

Machine Learning - Lecture 18

Repetition

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Bastian Leibe RWTH Aachen http://www.vision.rwth-aachen.de

leibe@vision.rwth-aachen.de



Announcements

- Today, I'll summarize the most important points from the lecture.
 - It is an opportunity for you to ask questions...
 - ...or get additional explanations about certain topics.
 - So, please do ask.

• Today's slides are intended as an index for the lecture.

- > But they are not complete, won't be sufficient as only tool.
- Also look at the exercises they often explain algorithms in detail.



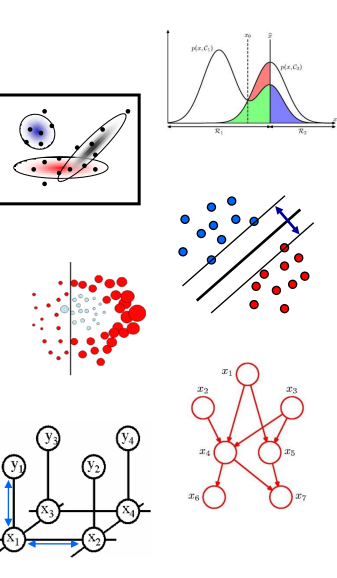
Announcements (2)

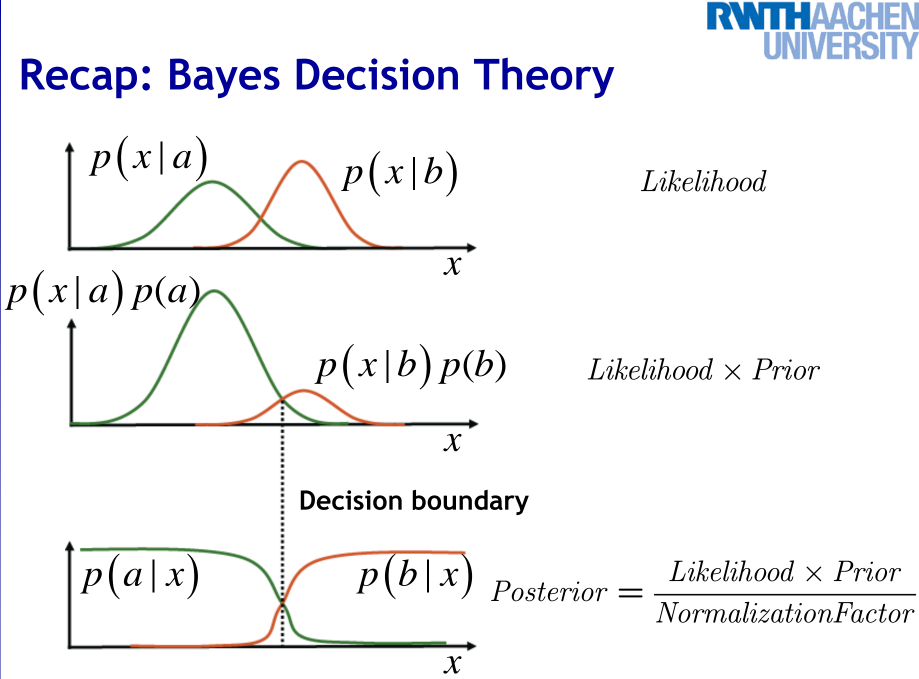
- Test exam on Thursday
 - > During the regular lecture slot
 - Duration: 1h (instead of 2h as for the real exam)
 - > Purpose: prepare you for the questions you can expect
 - All bonus points!

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Course Outline

- Fundamentals
 - Bayes Decision Theory
 - Probability Density Estimation
 - Mixture Models and EM
- Discriminative Approaches
 - > Linear Discriminant Functions
 - Statistical Learning Theory & SVMs
 - Ensemble Methods & Boosting
 - Decision Trees & Randomized Trees
- Generative Models
 - Bayesian Networks
 - Markov Random Fields
 - > Exact Inference





Slide credit: Bernt Schiele

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5 Image source: C.M. Bishop, 2006



Recap: Bayes Decision Theory

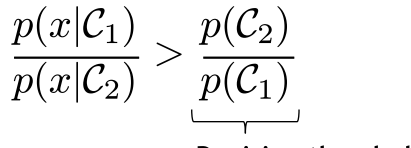
- Optimal decision rule
 - Decide for C₁ if

$$p(\mathcal{C}_1|x) > p(\mathcal{C}_2|x)$$

> This is equivalent to

$$p(x|\mathcal{C}_1)p(\mathcal{C}_1) > p(x|\mathcal{C}_2)p(\mathcal{C}_2)$$

Which is again equivalent to (Likelihood-Ratio test)



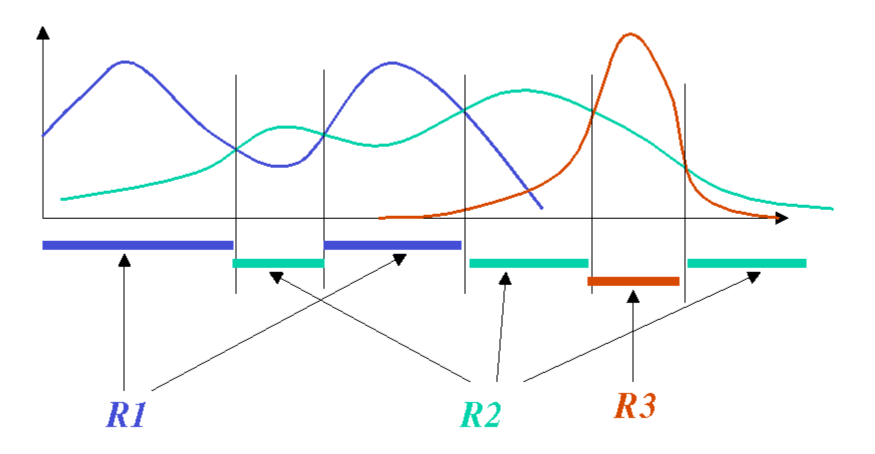
Decision threshold $\boldsymbol{\theta}$

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Recap: Bayes Decision Theory

• Decision regions: \mathcal{R}_1 , \mathcal{R}_2 , \mathcal{R}_{3^c} ...



Recap: Classifying with Loss Functions

- In general, we can formalize this by introducing a loss matrix ${\cal L}_{kj}$

$$L_{kj} = loss for decision C_j if truth is C_k.$$

Example: cancer diagnosis $\begin{array}{c} \text{Decision} \\ \text{cancer normal} \\ L_{cancer diagnosis} = \underbrace{\textbf{F}}_{\textbf{P}} \begin{array}{c} \text{cancer} \\ \text{normal} \end{array} \begin{pmatrix} 0 & 1000 \\ 1 & 0 \end{pmatrix} \end{array}$

Recap: Minimizing the Expected Loss

- Optimal solution minimizes the loss.
 - But: loss function depends on the true class, which is unknown.
- Solution: Minimize the expected loss

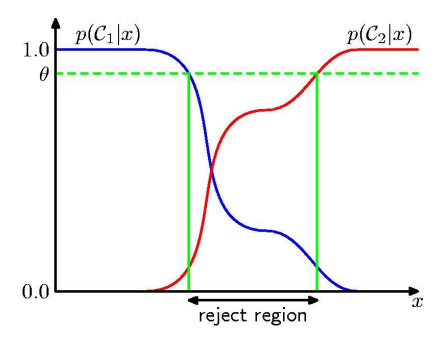
$$\mathbb{E}[L] = \sum_{k} \sum_{j} \int_{\mathcal{R}_{j}} L_{kj} p(\mathbf{x}, \mathcal{C}_{k}) \, \mathrm{d}\mathbf{x}$$

• This can be done by choosing the regions \mathcal{R}_j such that $\mathbb{E}[L] = \sum_k L_{kj} p(\mathcal{C}_k | \mathbf{x})$

which is easy to do once we know the posterior class probabilities $p(C_k|\mathbf{x})$.



Recap: The Reject Option

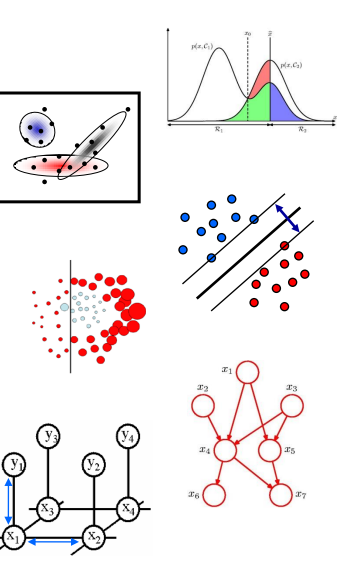


- Classification errors arise from regions where the largest posterior probability $p(C_k|\mathbf{x})$ is significantly less than 1.
 - These are the regions where we are relatively uncertain about class membership.
 - For some applications, it may be better to reject the automatic decision entirely in such a case and e.g. consult a human expert.

