

Part 3:

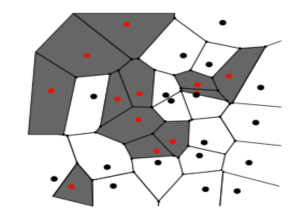
Elements of Non-parametric Techniques: the k-Nearest Neighbor (kNN) Classifier

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Introduction

- In Part 2 we assumed that the forms of the probability density functions were known
 - However, this assumption cannot be made in some pattern recognition applications
- In this chapter, we shall examine a nonparametric method that can be used with arbitrary distributions and without the assumption that the forms of the underlying densities are known
- We will discuss the k-Nearest Neighbor (kNN) pattern classifier, which allows:
 - estimating the density function $p(\mathbf{x} \mid \omega_i)$
 - estimating the posterior probability $P(\omega_i | \mathbf{x})$

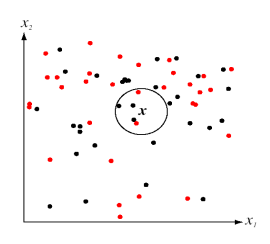
- Nonparametric classification is often associated with the notion of prototype
- We can think of a prototype as a representative element from a class
- The class label assigned to an example is based on the similarity of this example to one or more prototypes
- Similarity is defined in a geometrical sense, that is, based on a certain distance
 - The smaller the distance, the higher the similarity between x and the prototype



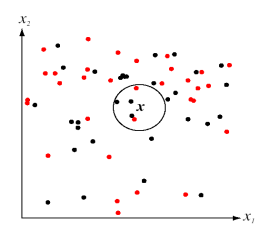
- kNN is one of the most theoretically elegant and simple classification techniques (L. Kuncheva, 2004)
- Let D be a labeled training set containing n points, called prototypes
- Each prototype belongs to one of the "c" classes, i.e., \mathbf{x}_i ϵ ω_j j=1,...,c

$$\mathbf{D} = [\mathbf{x}_1, \mathbf{x}_2,, \mathbf{x}_n]$$

 $\mathbf{x}_i = (\mathbf{x}_{i1}, \mathbf{x}_{i2},, \mathbf{x}_{id}) i=1,...,n$



- To classify an input \mathbf{x} , the k nearest prototypes are retrieved from \mathbf{D} together with their class labels
- The input x is labeled to the most represented class label amongst the k nearest neighbors
- In the figure:
 - x is the pattern to be classified
 - we consider a "region" R of the feature space containing the k nearest prototypes to x
 - we classify \mathbf{x} as belonging to the most represented class label amongst the k nearest neighbors within the region R.



 It can be shown (we see that later) that the kNN method estimates the posterior probabilities as:

$$\widehat{P}(\omega_i|\mathbf{x}) = \frac{k_i}{k}$$

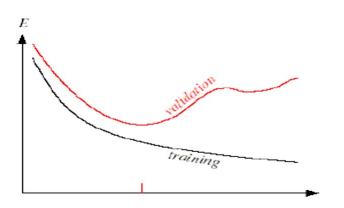
- k_i is the number of nearest neighbors belonging to the class ω_i within region R
- k is the number of the k nearest prototypes of x within region R
- The minimum (Bayes) error classifier using the approximations above will assign \mathbf{x} to the class with the highest posterior probability, i.e., the class most represented among the k nearest neighbors of \mathbf{x} .

How to Select the Right Value of the Hyperparameter *k*

- One simple rule of thumb (heuristic rule) is to select k as $k = \sqrt{n}$
 - n is the number of examples in our training set D
- Note that this rule imposes that the number k of nearest neighbors of the pattern \mathbf{x} that fall within the region R is smaller than the total number n of training set examples
- In binary (two-class) classification problems or problems with an **even** number of classes, it is helpful to choose k to be an odd number as this avoids **tie-breaks**

How to Select the Right Value of the Hyperparameter k

- An experimental method for selecting the value of the «k» parameter is «cross validation» (we see it later in Part 6).
- We subdivide our original data set D into three subsets:
 - training set, validation set, and test set
- We use the training set as the set containing the "prototypes"
- Simple method: we evaluate error E using different values of the «k» parameter with the validation set (more on this later)



• In the next slides, we see the theoretical concepts behind the k-NN method and how one can arrive at the approximation below:

$$\widehat{P}\left(\omega_i|\mathbf{x}\right) = \frac{k_i}{k}$$

Density Estimation for the kNN Method

• The basic idea underlying many non-parametric methods is very simple, and it can be illustrated as follows

