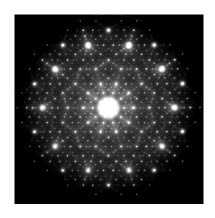
# LTAT.02.004 MACHINE LEARNING II

# Basics of probabilistic modelling

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# What is probability?







Probability is a measure of uncertainty which can rise in several ways

- ▷ Intrinsic uncertainty in the system
- ▷ Uncertainty caused by inherent instability of the system
- ▷ Uncertainty caused by lack of knowledge or control over the system

# Frequentistic interpretation of probability



Probability is an average occurrence rate in long series of experiments.

- ▷ Probability is a collective property
- > Probabilities can be assigned only to future events

# Bayesian interpretation of probability



Probability reflects persons individual beliefs on future or unknown events.

- ▷ Belief updates through the Bayes rule
- > Probability is an inherently subjective property
- > Probabilities can be assigned to past, present and future events

# Ultra-frequentistic interpretation of probability



Events with small enough probability do not occur

- > The main tool in classical statistics
- > Errors in judgement does not matter if a gamma ray pulse kills us.
- ▷ One must avoid the lottery paradox in the reasoning

# The goal of statistical inference

### Frequentist goal

- ▶ The aim of statistics is to design algorithms that work well on average.
- ▶ For that one needs to specify probabilistic model for data sources.
- ▷ Confidence is the fraction of cases the algorithm works as specified.

### Bayesian goal

- ▶ The aim of statistics is to design algorithms that allow rational individuals
   to reliably update their beliefs through Bayes formula
- Besides the data source model one has to provide model for initial beliefs.
- ▷ Correctness of an algorithm does not make sense.

Frequentistic methods

### Causation between zero-one events

Assume that condition A causes the event B=1 with probability p, i.e.,

$$\Pr\left[B=1|A\right]=p$$

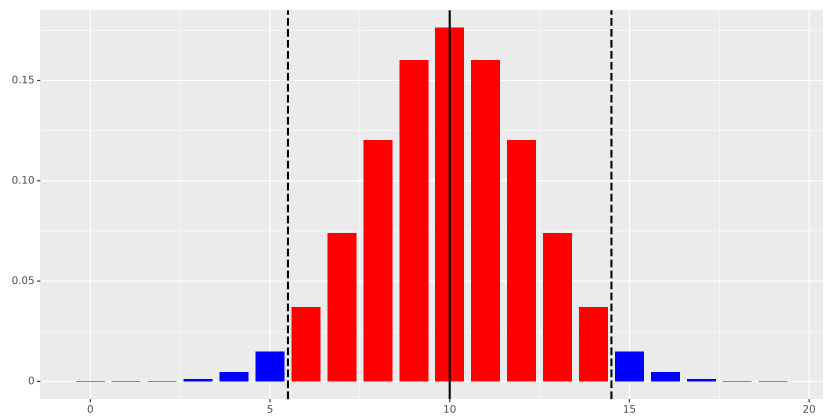
Then the probability is to get k ones in n independent trials is

$$\Pr[B_1 + \dots + B_n = k | A] = \binom{n}{k} p^k (1-p)^{n-k}$$

The number of ones in known to have a binomial distribution

$$B_1 + \cdots + B_n \sim \text{Bin}(n, p)$$

# Illustration



The distribution of  $B_1 + \ldots + B_n$  depends solely on the number of trials n and the probability p. Some values of  $B_1 + \ldots + B_n$  are very unlikely.

### How to build a statistical test

### I. Null hypothesis:

 $\triangleright$  The probability of heads in a coinflip is  $\Pr[B_i = 1] = p$ .

### II. Choose value to compute aka test statistic:

 $\triangleright$  Our test statistic will be  $B_1 + \ldots + B_n$ .

### III. Consequences on the observations:

- $\triangleright$  The observed sum  $B_1 + \ldots + B_n \sim \text{Bin}(n = 20, p = 0.5)$ .
- $\triangleright$  Limit on the tail probability  $\Pr\left[|B_1+\ldots+B_n-10|\geq 6\right]\leq 5\%$

### IV. Test procedure

 $\triangleright$  Reject null hypotesis at *significance level* 5% if  $|B_1 + \ldots + B_n - 10| \ge 6$ .