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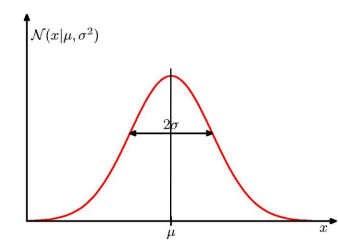
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RWTHAACHEN UNIVERSITY Recap: Gaussian (or Normal) Distribution

- One-dimensional case
 - > Mean μ
 - > Variance σ^2

$$\mathcal{N}(x|\mu,\sigma^2) = \frac{1}{\sqrt{2\pi\sigma}} \exp\left\{-\frac{(x-\mu)^2}{2\sigma^2}\right\}$$



0.16

0.14 0.12 0.1

0.08 0.06 0.04

0.02

- Multi-dimensional case
 - > Mean μ
 - \succ Covariance \varSigma

$$\mathcal{N}(\mathbf{x}|\boldsymbol{\mu}, \boldsymbol{\Sigma}) = \frac{1}{(2\pi)^{D/2} |\boldsymbol{\Sigma}|^{1/2}} \exp\left\{-\frac{1}{2}(\mathbf{x}-\boldsymbol{\mu})^{\mathrm{T}} \boldsymbol{\Sigma}^{-1}(\mathbf{x}-\boldsymbol{\mu})\right\}$$

Recap: Maximum Likelihood Approach

- Computation of the likelihood
 - \succ Single data point: $p(x_n| heta)$
 - Assumption: all data points $X = \{x_1, \dots, x_n\}$ are independent

$$L(\theta) = p(X|\theta) = \prod_{n=1}^{N} p(x_n|\theta)$$

Log-likelihood

$$E(\theta) = -\ln L(\theta) = -\sum_{n=1}^{\infty} \ln p(x_n | \theta)$$

- Estimation of the parameters heta (Learning)
 - Maximize the likelihood (= minimize the negative log-likelihood)

N

 \Rightarrow Take the derivative and set it to zero.

$$\frac{\partial}{\partial \theta} E(\theta) = -\sum_{n=1}^{N} \frac{\frac{\partial}{\partial \theta} p(x_n | \theta)}{p(x_n | \theta)} \stackrel{!}{=} 0$$

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Recap: Bayesian Learning Approach

- Bayesian view:
 - > Consider the parameter vector θ as a random variable.
 - > When estimating the parameters, what we compute is

$$p(x|X) = \int p(x,\theta|X)d\theta$$
Assumption: given θ , this
doesn't depend on X anymore

$$p(x,\theta|X) = p(x|\theta,X)p(\theta|X)$$

$$p(x|X) = \int p(x|\theta)p(\theta|X)d\theta$$

 $p(x|\Lambda) = \int \frac{p(x|\sigma)p(\sigma|\Lambda)d\sigma}{\int \frac{1}{\sqrt{1-1}}}$ This is entirely determined by the parameter θ

(i.e. by the parametric form of the pdf).

Recap: Bayesian Learning Approach

Discussion Likelihood of the parametric form θ given the data set X.

Estimate for x based on parametric form θ

Prior for the parameters θ

$$p(x|X) = \int \frac{p(x|\theta)L(\theta)p(\theta)}{\int L(\theta)p(\theta)d\theta} d\theta$$

Normalization: integrate over all possible values of $\boldsymbol{\theta}$

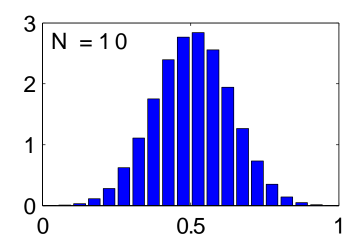
The more uncertain we are about θ , the more we average over all possible parameter values.



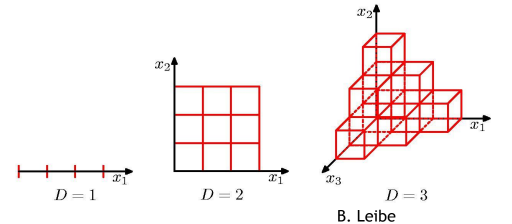
Recap: Histograms

- Basic idea:
 - > Partition the data space into distinct bins with widths Δ_i and count the number of observations, n_i , in each bin.

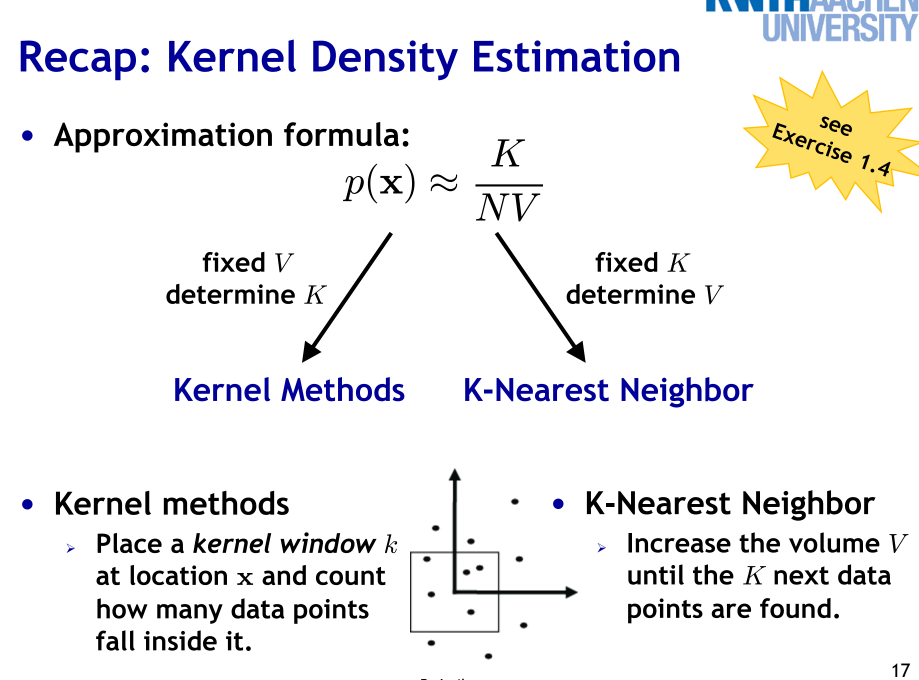
$$p_i = \frac{n_i}{N\Delta_i}$$



- > Often, the same width is used for all bins, $\Delta_i = \Delta$.
- This can be done, in principle, for any dimensionality D...



...but the required number of bins grows exponentially with D!



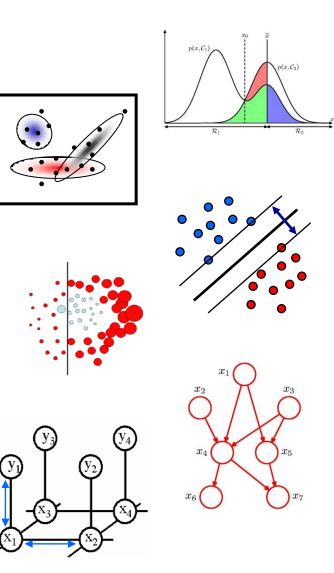
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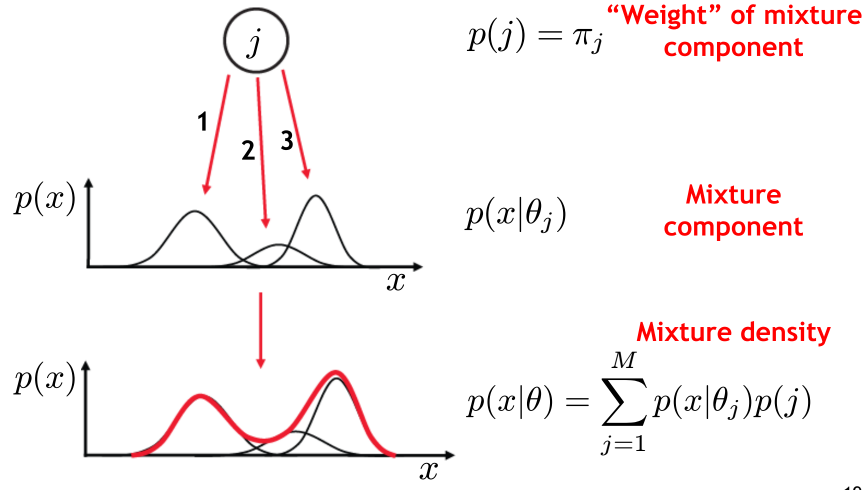
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Recap: Mixture of Gaussians (MoG)