• The probability **P** that a vector **x** will fall in a region \mathcal{R} of the feature space is:

$$P = \int_{\Re} p(\mathbf{x'}) d\mathbf{x'}$$

- if R is a small region, P can be regarded as a smoothed or averaged version of the density function p(x)
- We can thus estimate this smoothed value of p(x) with P

Density Estimation

- Suppose that n samples $\mathbf{x}_1,...,\mathbf{x}_n$ are drawn independently and identically distributed (i.i.d.) according to the probability law $p(\mathbf{x})$.
- If we know that \mathbf{k} samples of these \mathbf{n} fall in R, then \mathbf{P} can be estimated simply as $\mathbf{P} = \mathbf{k}/\mathbf{n}$. In general, this can be proved as follows.
- The probability that \mathbf{k} of the \mathbf{n} samples fall in R is given by the binomial law:

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

• We know that the expected value of the binomial law for k is : $\varepsilon(k) = nP$

Density Estimation

• If we consider the ratio k/n as argument of the binomial distribution, we can rewrite the expected value for k/n as:

$$\varepsilon(k/n) = P$$

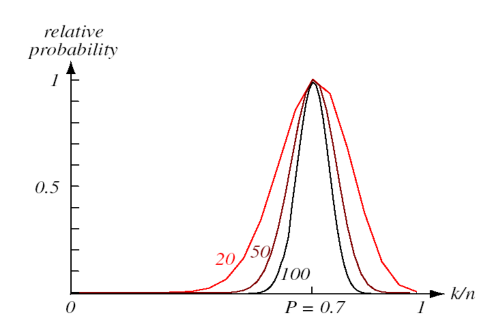
- We know that the variance of the binomial law is var(k/n)=P(1-P)/n
- If the number of samples n increases, the limit when it goes to infinity is:

$$n \to +\infty \to \varepsilon(k/n) = P$$
 and $var(k/n)=0$

- Therefore, we can say that k/n is an asymptotically unbiased estimator of P
- We expect that the ratio k/n will be a very good estimate for the probability P, and hence for the smoothed density function if n is very large
 - Indeed, this estimate is especially accurate when n is very large (see next Figure)

Density Estimation: Binomial Distribution

- The plot shows the probability P_k of finding k samples in a region where the averaged probability is P=0.7, as a function of k/n
- Each curve is labeled by the total number of patterns n
- For large n, such binomial distributions peak strongly at k/n = P (here chosen to be 0.7)



Density Estimation

• If we now assume that p(x) is continuous and that **the region R is so small** that p(x) does not vary appreciably within it, we can write:

$$\int_{\Re} p(\mathbf{x'}) d\mathbf{x'} \cong p(\mathbf{x}) V = P$$

- where x is a point within R and V is the volume enclosed by R.
- Combining these two estimates, we get the following expression for p(x):

$$\int_{\Re} p(\mathbf{x}') d\mathbf{x}' \cong p(\mathbf{x}) V \cong k / n \rightarrow p(\mathbf{x}) \cong \frac{k / n}{V}$$

Density Estimation

C. Bishop, Pattern Recognition and Machine Learning, pp. 121-122, 2006

$$p(\mathbf{x}) = \frac{k}{nV}$$

- The validity of the above estimate depends on two contradictory assumptions:
 - 1. the region R has to be small enough that the density is approximately constant over the region
 - 2. and yet sufficiently large (in relation to the value of that density) that the number k of points falling inside the region is sufficient for the binomial distribution to be sharply peaked
- We can exploit this result in two different ways
 - Fix k and determine the value of V from the data, which gives rise to the kNN method
 - Fix V and determine k from data, giving rise to kernel density estimators (Parzen windows)
- It can be shown that both the kNN density estimator and the kernel density estimator converge to the true probability density in the limit $n \to \infty$, provided that V shrinks suitably with n and k grows with n (Duda and Hart, 1973).

