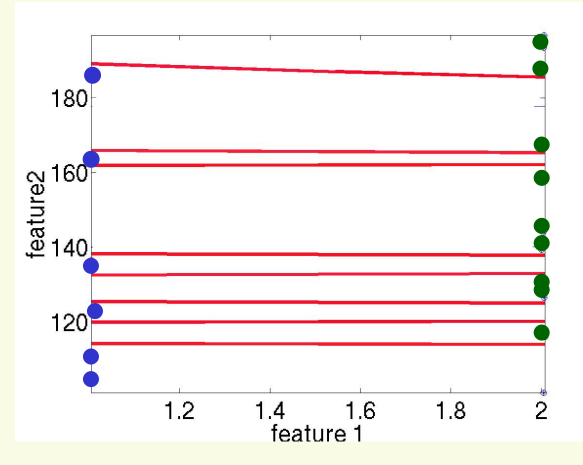
kNN: Selection of Distance

 So far we assumed we use Euclidian Distance to find the nearest neighbor:

$$D(a,b) = \sqrt{\sum_{k} (a_k - b_k)^2}$$

- However some features (dimensions) may be much more discriminative than other features (dimensions)
- Eucleadian distance treats each feature as equally important

kNN: Extreme Example of Distance Selection



- decision boundaries for blue and green classes are in red
- These boundaries are really bad because
 - feature 1 is discriminative, but it's scale is small
 - feature 2 gives no class information (noise) but its scale is large

kNN: Selection of Distance

- Extreme Example
 - feature 1 gives the correct class: 1 or 2
 - feature 2 gives irrelevant number from 100 to 200
- Suppose we have to find the class of x=[1 100] and we have 2 samples [1 150] and [2 110]

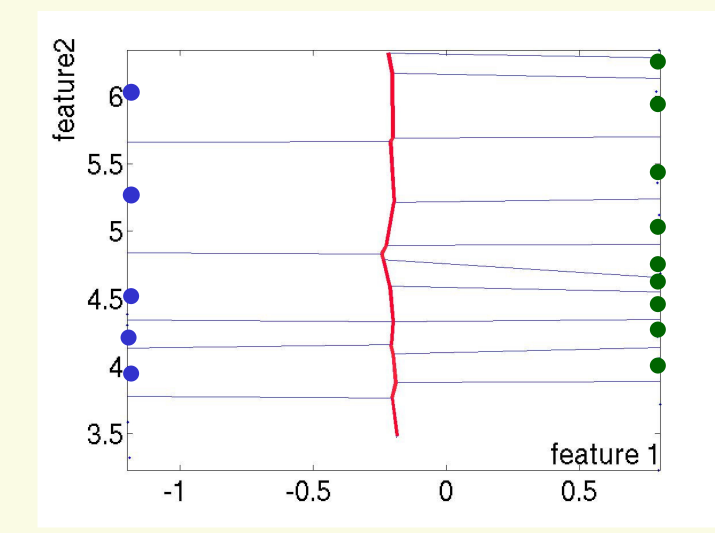
 $D(\begin{bmatrix} 1\\100 \end{bmatrix}, \begin{bmatrix} 1\\150 \end{bmatrix}) = \sqrt{(1-1)^2 + (100-150)^2} = 50 \qquad D(\begin{bmatrix} 1\\100 \end{bmatrix}, \begin{bmatrix} 2\\110 \end{bmatrix}) = \sqrt{(1-2)^2 + (100-110)^2} = 10.5$

- x = [1 100] is misclassified!
- The denser the samples, the less of the problem
 - But we rarely have samples dense enough

kNN: Selection of Distance

- Notice the 2 features are on different scales:
 - feature 1 takes values between 1 or 2
 - feature 2 takes values between 100 to 200
- We could normalize each feature to be between of mean 0 and variance 1
- If **X** is a random variable of mean μ and varaince σ^2 , then $(\mathbf{X} \mu)/\sigma$ has mean 0 and variance 1
- Thus for each feature vector x_i, compute its sample mean and variance, and let the new feature be [x_i mean(x_i)]/sqrt[var(x_i)]
- Let's do it in the previous example

kNN: Normalized Features



The decision boundary (in red) is very good now!

kNN: Selection of Distance

 However in high dimensions if there are a lot of irrelevant features, normalization will not help

$$D(a,b) = \sqrt{\sum_{k} (a_{k} - b_{k})^{2}} = \sqrt{\sum_{i} (a_{i} - b_{i})^{2} + \sum_{j} (a_{j} - b_{j})^{2}}$$

discriminative noisy
feature features

 If the number of discriminative features is smaller than the number of noisy features, Euclidean distance is dominated by noise

kNN: Feature Weighting

 Scale each feature by its importance for classification

$$D(a,b) = \sqrt{\sum_{k} w_{k} (a_{k} - b_{k})^{2}}$$

- Can learn the weights w_k from the training data
 - Increase/decrease weights until classification improves

kNN Summary

- Advantages
 - Can be applied to the data from any distribution
 - Very simple and intuitive
 - Good classification if the number of samples is large enough
- Disadvantages
 - Choosing best k may be difficult
 - Computationally heavy, but improvements possible
 - Need large number of samples for accuracy
 - Can never fix this without assuming parametric distribution