"Generative model"



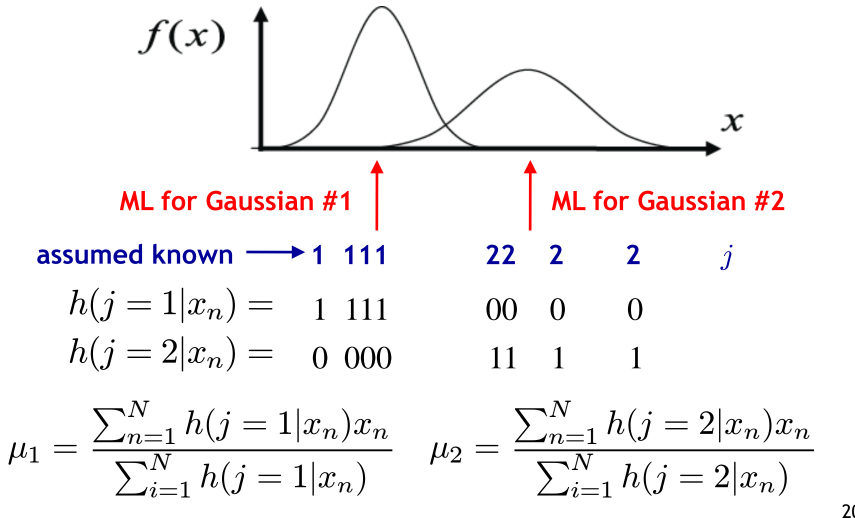
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Recap: MoG - Iterative Strategy

• Assuming we knew the values of the hidden variable...



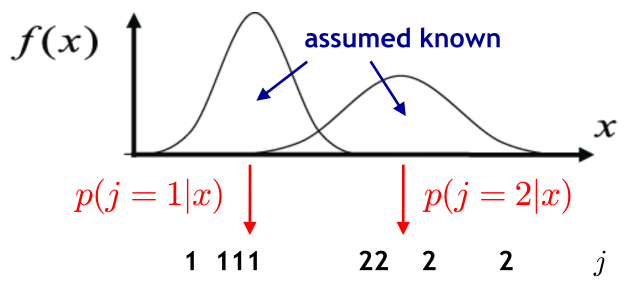
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Recap: MoG - Iterative Strategy

• Assuming we knew the mixture components...



• Bayes decision rule: Decide j = 1 if

$$p(j=1|x_n) > p(j=2|x_n)$$

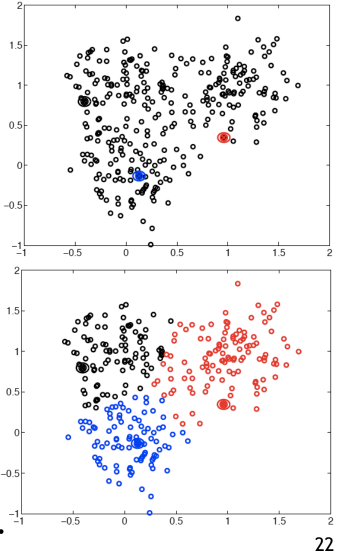


Recap: K-Means Clustering

- Iterative procedure
 - **1.** Initialization: pick *K* arbitrary centroids (cluster means)
 - 2. Assign each sample to the closest centroid.
 - 3. Adjust the centroids to be the means of the samples assigned to them.
 - 4. Go to step 2 (until no change)
 - Algorithm is guaranteed to converge after finite #iterations.
 - Local optimum
 - Final result depends on initialization.



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Recap: EM Algorithm

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- Expectation-Maximization (EM) Algorithm
 - E-Step: softly assign samples to mixture components

$$\gamma_j(\mathbf{x}_n) \leftarrow \frac{\pi_j \mathcal{N}(\mathbf{x}_n | \boldsymbol{\mu}_j, \boldsymbol{\Sigma}_j)}{\sum_{k=1}^N \pi_k \mathcal{N}(\mathbf{x}_n | \boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k)} \quad \forall j = 1, \dots, K, \ n = 1, \dots, N$$

M-Step: re-estimate the parameters (separately for each mixture component) based on the soft assignments

$$\hat{N}_{j} \leftarrow \sum_{n=1}^{N} \gamma_{j}(\mathbf{x}_{n}) = \text{soft number of samples labeled}$$

$$\hat{\pi}_{j}^{\text{new}} \leftarrow \frac{\hat{N}_{j}}{N}$$

$$\hat{\mu}_{j}^{\text{new}} \leftarrow \frac{1}{\hat{N}_{j}} \sum_{n=1}^{N} \gamma_{j}(\mathbf{x}_{n}) \mathbf{x}_{n}$$

$$\hat{\Sigma}_{j}^{\text{new}} \leftarrow \frac{1}{\hat{N}_{j}} \sum_{n=1}^{N} \gamma_{j}(\mathbf{x}_{n}) (\mathbf{x}_{n} - \hat{\mu}_{j}^{\text{new}}) (\mathbf{x}_{n} - \hat{\mu}_{j}^{\text{new}})^{\text{T}}$$

Slide adapted from Bernt Schiele

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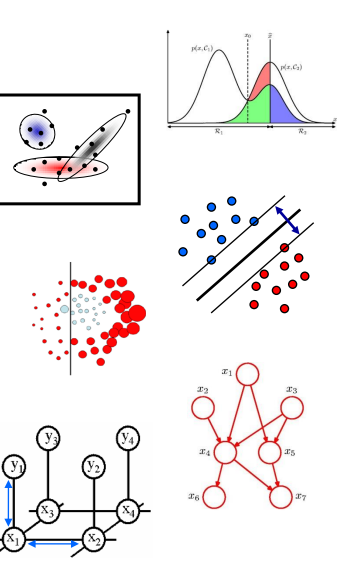
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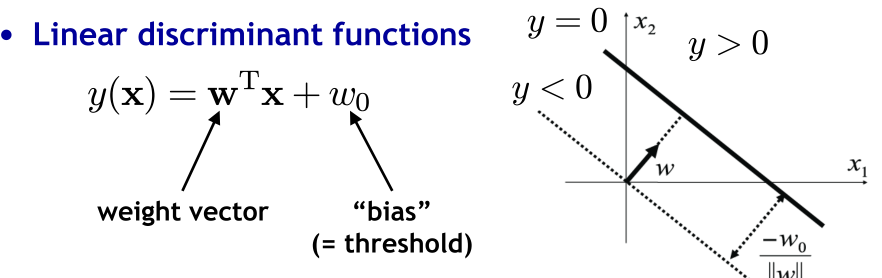
Recap: Linear Discriminant Functions

• Basic idea

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- Directly encode decision boundary
- Minimize misclassification probability directly.



- \succ w, w_{o} define a hyperplane in \mathbb{R}^{D} .
- If a data set can be perfectly classified by a linear discriminant, then we call it linearly separable.

Slide adapted from Bernt Schiele

Recap: Least-Squares Classification

- Simplest approach
 - » Directly try to minimize the sum-of-squares error

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$$E(\mathbf{w}) = \sum_{n=1}^{N} (y(\mathbf{x}_n; \mathbf{w}) - \mathbf{t}_n)^2$$
$$E_D(\widetilde{\mathbf{W}}) = \frac{1}{2} \operatorname{Tr} \left\{ (\widetilde{\mathbf{X}} \widetilde{\mathbf{W}} - \mathbf{T})^{\mathrm{T}} (\widetilde{\mathbf{X}} \widetilde{\mathbf{W}} - \mathbf{T}) \right\}$$

Setting the derivative to zero yields

$$\widetilde{\mathbf{W}} \,=\, (\widetilde{\mathbf{X}}^{\mathrm{T}}\widetilde{\mathbf{X}})^{-1}\widetilde{\mathbf{X}}^{\mathrm{T}}\mathbf{T} = \widetilde{\mathbf{X}}^{\dagger}\mathbf{T}$$

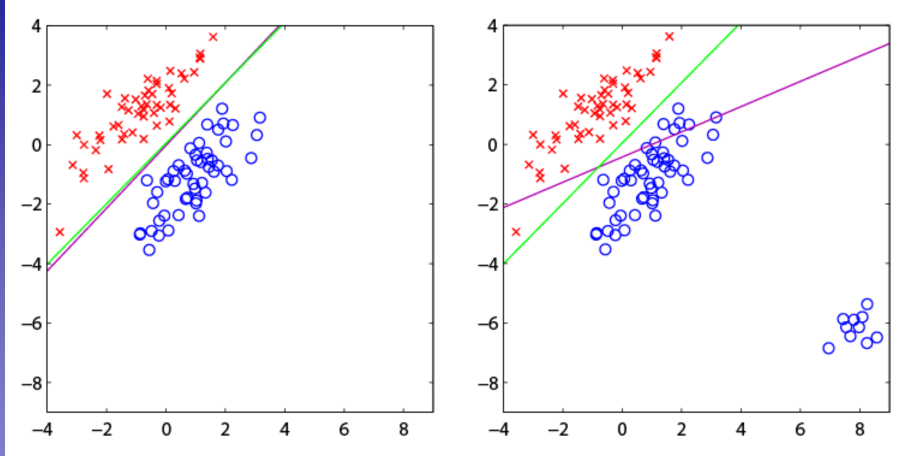
We then obtain the discriminant function as

$$\mathbf{y}(\mathbf{x}) = \widetilde{\mathbf{W}}^{\mathrm{T}} \widetilde{\mathbf{x}} = \mathbf{T}^{\mathrm{T}} \left(\widetilde{\mathbf{X}}^{\dagger} \right)^{\mathrm{T}} \widetilde{\mathbf{x}}$$

➤ ⇒ Exact, closed-form solution for the discriminant function parameters.