Density Estimation: Contradictory Assumptions

$$\int_{\Re} p(\mathbf{x}') d\mathbf{x}' \cong p(\mathbf{x}) V \cong k / n \rightarrow p(\mathbf{x}) \cong \frac{k / n}{V}$$

• If we fix the volume V and take more and more training samples, the ratio k/n will converge (in probability) as desired, but we have only obtained an estimate of the space-averaged value of p(x):

$$\frac{P}{V} = \frac{\int_{\Re} p(\mathbf{x'}) d\mathbf{x'}}{\int_{\Re} d\mathbf{x'}}$$

if we want to obtain p(x) rather than just a smoothed version of it, V has to tend to zero

Density Estimation

- From a practical standpoint, we note that the number of samples is always limited. Thus, the volume V can not be allowed to become arbitrarily small
 - One will have to accept a certain amount of variance in the ratio k/n and a certain amount of averaging of the density p(x)
- From a theoretical standpoint, it is interesting to ask how these limitations can be circumvented if an unlimited number of samples is available
 - We will discuss that in the next slides

- To estimate the density at x, we form a sequence of regions R₁, R₂,... R_n, containing x
 the first region to be used with one sample, the second with two, and so on.
- Let V_n be the volume of R_n , k_n be the number of samples falling in R_n , and $p_n(x)$ be the nth estimate for p(x):

$$p_n(\mathbf{x}) = \frac{k_n/n}{V_n}$$

• If $p_n(x)$ is to converge to p(x), three conditions appear to be required:

$$\lim_{n \to \infty} V_n = 0$$

$$\lim_{n \to \infty} k_n = \infty$$

$$\lim_{n \to \infty} \frac{k_n}{n} = 0$$

- The first condition assures us that the space averaged P/V will converge to p(x), provided that the regions shrink uniformly and that $p(\cdot)$ is continuous at x.
- The second condition, which only makes sense if p(x)≠0, assures us that the frequency ratio will converge (in probability) to the probability P
- The third condition is necessary if $p_n(x)$ is to converge at all. It also says that although a huge number of samples will eventually fall within the small region R_n , they will form a negligibly small fraction of the total number of samples.

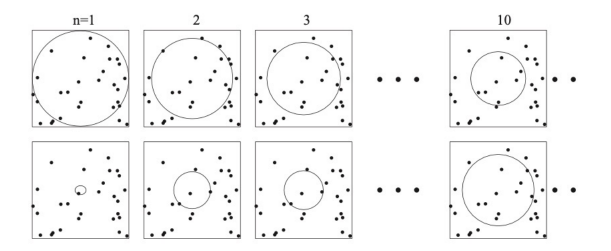
$$\lim_{n\to\infty}V_n=0$$

$$\lim_{n\to\infty} k_n = \infty$$

$$\lim_{n\to\infty}\frac{k_n}{n}=0$$

- There are two common ways of obtaining sequences of regions that satisfy these conditions
- One is to shrink an initial region by specifying the volume V_n as some function of n, such as $V_n = 1/\sqrt{n}$. It then must be shown that the random variables k_n and k_n/n behave properly, or more to the point, that $p_n(x)$ converges to p(x)
 - This is the Parzen-window / kernel density estimation (KDE) method
- The second method is to specify k_n as some function of n, such as $k_n = \sqrt{n}$. Here, the volume V_n is grown until it encloses k_n neighbors of x
 - This is the kNN method
- Both these methods converge in the asymptotic regime of infinite samples, but it is hard to give guarantees in the finite-sample regime

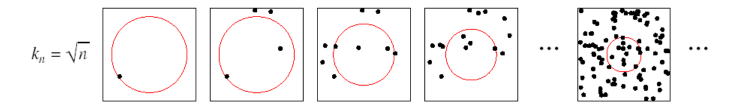
Sequence of Regions for KDE and kNN



k-Nearest Neighbor Method: Value of k

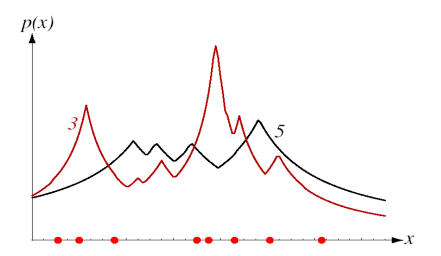
kNN non-parametric method

- This method specifies k_n as some function of n, such as $k_n = \sqrt{n}$. Here, the volume Vn grows until it encloses k_n neighbors of x.
- **Key concept:** the region (volume) is specified considering the number k of the samples that fall into it.