# Properties of statistical tests

Statistical test is a classification algorithm designed to distinguish a fixed distribution of negative examples specified by a null hypothesis.

Any *static* classification algorithm can be converted to a statistical test by finding out the percentage of false positives aka *p-value*:

- > There might exists a closed form solution.
- ▶ We can always estimate p-values using simulations.
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Testing several hypothesis in parallel increases the number of false positives. Several p-value adjustment methods are used to correct the issue:

- ▷ Bonferroni correction is almost optimal
- > FDR correction controls the expected number false positives

### How to build confidence intervals

### I. Construct a family of statistical tests:

- $\triangleright$  Define a statistical test  $T_p$  for all possible parameter values p.
- ▷ All tests should share the same test statistic.

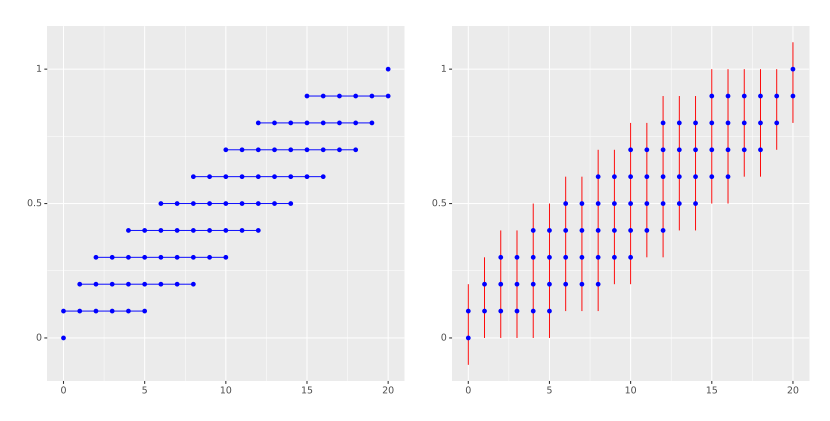
### II. Perform multiple hypotesis testing for all parameter values:

- $\triangleright$  Accept all parameters values for which p-value is greater than  $1-\alpha$ .
- Dutput a minimal interval that covers all accepted parameter values.

### Rationale

- $\triangleright$  The true parameter value is accepted on  $\alpha$ -fraction of possible observations.
- Otherwise, the true value is inside the predicted interval.

### Illustration



- ▷ Acceptance ranges for different parameter values on the left.
- > Extended parameter ranges covering all accepted parameters on the right.
- > These ranges are the desired confidence intervals.

# Interpretation of confidence intervals

**Definition.** Confidence interval for a parameter p is an outcome of an approximation algorithm. The algorithm must output an interval  $[\hat{p}-\varepsilon,\hat{p}+\varepsilon]$  such that the true estimate is in the range on  $\alpha$ -fraction of cases.

### Paradoxical inapplicability

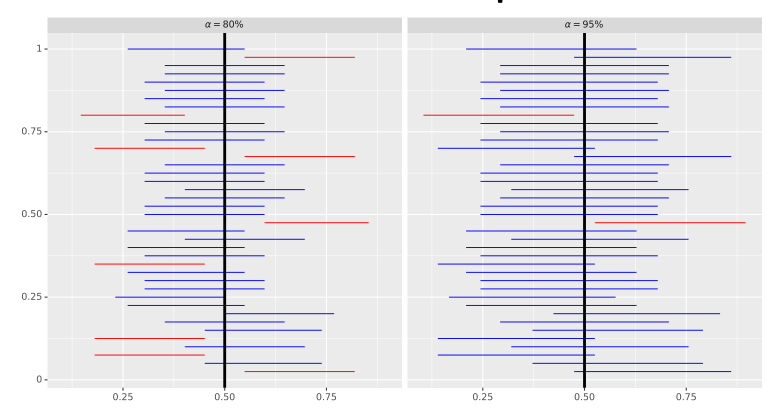
The definition does not state that the probability  $p \in [\hat{p} - \varepsilon, \hat{p} + \varepsilon]$  is  $\alpha!$ 

- ho The statement  $p \in [\hat{p} \varepsilon, \hat{p} + \varepsilon]$  is either true or false.
- ▶ There is no probability left. We just do not know the answer!