Recap: Problems with Least Squares



Least-squares is very sensitive to outliers!

> The error function penalizes predictions that are "too correct".

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Recap: Generalized Linear Models

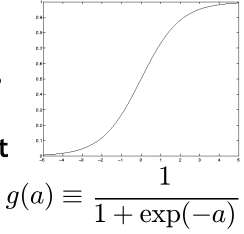
Generalized linear model

 $y(\mathbf{x}) = g(\mathbf{w}^{\mathrm{T}}\mathbf{x} + w_0)$

- > $g(\cdot)$ is called an activation function and may be nonlinear.
- The decision surfaces correspond to

 $y(\mathbf{x}) = const. \quad \Leftrightarrow \quad \mathbf{w}^{\mathrm{T}}\mathbf{x} + w_0 = const.$

- > If g is monotonous (which is typically the case), the resulting decision boundaries are still linear functions of x.
- Advantages of the non-linearity
 - Can be used to bound the influence of outliers and "too correct" data points.
 - > When using a sigmoid for $g(\cdot)$, we can interpret the $y(\mathbf{x})$ as posterior probabilities.

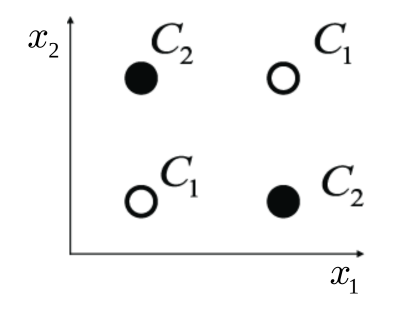




Recap: Linear Separability

- Up to now: restrictive assumption
 - Only consider linear decision boundaries

• Classical counterexample: XOR



Recap: Extension to Nonlinear Basis Fcts.

- Generalization
 - > Transform vector \mathbf{x} with M nonlinear basis functions $\phi_j(\mathbf{x})$:

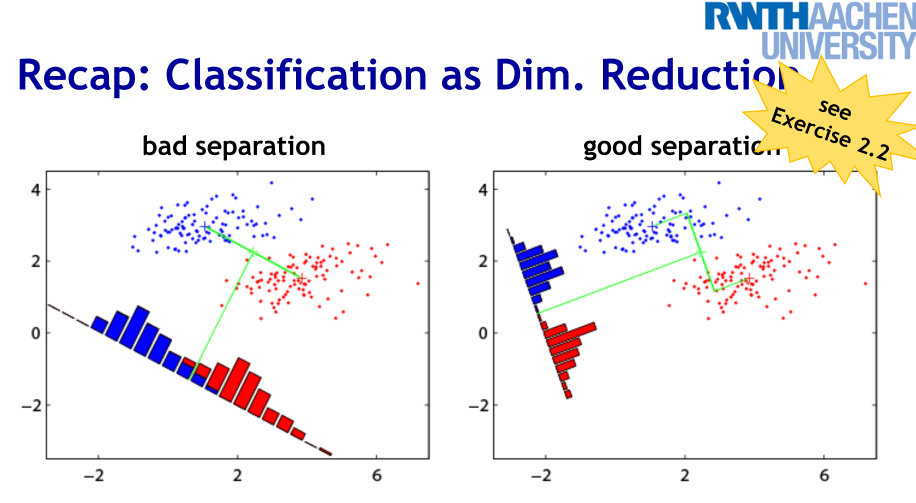
$$y_k(\mathbf{x}) = \sum_{j=1}^{M} w_{ki} \phi_j(\mathbf{x}) + w_{k0}$$

Advantages

- > Transformation allows non-linear decision boundaries.
- > By choosing the right ϕ_j , every continuous function can (in principle) be approximated with arbitrary accuracy.

Disadvatage

- The error function can in general no longer be minimized in closed form.
- \Rightarrow Minimization with Gradient Descent

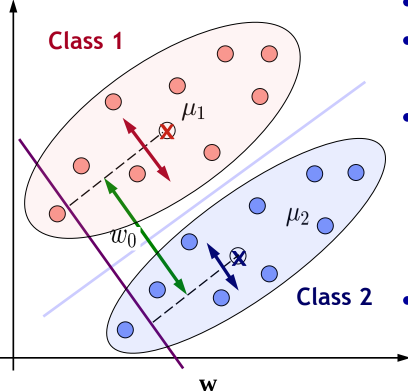


- Classification as dimensionality reduction
 - > Interpret linear classification as a projection onto a lower-dim. space. $y = \mathbf{w}^{\mathrm{T}} \mathbf{x}$
 - \Rightarrow Learning problem: Try to find the projection vector w that maximizes class separation.

Image source: C.M. Bishop, 2006

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RWTHAACHEN UNIVERSITY Recap: Fisher's Linear Discriminant Analysis



- Maximize distance between classes
- Minimize distance within a class

• Criterion:
$$J(\mathbf{w}) = \frac{\mathbf{w}^{\mathrm{T}} \mathbf{S}_{B} \mathbf{w}}{\mathbf{w}^{\mathrm{T}} \mathbf{S}_{W} \mathbf{w}}$$

 $S_B \dots$ between-class scatter matrix $S_W \dots$ within-class scatter matrix

• The optimal solution for w can be obtained as:

$$\mathbf{w} \propto \mathbf{S}_W^{-1}(\mathbf{m}_2-\mathbf{m}_1)$$
 ,

• Classification function:

$$y(\mathbf{x}) = \mathbf{w}^T \mathbf{x} + w_0 \mathop{\gtrless}\limits_{ ext{Class 1}}^{ ext{Class 1}} 0$$

where $w_0 = -\mathbf{w}^T \mathbf{m}$

Slide adapted from Ales Leonardis

RWTHAACHEN UNIVERSITY Recap: Probabilistic Discriminative Models

Consider models of the form

with
$$p(\mathcal{C}_1|oldsymbol{\phi}) ~=~ y(oldsymbol{\phi}) = \sigma(\mathbf{w}^Toldsymbol{\phi})$$

 $p(\mathcal{C}_2|oldsymbol{\phi}) ~=~ 1 - p(\mathcal{C}_1|oldsymbol{\phi})$

• This model is called logistic regression.

Properties

- Probabilistic interpretation
- > But discriminative method: only focus on decision hyperplane
- > Advantageous for high-dimensional spaces, requires less parameters than explicitly modeling $p(\phi | C_k)$ and $p(C_k)$.



Recap: Logistic Regression

• Let's consider a data set $\{\phi_n, t_n\}$ with n = 1, ..., N, where $\phi_n = \phi(\mathbf{x}_n)$ and $t_n \in \{0, 1\}$, $\mathbf{t} = (t_1, ..., t_N)^T$.

• With
$$y_n = p(\mathcal{C}_1 | \phi_n)$$
, we can write the likelihood as $p(\mathbf{t} | \mathbf{w}) = \prod_{n=1}^N y_n^{t_n} \{1 - y_n\}^{1 - t_n}$

- Define the error function as the negative log-likelihood $E(\mathbf{w}) = -\ln p(\mathbf{t}|\mathbf{w})$ $= -\sum_{n=1}^{N} \{t_n \ln y_n + (1 - t_n) \ln(1 - y_n)\}$
 - > This is the so-called cross-entropy error function.