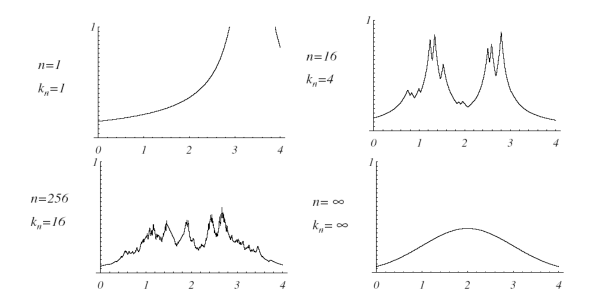
An Example of Density Estimation with kNN

- Eight points in one dimension and the kNN density estimates for k=3 and k=5.
- The discontinuities in the slopes in the estimates generally occur away from the positions of the points themselves



An Example of Density Estimation with kNN

- Several kNN estimates of a Gaussian distribution.
- Note how the finite n estimates can be quite "spiky"



Kernel Density Estimation

Recall that:

$$p(\mathbf{x}) = \frac{k}{nV}$$

- The validity of the above estimate depends on two contradictory assumptions:
 - 1. the region R has to be small enough that the density is approximately constant over the region
 - 2. and yet sufficiently large (in relation to the value of that density) that the number k of points falling inside the region is sufficient for the binomial distribution to be sharply peaked
- We can exploit this result in two different ways
 - Fix k and determine the value of V from the data, which gives rise to the kNN method
 - Fix V and determine k from data, giving rise to kernel density estimators (Parzen windows)

Kernel Density Estimation

C. Bishop, PRML, pp. 123-124, 2006

- Let's consider R to be a small hypercube centered on the origin
 - This is referred to as a kernel function or Parzen window

$$k(\mathbf{u}) = \begin{cases} 1, & |u_i| \leq 1/2, \quad i = 1, \dots, D, \\ 0, & \text{otherwise} \end{cases}$$

- The quantity $k\left(\frac{x-x_n}{h}\right)$ will be one if the data point x_n lies inside a cube of side h centered on x, and zero otherwise
- The total number of data points lying inside this cube will therefore be

$$K = \sum_{n=1}^{N} k \left(\frac{\mathbf{x} - \mathbf{x}_n}{h} \right)$$

Kernel Density Estimation

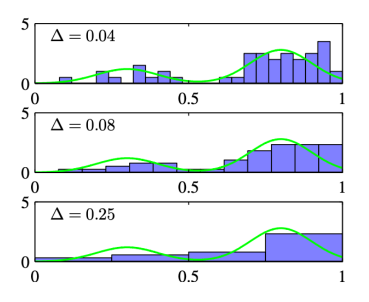
We can thus estimate p(x) as

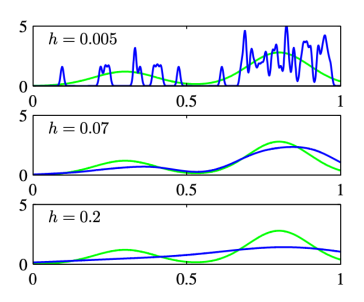
$$p(\mathbf{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{h^{D}} k \left(\frac{\mathbf{x} - \mathbf{x}_{n}}{h} \right)$$

If we consider a Gaussian kernel instead of a hypercube:

$$p(\mathbf{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{(2\pi h^2)^{1/2}} \exp\left\{-\frac{\|\mathbf{x} - \mathbf{x}_n\|^2}{2h^2}\right\}$$

Kernel Density Estimation: 1D Examples





We have discussed density estimation so far. How about classification?

From Density Estimation to Classification

- We can use the kNN or KDE estimators to get an estimate of the likelihood $p(\mathbf{x} \mid y)$
- Then estimate priors to compute the posterior probabilities, using the Bayes' theorem
- ... and decide for the class exhibiting the maximum support (MAP criterion)

• In the previous slides, we have shown that:

$$p_n(\mathbf{x}) = \frac{k_n/n}{V_n}$$

It is easy to show that the kNN method estimates the posterior probabilities as

$$P(\omega_i|\mathbf{x}) = \frac{k_i}{k}$$

• where k_i is the number of nearest neighbors belonging to the class ω_i within the region R