### Ultra-frequentistic resolution

 $\triangleright$  If  $1-\alpha$  is small enough say 5% then the algorithm is always correct.

# Illustrative example



By increasing the length of the interval we increase the fraction of runs for which the true value of p lies in the interval.

### Problems with confidence intervals

### Inability to capture background knowledge

- $\triangleright$  What if I know that  $p \in [0.1, 0.2]$  and observe  $B_1 = \ldots = B_N = 1$ ?
- $\triangleright$  Then the estimate  $[\hat{p} \varepsilon, \hat{p} + \varepsilon]$  is clearly wrong although on average this confidence interval is reasonable.

### Multiple hypothesis testing

- □ Using several confidence intervals in parallel increases the fraction of cases where some true estimate is out of the predicted range.
- > We can use p-value adjustment methods are used to correct the issue.

### **Prediction intervals**

Even if we know the true relation y = f(x) we cannot predict the observation  $y_i = f(x_i) + \varepsilon_i$ , as the noise term  $\varepsilon_i$  is not known ahead.

 $\triangleright$  We cannot give upper and lower bounds for  $y_i$  which always hold.

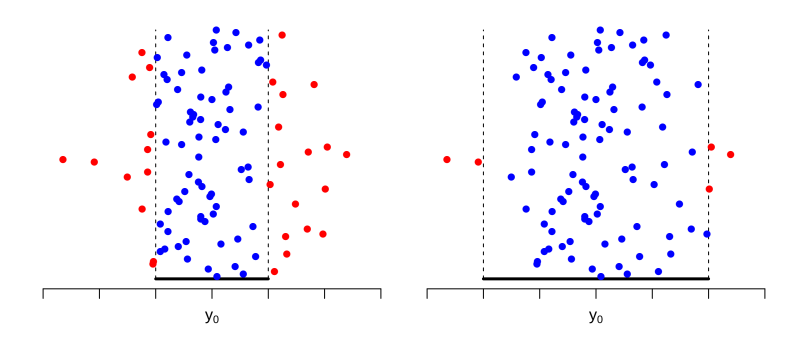
Instead, we can specify a prediction interval  $[y_* - \varepsilon, y_* + \varepsilon]$  so that with probability 95% the resulting measurement  $y_i$  is in the range.

▶ Usually, the analysis is similar to confidence interval derivation.

Interpretation of prediction intervals is different from confidence intervals.

▶ The probability estimate holds for the particular interval.

# Illustrative example



By increasing the length of the prediction interval we increase the fraction of future measurements which fall into interval.

# **Confidence envelopes**

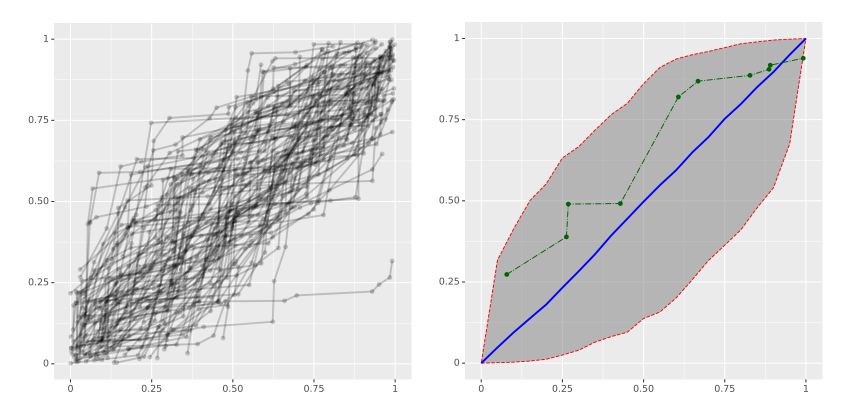
Confidence intervals is a good way to visualise uncertainty of a particular parameter. However, we are sometimes interested in the uncertainty many parameters or in the uncertainty of a function:

- hd How a predictor  $f:[0,1] 
  ightarrow \mathbb{R}$  depends on the training set
- hd How a ROC curve Roc: [0,1] 
  ightarrow [0,1] depends on the test set
- → How should a quantile-quantile plot be distributed.