Recap: Iterative Methods for Estimation

• Gradient Descent (1st order)

$$\mathbf{w}^{(\tau+1)} = \mathbf{w}^{(\tau)} - \eta \left. \nabla E(\mathbf{w}) \right|_{\mathbf{w}^{(\tau)}}$$

- Simple and general
- Relatively slow to converge, has problems with some functions
- Newton-Raphson (2nd order) $\mathbf{w}^{(\tau+1)} = \mathbf{w}^{(\tau)} - \eta \mathbf{H}^{-1} \nabla E(\mathbf{w}) \big|_{\mathbf{w}^{(\tau)}}$

where $\mathbf{H} = \nabla \nabla E(\mathbf{w})$ is the Hessian matrix, i.e. the matrix of second derivatives.

- Local quadratic approximation to the target function
- Faster convergence

Recap: Iteratively Reweighted Least Squares

• Update equations

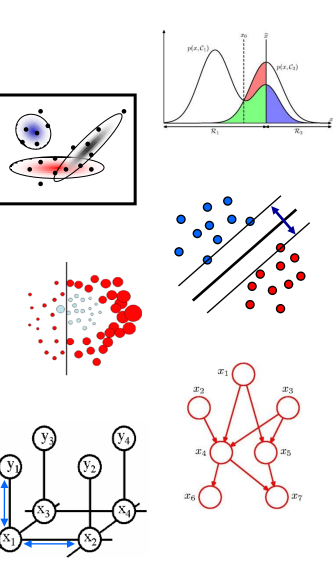
$$\begin{split} \mathbf{w}^{(\tau+1)} &= \mathbf{w}^{(\tau)} - (\mathbf{\Phi}^T \mathbf{R} \mathbf{\Phi})^{-1} \mathbf{\Phi}^T (\mathbf{y} - \mathbf{t}) \\ &= (\mathbf{\Phi}^T \mathbf{R} \mathbf{\Phi})^{-1} \left\{ \mathbf{\Phi}^T \mathbf{R} \mathbf{\Phi} \mathbf{w}^{(\tau)} - \mathbf{\Phi}^T (\mathbf{y} - \mathbf{t}) \right\} \\ &= (\mathbf{\Phi}^T \mathbf{R} \mathbf{\Phi})^{-1} \mathbf{\Phi}^T \mathbf{R} \mathbf{z} \\ & \text{with} \quad \mathbf{z} = \mathbf{\Phi} \mathbf{w}^{(\tau)} - \mathbf{R}^{-1} (\mathbf{y} - \mathbf{t}) \end{split}$$

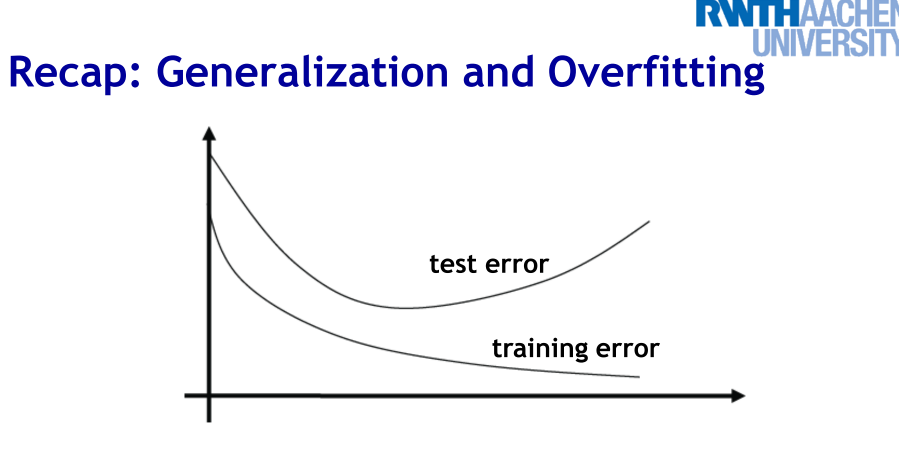
- Very similar form to pseudo-inverse (normal equations)
 - \succ But now with non-constant weighing matrix ${f R}$ (depends on ${f w}$).
 - Need to apply normal equations iteratively.
 - \Rightarrow Iteratively Reweighted Least-Squares (IRLS)

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- Goal: predict class labels of new observations
 - > Train classification model on limited training set.
 - The further we optimize the model parameters, the more the training error will decrease.
 - However, at some point the test error will go up again.
 - \Rightarrow Overfitting to the training set!



Recap: Risk

- Empirical risk
 - Measured on the training/validation set

$$R_{emp}(\alpha) = \frac{1}{N} \sum_{i=1}^{N} L(y_i, f(\mathbf{x}_i; \alpha))$$

- Actual risk (= Expected risk)
 - Expectation of the error on all data.

$$R(\alpha) = \int L(y_i, f(\mathbf{x}; \alpha)) dP_{X,Y}(\mathbf{x}, y)$$

- > $P_{X,Y}(\mathbf{x},y)$ is the probability distribution of (\mathbf{x},y) . It is fixed, but typically unknown.
- \Rightarrow In general, we can't compute the actual risk directly!

Recap: Statistical Learning Theory

• Idea

Compute an upper bound on the actual risk based on the empirical risk

$$R(\alpha) \cdot R_{emp}(\alpha) + \epsilon(N, p^*, h)$$

- > where
 - $N\!\!:\!$ number of training examples
 - \boldsymbol{p}^* : probability that the bound is correct
 - *h*: capacity of the learning machine ("VC-dimension")

Recap: VC Dimension

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- Vapnik-Chervonenkis dimension
 - > Measure for the capacity of a learning machine.
- Formal definition:
 - > If a given set of ℓ points can be labeled in all possible 2^{ℓ} ways, and for each labeling, a member of the set $\{f(\alpha)\}$ can be found which correctly assigns those labels, we say that the set of points is shattered by the set of functions.
 - > The VC dimension for the set of functions $\{f(\alpha)\}$ is defined as the maximum number of training points that can be shattered by $\{f(\alpha)\}$.





Recap: Upper Bound on the Risk

- Important result (Vapnik 1979, 1995)
 - » With probability $(1-\eta)$, the following bound holds

$$R(\alpha) \cdot R_{emp}(\alpha) + \sqrt{\frac{h(\log(2N/h) + 1) - \log(\eta/4)}{N}}$$

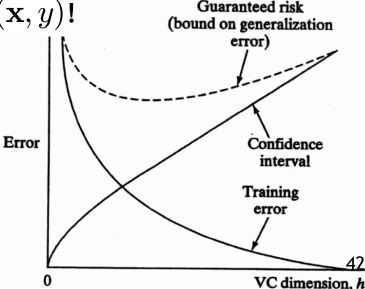
"VC confidence"

- > This bound is independent of $P_{X,Y}(\mathbf{x},y)$!
- If we know h (the VC dimension), we can easily compute the risk bound

$$R(\alpha) \cdot R_{emp}(\alpha) + \epsilon(N, p^*, h)$$

Slide adapted from Bernt Schiele





Recap: Structural Risk Minimization

How can we implement Structural Risk Minimization?

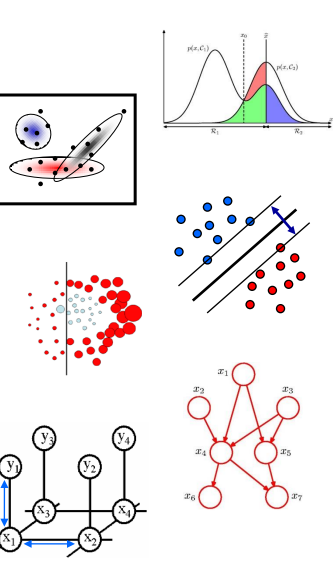
$$R(\alpha) \cdot R_{emp}(\alpha) + \epsilon(N, p^*, h)$$

- Classic approach
 - » Keep $\epsilon(N,p^*,h)$ constant and minimize $R_{emp}(lpha)$.
 - $\succ \epsilon(N,p^*,h)$ can be kept constant by controlling the model parameters.
- Support Vector Machines (SVMs)
 - $ightarrow \, {\sf Keep} \, R_{emp}(lpha)$ constant and minimize $\epsilon(N,p^*,h)$.
 - > In fact: $R_{emp}(\alpha)=0$ for separable data.
 - > Control $\epsilon(N, p^*, h)$ by adapting the VC dimension (controlling the "capacity" of the classifier).