• To arrive at this classification method, we fix k_n and n and allow for a variable V_n . Assuming Euclidean distance, let R be the region containing exactly k_n of the elements of the training set D. We know that the unconditional p.d.f. can be approximated as

$$p_n(\mathbf{x}) = \frac{k_n/n}{V_n}$$

• Denoting by k_i the number of elements in R from class ω_i , the class-conditional pdf for ω_i , i=1,...,c, can be approximated in R, as

$$p(\mathbf{x}/\boldsymbol{\omega}_i) = \frac{k_i/n_i}{V_n}$$

• The posterior probabilities are obtained as

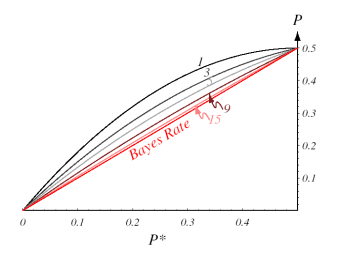
$$P_n(\boldsymbol{\omega}_i \mid \mathbf{x}) = \frac{p_n(\mathbf{x} / \boldsymbol{\omega}_i) P(\boldsymbol{\omega}_i)}{p_n(\mathbf{x})} = \frac{\frac{k_i / n_i}{V} \frac{n_i}{n}}{\frac{k / n}{V}} = \frac{k_i}{k}$$

- The minimum error (Bayes) classifier using the approximations above will assign x to the class with the highest posterior probability, i.e., the class most represented among the k nearest neighbors of x
- The region R and the volume V are specific for each x
 - The kNN classification rule, however, assigns the class label using only the numbers k_i , so the winning label does not depend on V

• kNN is Bayes-optimal when:

$$\lim_{n\to\infty} k_n = \infty$$

$$\lim_{n\to\infty}\frac{k_n}{n}=0$$



• The error rate for the kNN rule for a two-category problem. Each curve is labeled by k; when $k=\infty$, the estimated probabilities match the true probabilities, and thus, the error rate is equal to the Bayes rate, i.e., $P = P_{Bayes}$

The Nearest-Neighbor Rule (1NN)

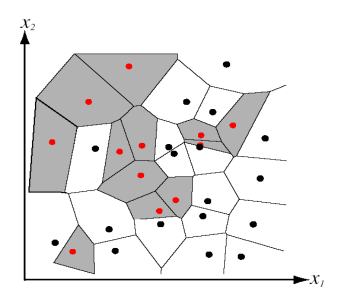
- Let be $D^n=\{x_1,...,x_n\}$ a training set of samples belonging to the "c" classes $\omega_1,...,\omega_c$ and be \mathbf{x} 's D^n the nearest sample to the unknown sample \mathbf{x}
- Decision rule: we assign x to the class of x'
- The nearest-neighbor rule is a sub-optimal procedure
 - that usually leads to an error rate greater than the minimum possible, the Bayes rate.
 - However, with infinite prototypes, the error rate is never worse than twice the Bayes rate.

The Nearest-Neighbor Rule (1NN)

- Let be ω ' the true class of \mathbf{x} '. If "n" is very large, we can assume that \mathbf{x} ' is very close to \mathbf{x} , so that $P(\omega' | \mathbf{x}') = P(\omega_i | \mathbf{x})$
 - Here, the nearest-neighbor decision rule is in good agreement with the MAP rule.
- In general, if we specify $\omega_m(\mathbf{x})$ by $P(\boldsymbol{\omega}_m \mid \mathbf{x}) = \max_i P(\boldsymbol{\omega}_i \mid \mathbf{x})$
- The MAP rule assigns the pattern to the class ω_{m}
 - If \mathbf{x}' has been assigned to the class \mathbf{j} , then we should assume that $\omega_{m}(\mathbf{x}') = \omega_{i}$
- This rule partitions the feature space into cells that contain the closest points to a given training point \mathbf{x}'
 - All points in such a cell are thus labeled as the same class of the training point a so-called Voronoi tesselation of the space (see next slide)

1NN and Voronoi Tesselation: A 2D Example

• In two dimensions, the nearest-neighbor algorithm with k=1 (1-NN) partitions the input space into Voronoi cells, each labeled by the category of the training point it contains



The Nearest-Neighbor Rule: Final Remarks

- If $P(\omega_m \mid \mathbf{x}) \cong 1$, the nearest-neighbor rule is close to the optimal rule, as it is very unlikely that the posterior probability changes abruptly.
 - When the minimum probability of error is small, the nearest-neighbor rule's probability of error is also small.

- If $P(\omega_m \mid \mathbf{x}) \cong 1/c$, the classes have the same probabilities, and the nearest-neighbor rule is likely suboptimal
 - In this case, the error probability is about 1-1/c for both methods.

References

- Sections 4.1, 4.2, 4.3, 4.4, 4.5, Pattern Classification, R. O. Duda, P. E. Hart, D. G. Stork, John Wiley & Sons, 2000
- Chris Bishop, Machine Learning and Pattern Recognition, 2006.
- L. Kuncheva, Combining pattern classifiers, Wiley, 2004.