Confidence bands are generalisations of confidence intervals

- > Pointwise confidence band is a collection of confidence intervals
- $\triangleright$  Simultaneous confidence band must enclose  $\alpha$ -fraction of functions.
- > Simultaneous confidence bands are much wider than pointwise bands.

# Illustrative example



- Distribution of qq-lines visualised through a sample on the left.
- $\triangleright$  A simulation based pointwise 95% confidence envelope on the right.
- $\triangleright$  The significance level that qq-line is inside the envelope is ca 50%.

### **Permutation tests**

### **Baseline problem:**

- > Achievable accuracy depends on the data distribution.
- > Artefacts in the dataset may bias performance measures.

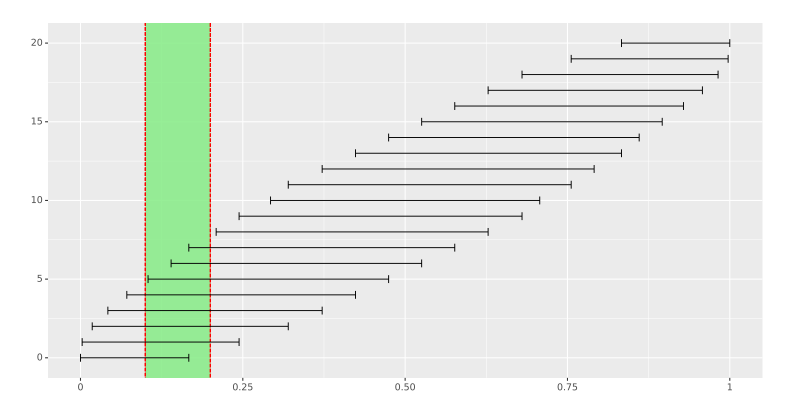
**Label permutation.** A random permutation  $\pi$  on outputs  $y_i$  destroys correlations between input-output pairs  $(\boldsymbol{x}_i, \boldsymbol{y}_{\pi(i)})$  but preserves marginal distribution of inputs and outputs.

**Permutation test.** Estimate how probable is to achieve equal or higher accuracy than was observed on the real data.

- ▷ If this probability is small then there must be signal in the data.
- > The test completely neglect the effect size, i.e., how much results differ.
- Statistical significance does not imply utility!

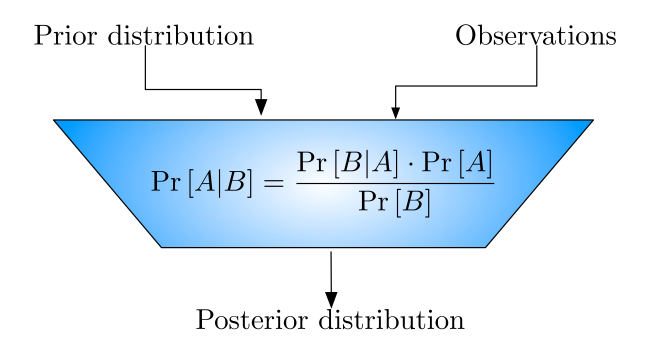
# Bayesian methods

# Confidence intervals vs background knowledge



- $\triangleright$  Confidence intervals do not capture background knowledge  $p \in [0.1, 0.2]$ .
- > Thus we must accept absurd or suboptimal parameter estimations.

# Bayesian inference procedure



- $\triangleright$  Prior distribution  $\Pr[A]$  encodes the background knowledge
- $\triangleright$  The model  $\Pr[B|A]$  determines how the posterior  $\Pr[A|B]$  is updated

### Prior and likelihood

Likelihood  $\mathcal{L}(\mathcal{D}|\mathcal{M})$  is a probability of observations  $\mathcal{D}$  when the data generation model  $\mathcal{M}$  is fixed. The model is fixed by the set of parameters.