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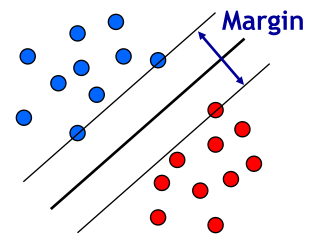
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Recap: Support Vector Machine (SVM)

- Basic idea
 - The SVM tries to find a classifier which maximizes the margin between pos. and neg. data points.
 - > Up to now: consider linear classifiers

$$\mathbf{w}^{\mathrm{T}}\mathbf{x} + b = 0$$



- Formulation as a convex optimization problem
 - Find the hyperplane satisfying

$$\operatorname*{arg\,min}_{\mathbf{w},b} \frac{1}{2} \|\mathbf{w}\|^2$$

under the constraints

$$t_n(\mathbf{w}^{\mathrm{T}}\mathbf{x}_n+b) \ge 1 \quad \forall n$$

based on training data points \mathbf{x}_n and target values $t_n \in \{-1, 1\}$.



Recap: SVM - Primal Formulation

Lagrangian primal form

$$L_{p} = \frac{1}{2} \|\mathbf{w}\|^{2} - \sum_{n=1}^{N} a_{n} \{t_{n}(\mathbf{w}^{\mathrm{T}}\mathbf{x}_{n} + b) - 1\}$$
$$= \frac{1}{2} \|\mathbf{w}\|^{2} - \sum_{n=1}^{N} a_{n} \{t_{n}y(\mathbf{x}_{n}) - 1\}$$

- The solution of L_p needs to fulfill the KKT conditions
 - Necessary and sufficient conditions

$$a_n \ge 0$$

 $t_n y(\mathbf{x}_n) - 1 \ge 0$

$$a_n \left\{ t_n y(\mathbf{x}_n) - 1 \right\} = 0$$

KKT:
$$\lambda \geq 0$$
 $f(\mathbf{x}) \geq 0$ $\lambda f(\mathbf{x}) = 0$

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Recap: SVM - Solution

- Solution for the hyperplane
 - Computed as a linear combination of the training examples

$$\mathbf{w} = \sum_{n=1}^{N} a_n t_n \mathbf{x}_n$$

- Sparse solution: $a_n \neq 0$ only for some points, the support vectors \Rightarrow Only the SVs actually influence the decision boundary!
- Compute b by averaging over all support vectors:

$$b = \frac{1}{N_{\mathcal{S}}} \sum_{n \in \mathcal{S}} \left(t_n - \sum_{m \in \mathcal{S}} a_m t_m \mathbf{x}_m^{\mathrm{T}} \mathbf{x}_n \right)$$



Recap: SVM - Support Vectors

Origi

- The training points for which a_n > 0 are called "support vectors".
- Graphical interpretation:
 - The support vectors are the points on the margin.
 - They define the margin and thus the hyperplane.
 - ⇒ All other data points can be discarded!

0

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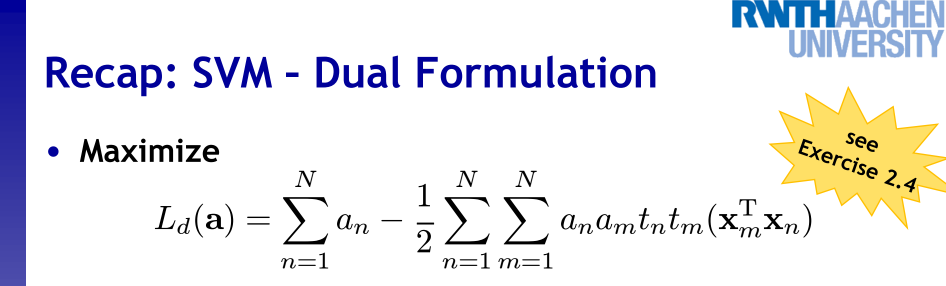
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w

Slide adapted from Bernt Schiele



under the conditions

$$a_n \geq 0 \quad orall n$$

 $\sum_{n=1}^N a_n t_n = 0$

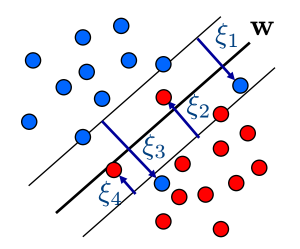
Comparison

- > L_d is equivalent to the primal form L_p , but only depends on a_n .
- > L_p scales with $\mathcal{O}(D^3)$.
- > L_d scales with $\mathcal{O}(N^3)$ in practice between $\mathcal{O}(N)$ and $\mathcal{O}(N^2)$.

Slide adapted from Bernt Schiele

Recap: SVM for Non-Separable Data

- Slack variables
 - > One slack variable $\xi_n \ge 0$ for each training data point.
- Interpretation
 - > $\xi_n = 0$ for points that are on the correct side of the margin.
 - > $\xi_n = |t_n y(\mathbf{x}_n)|$ for all other points.



Point on decision boundary: $\xi_n = 1$

Misclassified point: $\xi_n>1$

- We do not have to set the slack variables ourselves!
- \Rightarrow They are jointly optimized together with w.

Recap: SVM - New Dual Formulation

New SVM Dual: Maximize

SVM Dual: Maximize

$$L_d(\mathbf{a}) = \sum_{n=1}^{N} a_n - \frac{1}{2} \sum_{n=1}^{N} \sum_{m=1}^{N} a_n a_m t_n t_m(\mathbf{x}_m^{\mathrm{T}} \mathbf{x}_n)$$

λT

λT

under the conditions

$$0 \cdot a_n \cdot C$$
$$\sum_{n=1}^N a_n t_n = 0$$

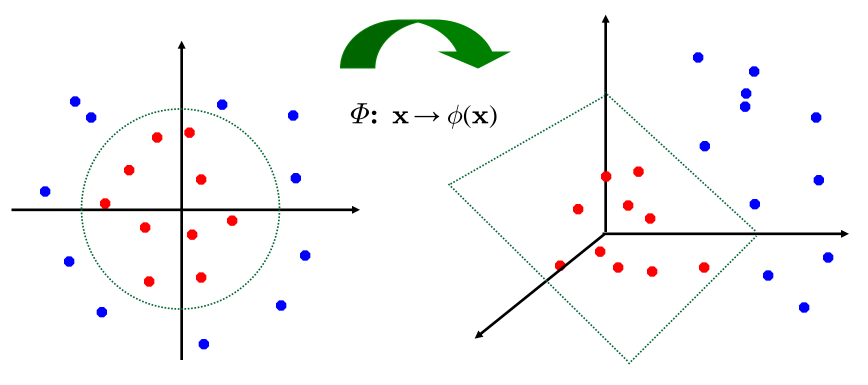
This is all that changed! See

This is again a quadratic programming problem \Rightarrow Solve as before...



Recap: Nonlinear SVMs

 General idea: The original input space can be mapped to some higher-dimensional feature space where the training set is separable:





Recap: The Kernel Trick

- Important observation
 - > $\phi(\mathbf{x})$ only appears in the form of dot products $\phi(\mathbf{x})^{\mathsf{T}}\phi(\mathbf{y})$:

$$y(\mathbf{x}) = \mathbf{w}^{\mathrm{T}} \phi(\mathbf{x}) + b$$
$$= \sum_{n=1}^{N} a_n t_n \phi(\mathbf{x}_n)^{\mathrm{T}} \phi(\mathbf{x}) + b$$

- > Define a so-called kernel function $k(\mathbf{x},\mathbf{y}) = \phi(\mathbf{x})^{\mathsf{T}}\phi(\mathbf{y})$.
- Now, in place of the dot product, use the kernel instead:

$$y(\mathbf{x}) = \sum_{n=1}^{N} a_n t_n k(\mathbf{x}_n, \mathbf{x}) + b$$

> The kernel function *implicitly* maps the data to the higherdimensional space (without having to compute $\phi(\mathbf{x})$ explicitly)!

RWTHAACHEN UNIVERSITY Recap: Kernels Fulfilling Mercer's Condition

Polynomial kernel

$$k(\mathbf{x}, \mathbf{y}) = (\mathbf{x}^{\mathrm{T}}\mathbf{y} + 1)^{p}$$

Radial Basis Function kernel

$$k(\mathbf{x}, \mathbf{y}) = \exp\left\{-rac{(\mathbf{x} - \mathbf{y})^2}{2\sigma^2}
ight\}$$
 e.g. Gaussian

Hyperbolic tangent kernel

$$k(\mathbf{x},\mathbf{y}) = anh(\kappa \mathbf{x}^{\mathrm{T}}\mathbf{y} + \delta)$$
 e.g. Sigmoid

And many, many more, including kernels on graphs, strings, and symbolic data...

RWTHAACHEN UNIVERSITY Recap: Kernels Fulfilling Mercer's Condition

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Hyperbolic tangent kernel

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 e.g. Sigmoid

Actually, that was wrong in the original SVM paper...

And many, many more, including kernels on graphs, strings, and symbolic data...

Recap: Nonlinear SVM - Dual Formulation

SVM Dual: Maximize

Dual: Maximize

$$L_d(\mathbf{a}) = \sum_{n=1}^{N} a_n - \frac{1}{2} \sum_{n=1}^{N} \sum_{m=1}^{N} a_n a_m t_n t_m k(\mathbf{x}_m, \mathbf{x}_n)$$

under the conditions

$$\begin{array}{rcl}
0 \cdot & a_n \cdot & C \\
\sum_{n=1}^N a_n t_n &= & 0
\end{array}$$

• Classify new data points using

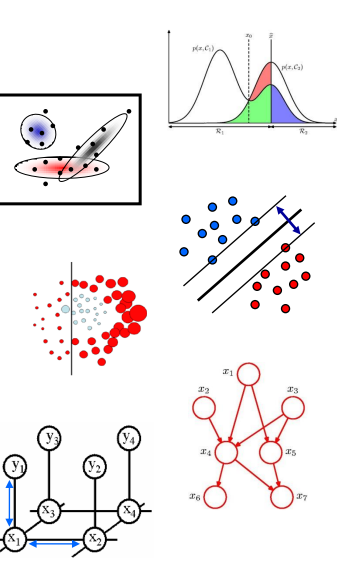
$$y(\mathbf{x}) = \sum_{n=1}^{N} a_n t_n \mathbf{k}(\mathbf{x}_n, \mathbf{x}) + b$$

See

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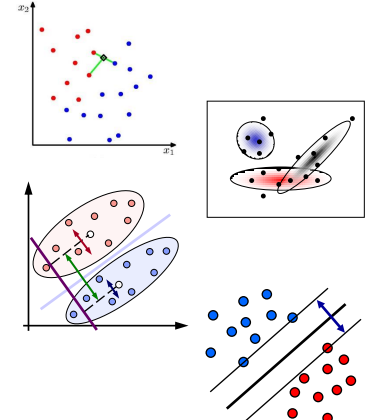




Recap: Classifier Combination

- We've seen already a variety of different classifiers
 - > k-NN
 - Bayes classifiers
 - Fisher's Linear Discriminant

> SVMs



- Each of them has their strengths and weaknesses...
 - Can we improve performance by combining them?



Combination

Classifier

Recap: Stacking

- Idea
 - > Learn L classifiers (based on the training data)
 - Find a meta-classifier that takes as input the output of the L first-level classifiers.
 Classifier 1

Data

Classifier 2

Classifier L



- Learn L classifiers with leave-one-out.
- > Interpret the prediction of the L classifiers as L-dimensional feature vector.
- Learn "level-2" classifier based on the examples generated this way.



Recap: Stacking

- Why can this be useful?
 - Simplicity
 - We may already have several existing classifiers available.
 - \Rightarrow No need to retrain those, they can just be combined with the rest.
 - Correlation between classifiers
 - The combination classifier can learn the correlation.
 - \Rightarrow Better results than simple Naïve Bayes combination.
 - Feature combination
 - E.g. combine information from different sensors or sources (vision, audio, acceleration, temperature, radar, etc.).
 - We can get good training data for each sensor individually, but data from all sensors together is rare.
 - \Rightarrow Train each of the L classifiers on its own input data. Only combination classifier needs to be trained on combined input.



Recap: Bayesian Model Averaging

- Model Averaging
 - Suppose we have H different models h = 1, ..., H with prior probabilities p(h).
 - Construct the marginal distribution over the data set

$$p(\mathbf{X}) = \sum_{h=1}^{n} p(\mathbf{X}|h) p(h)$$

- Average error of committee $\mathbb{E}_{COM} = \frac{1}{M} \mathbb{E}_{AV}$
 - > This suggests that the average error of a model can be reduced by a factor of M simply by averaging M versions of the model!
 - Unfortunately, this assumes that the errors are all uncorrelated.
 In practice, they will typically be highly correlated.

Recap: AdaBoost - "Adaptive Boosting"

• Main idea

[Freund & Schapire, 1996]

- Instead of resampling, reweight misclassified training examples.
 - Increase the chance of being selected in a sampled training set.
 - Or increase the misclassification cost when training on the full set.

Components

- > $h_m(\mathbf{x})$: "weak" or base classifier
 - Condition: <50% training error over any distribution
- > $H(\mathbf{x})$: "strong" or final classifier

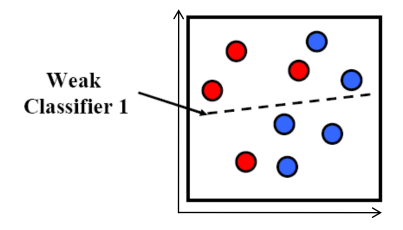
• AdaBoost:

Construct a strong classifier as a thresholded linear combination of the weighted weak classifiers:

$$H(\mathbf{x}) = sign\left(\sum_{\substack{m=1\\B,\ l \ eibe}}^{M} \alpha_m h_m(\mathbf{x})\right)$$



Recap: AdaBoost - Intuition



Consider a 2D feature space with positive and negative examples.

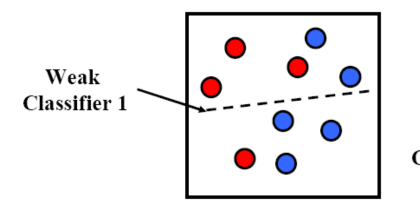
Each weak classifier splits the training examples with at least 50% accuracy.

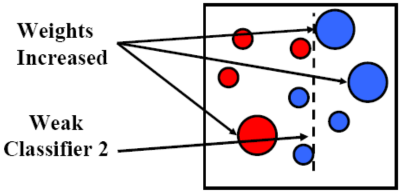
Examples misclassified by a previous weak learner are given more emphasis at future rounds.

Slide credit: Kristen Grauman



Recap: AdaBoost - Intuition

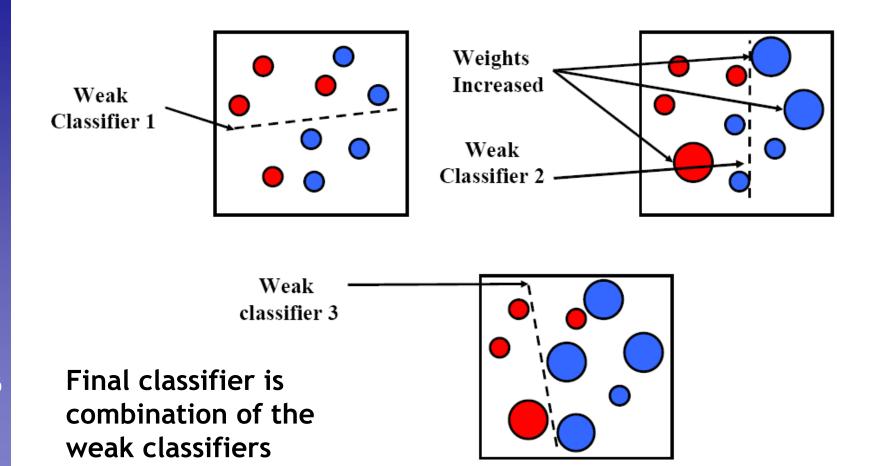




Slide credit: Kristen Grauman



Recap: AdaBoost - Intuition



Slide credit: Kristen Grauman

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Recap: AdaBoost - Algorithm

- **1.** Initialization: Set $w_n^{(1)} = \frac{1}{N}$ for n = 1, ..., N.
- **2.** For $m = 1, \ldots, M$ iterations
 - a) Train a new weak classifier $h_m(\mathbf{x})$ using the current weighting coefficients $\mathbf{W}^{(m)}$ by minimizing the weighted error function

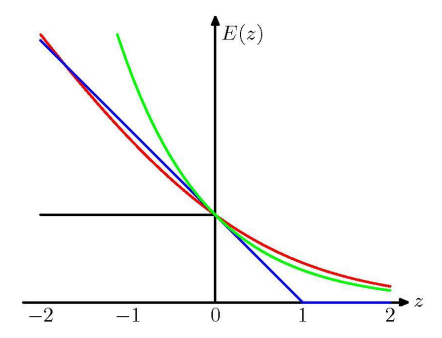
$$J_m = \sum_{n=1}^N w_n^{(m)} I(h_m(\mathbf{x}) \neq t_n) \qquad \qquad I(A) = \begin{cases} 1, & \text{if } A \text{ is true} \\ 0, & \text{else} \end{cases}$$

b) Estimate the weighted error of this classifier on ${f X}$:

$$\epsilon_m = \frac{\sum_{n=1}^N w_n^{(m)} I(h_m(\mathbf{x}) \neq t_n)}{\sum_{n=1}^N w_n^{(m)}}$$

- c) Calculate a weighting coefficient for $h_m(\mathbf{x})$: $\alpha_m = \ln \left\{ \frac{1 - \epsilon_m}{\epsilon_m} \right\}$
- d) Update the weighting coefficients: $w_n^{(m+1)} = w_n^{(m)} \exp \{\alpha_m I(h_m(\mathbf{x}_n) \neq t_n)\}$

Recap: Comparing Error Functions

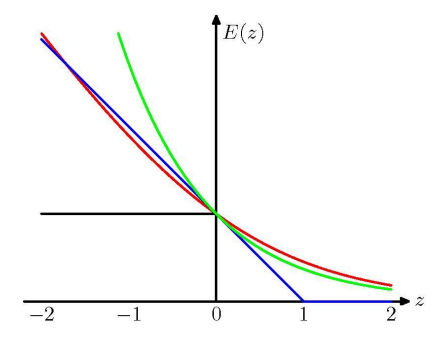


- Ideal misclassification error function
- "Hinge error" used in SVMs
- Exponential error function
 - Continuous approximation to ideal misclassification function.
 - Sequential minimization leads to simple AdaBoost scheme.
 - Disadvantage: exponential penalty for large negative values!
 - \Rightarrow Less robust to outliers or misclassified data points!