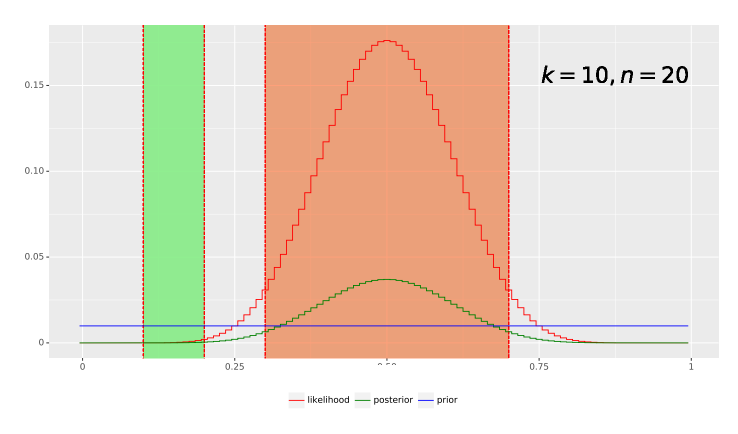
For coin flipping experiment the number of ones k is the observation and the coin bias p is the model parameter and thus

$$\mathcal{L}[k|p] = \binom{n}{k} p^k (1-p)^{n-k}$$

Prior is a distribution over models that encodes our preferences of models before we observe any data.

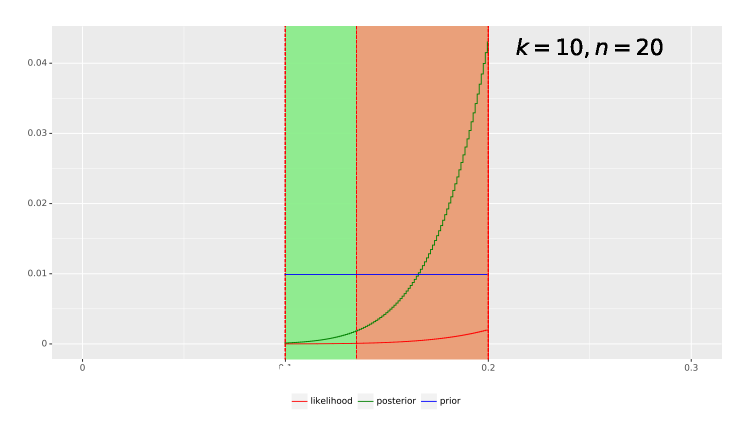
- ▶ Uninformative prior assigns uniform probability to all models.
- ▶ Uninformative prior is not well-defined for continuous parameters.

# Posterior of an uninformed person



- $\triangleright$  Credibility interval  $p \in [0.3, 0.7]$  contains 95% of posterior probability.

# Posterior of an informed person



- $\triangleright$  Credibility interval  $p \in [0.135, 0.2]$  contains 95% of posterior probability.

## Beta distribution as a posterior

By increasing the number of grid points in the non-informative prior we reach a continuous distribution with a density function

$$p[p|k] = \frac{\Gamma(n+2)}{\Gamma(k+1)\Gamma(n-k+1)} \cdot p^k (1-p)^{n-k} .$$

This distribution is known as *beta distribution* Beta $(\alpha = k+1, \beta = n-k+1)$ . The parameter value that maximises the posterior is

$$p_* = \frac{\alpha - 1}{\beta - \alpha} = \frac{k}{n} .$$

# Dice throwing vs coin flipping

A behaviour of a dice with faces  $\{1,\ldots,m\}$  is determined by probabilities

$$p_1 = \Pr[D_i = 1], \dots, p_m = \Pr[D_i = m]$$

### Reduction to coin flipping

- $\triangleright$  Let  $B_i$  denote the event that  $D_i = j$ .
- $\triangleright$  Then  $B_1, \ldots, B_n$  is a coinflipping sequence with bias  $\Pr[B_i = 1] = p_j$ .
- ▶ Non-informative prior for dice throwing goes to the non-informative prior.
- ▷ Informative priors can be marginalised to the right format.
- > The same reduction can be done for all faces of the dice.

Caution: Marginal posteriors do not determine the full posterior in general.

# Laplace smoothing

Assume that we throw a dice with m faces and  $B_i$  encodes the event that the dice lands on a specific face. Then it is natural to assign the maximum prior probability to the parameter value  $p_* = \frac{1}{m}$ .

Such prior can be defined through a following though experiment:

- ▶ We start with non-informative prior.
- $\triangleright$  We observe all possible outcomes of the dice  $\alpha$  times.
- ▶ We use the resulting posterior as a prior for real observations.

Thus the posterior can be obtained by starting with non-informative prior and observing  $k + \alpha$  ones among  $n + m\alpha$  throws.

 $\triangleright$  The ratio  $p = \frac{k+\alpha}{n+m\alpha}$  is the maximal aposteriori estimate for p.

### Markov chains

**Definition.** Markov chain with order m is an outcome of a process that outputs correlated observations  $X_1, X_2, \ldots$  in such a way that the probability of the observation  $X_{m+i}$  depends only on the observations  $X_{m+i-1}, \ldots, X_m$ 

**Log-likelihood.** Let  $\mathbf{x} = (x_1, \dots, x_{m+n})$  be a sequence of observations. Then the log-likelihood  $\ell[\mathbf{x}]$  can be expressed

$$\ell[x] = \log \Pr[x_1, \dots, x_m] + \sum_{i=1}^n \log \Pr[x_{m+i} | x_{m+i-1}, \dots, x_i]$$

$$\beta[x_1, \dots, x_m] + \sum_{i=1}^n \log \Pr[x_{m+i} | x_{m+i-1}, \dots, x_i]$$

#### where

- $\triangleright$  the tensor  $\beta[\ldots]$  determines initial probabilities;
- $\triangleright$  the tensor  $\alpha[\ldots]$  determines transition probabilities.

# Reduction to the dice throwing experiment

Let  $k(u_1, \ldots, u_{m+1})$  is the count of subsequences  $u_1, \ldots, u_{m+1}$  then

$$\ell[\mathbf{x}] = \log \beta[x_1, \dots, x_m] + \sum_{\mathbf{u}} k(\mathbf{u}) \log \alpha[\mathbf{u}]$$

Let us assume that the posterior is defined

- $\triangleright$  fixing probabilities for slices  $\alpha[u_1,\ldots,u_m,*]$
- $\triangleright$  multiplying these probabilities to get the probability of  $\alpha[\ldots]$

The logarithm of the posterior decomposes into sum of independent terms

$$\sum_{u_{m+1}} k(u_1, \dots, u_{m+1}) \cdot \log \alpha[u_1, \dots, u_{m+1}] + \log p(\alpha[u_1, \dots, u_m, *])$$

This is equivalent to inferring probabilities in a dice throws

### **Hidden Markov models**

**Definition.** Let  $X_1, X_2, \ldots$  be hidden states that form a Markov chain and let  $Y_1, Y_2, \ldots$  be observations that the probability of  $Y_i$  depends only on the state  $X_i$ . Then the entire process is known as Hidden Markov Model.

**Log-likelihood.** Let  $y = (y_1, \dots, y_{m+n})$  and  $x = (x_1, \dots, x_{m+n})$  be the observations and hidden states. Then the complete log likelihood is

$$\ell[x, y] = \log \beta[x_1, \dots, x_m] + \sum_{i=1}^n \log \alpha[x_i, \dots, x_{i+m-1}, x_{m+i}] + \log \delta[x_i, y_i]$$

#### where

- $\triangleright$  the tensor  $\beta[\ldots]$  determines initial probabilities;
- $\triangleright$  the tensor  $\alpha[\ldots]$  determines transition probabilities;
- $\triangleright$  the tensor  $\delta[\ldots]$  determines emission probabilities.

# Reduction to the previous building blocks

The problem simplifies when we know the vector of hidden states x:

- $\triangleright$  Inference of emission probabilities  $\delta[\ldots]$  reduce to dice throwing.
- $\triangleright$  Inference of the chain parameters  $\alpha[\ldots]$  and  $\beta[\ldots]$  is also possible.

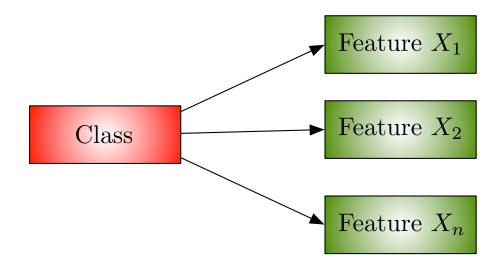
We can use Viterbi algorithm for finding the most probable hidden state x is easy if all parameters are known.

**Naive inference algorithm:** Fix random parameters and repeat steps:

- $\triangleright$  Given parameters  $\alpha, \beta, \delta$  learn the most probable hidden state x.
- $\triangleright$  Given the most probable hidden state  $\boldsymbol{x}$  learn model parameters  $\alpha, \beta, \delta$ .

This algorithm overfits as all hidden states can have similar probability.

# Model behind naive Bayes classifier



Underlying class value determines observed attributes

- $\triangleright$  Each attribute  $X_i$  is binary
- > All variables are independent if class is fixed
- > Sometimes we just ignore dependancies for easier modelling

### Likelihood of the data

Let us assume that we know the probabilities

$$p_i = \Pr\left[X_i = 1 \middle| Class = 0\right]$$

$$q_i = \Pr\left[X_i = 1 \middle| Class = 1\right]$$

Then using the independence assumption we get

$$\Pr\left[X_1 = a_1, \dots, X_n = a_n | Class = 0\right] = \prod_{i=1}^n p_i^{a_i} (1 - p_i)^{1 - a_i}$$

$$\Pr\left[X_1 = a_1, \dots, X_n = a_n | Class = 1\right] = \prod_{i=1}^n q_i^{a_i} (1 - q_i)^{1 - a_i}$$

## Prior and posterior for the class labels

Now it is straightforward to derive

$$\Pr\left[Class = 0 | \boldsymbol{X} = \boldsymbol{a}\right] = \frac{\prod_{i=1}^{n} p_i^{a_i} (1 - p_i)^{1 - a_i} \cdot \Pr\left[Class = 0\right]}{\Pr\left[\boldsymbol{X} = \boldsymbol{a}\right]}$$

$$\Pr\left[Class = 1 | \boldsymbol{X} = \boldsymbol{a}\right] = \frac{\prod_{i=1}^{n} q_i^{a_i} (1 - q_i)^{1 - a_i} \cdot \Pr\left[Class = 1\right]}{\Pr\left[\boldsymbol{X} = \boldsymbol{a}\right]}$$

which gives an odd ratio

$$\frac{\Pr[Class = 0 | \mathbf{X} = \mathbf{a}]}{\Pr[Class = 1 | \mathbf{X} = \mathbf{a}]} = \frac{\Pr[Class = 0]}{\Pr[Class = 1]} \cdot \frac{\prod_{i=1}^{n} p_i^{a_i} (1 - p_i)^{1 - a_i}}{\prod_{i=1}^{n} q_i^{a_i} (1 - q_i)^{1 - a_i}}$$

# The resulting classifier is a linear classifer

By taking logarithm form the odd ratio we get

$$\log \left( \frac{\Pr\left[Class = 0 | \boldsymbol{X} = \boldsymbol{a}\right]}{\Pr\left[Class = 1 | \boldsymbol{X} = \boldsymbol{a}\right]} \right) = w_0 + \sum_{i=1}^n w_i a_i$$

where

$$w_0 = \log\left(\frac{\Pr\left[Class = 0\right]}{\Pr\left[Class = 1\right]}\right) + \sum_{i=1}^n \log\left(\frac{1 - p_i}{1 - q_i}\right)$$
$$w_i = \log\left(\frac{p_i}{1 - p_i} \cdot \frac{1 - q_i}{q_i}\right)$$

### How to train the classifier?

A frequentistic approach is to fix probabilities from the training sample

$$p_i = \frac{\# \left\{ \text{data points form class 0 with } X_i = 1 \right\}}{\# \left\{ \text{data points form class 0} \right\}}$$
 
$$q_i = \frac{\# \left\{ \text{data points form class 1 with } X_i = 1 \right\}}{\# \left\{ \text{data points form class 1} \right\}}$$

However if some value does not occur for  $X_i$  in the training sample we get overly confident results. Thus, Bayesian mean estimate is better alternative

$$p_i = \frac{\# \left\{ \text{data points form class 0 with } X_i = 1 \right\} + 1}{\# \left\{ \text{data points form class 0} \right\} + 2}$$
 
$$q_i = \frac{\# \left\{ \text{data points form class 1 with } X_i = 1 \right\} + 1}{\# \left\{ \text{data points form class 1} \right\} + 2}$$