Image source: Bishop, 2006

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Recap: Comparing Error Functions



- Ideal misclassification error function
- "Hinge error" used in SVMs
- Exponential error function

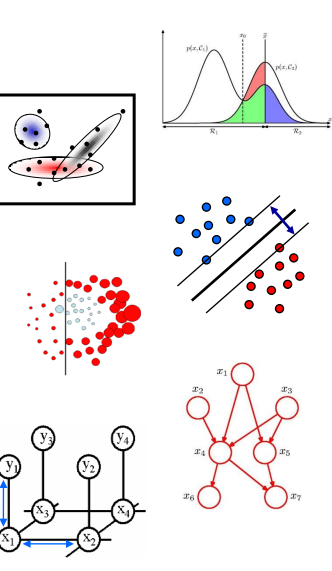
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- "Cross-entropy error" $E = -\sum \{t_n \ln y_n + (1 t_n) \ln(1 y_n)\}$
 - Similar to exponential error for $\overline{z>0}$.
 - Only grows linearly with large negative values of z.
 - $\Rightarrow Make AdaBoost more robust by switching \Rightarrow "GentleBoost" 68$ B. Leibe Image source: Bishop, 2006

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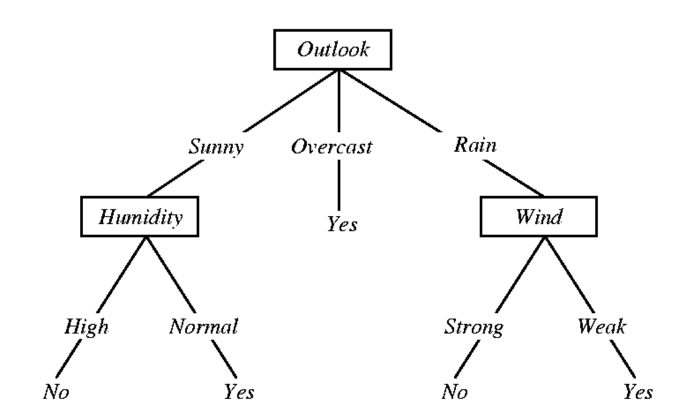
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Recap: Decision Trees



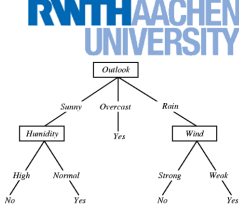
• Example:

Classify Saturday mornings according to whether they're suitable for playing tennis."

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Recap: CART Framework

- Six general questions
 - 1. Binary or multi-valued problem?
 - I.e. how many splits should there be at each node?
 - 2. Which property should be tested at a node?
 - I.e. how to select the query attribute?
 - 3. When should a node be declared a leaf?
 - I.e. when to stop growing the tree?
 - 4. How can a grown tree be simplified or pruned?
 - Goal: reduce overfitting.
 - 5. How to deal with impure nodes?
 - I.e. when the data itself is ambiguous.
 - 6. How should missing attributes be handled?



Recap: Picking a Good Splitting Feature

- Goal
 - Select the query (=split) that decreases impurity the most \geq

$$\triangle i(N) = i(N) - P_L i(N_L) - (1 - P_L)i(N_R)$$

i(P)

Impurity measures

Entropy impurity (information gain):

$$i(N) = -\sum_{j} p(\mathcal{C}_{j}|N) \log_2 p(\mathcal{C}_{j}|N)$$

Gini impurity: ≻

$$i(N) = \sum_{i \neq j} p(\mathcal{C}_i|N) p(\mathcal{C}_j|N) = \frac{1}{2} \left[1 - \sum_j p^2(\mathcal{C}_j|N) \right]$$

Г

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Image source: R.O. Duda, P.E. Hart, D.G. Stork, 2001

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Exercise 3.2

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Recap: Computational Complexity

- Given
 - > Data points $\{\mathbf{x}_1, \dots, \mathbf{x}_N\}$
 - \succ Dimensionality D
- Complexity
 - > Storage: O(N)
 - > Test runtime: $O(\log N)$
 - > Training runtime: $O(DN^2 \log N)$
 - Most expensive part.
 - Critical step: selecting the optimal splitting point.
 - Need to check ${\cal D}$ dimensions, for each need to sort N data points.

 $O(DN \log N)$

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Recap: Decision Trees - Summary

• Properties

- Simple learning procedure, fast evaluation.
- Can be applied to metric, nominal, or mixed data.
- > Often yield interpretable results.

Recap: Decision Trees - Summary

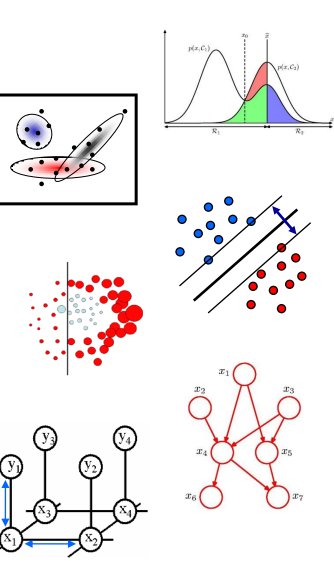
Limitations

- > Often produce noisy (bushy) or weak (stunted) classifiers.
- > Do not generalize too well.
- Training data fragmentation:
 - As tree progresses, splits are selected based on less and less data.
- > Overtraining and undertraining:
 - Deep trees: fit the training data well, will not generalize well to new test data.
 - Shallow trees: not sufficiently refined.
- Stability
 - Trees can be very sensitive to details of the training points.
 - If a single data point is only slightly shifted, a radically different tree may come out!
 - \Rightarrow Result of discrete and greedy learning procedure.
- Expensive learning step
 - Mostly due to costly selection of optimal split.

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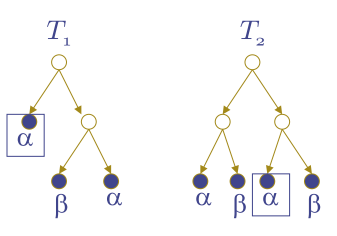
Recap: Randomized Decision Trees

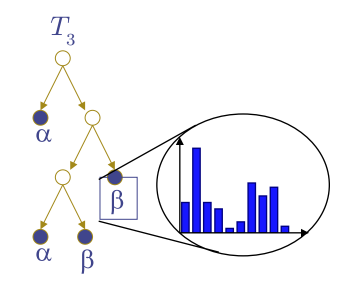
- Decision trees: main effort on finding good split
 - > Training runtime: $O(DN^2 \log N)$
 - This is what takes most effort in practice.
 - > Especially cumbersome with many attributes (large D).
- Idea: randomize attribute selection
 - > No longer look for globally optimal split.
 - > Instead randomly use subset of K attributes on which to base the split.
 - Choose best splitting attribute e.g. by maximizing the information gain (= reducing entropy):

$$\triangle E = \sum_{k=1}^{K} \frac{|S_k|}{|S|} \sum_{j=1}^{N} p_j \log_2(p_j)$$



Recap: Ensemble Combination





- Ensemble combination
 - > Tree leaves (l,η) store posterior probabilities of the target classes. $p_{l,\eta}(\mathcal{C}|\mathbf{x})$
 - Combine the output of several trees by averaging their posteriors (Bayesian model combination)

$$p(\mathcal{C}|\mathbf{x}) = rac{1}{L} \sum_{l=1}^{L} p_{l,\eta}(\mathcal{C}|\mathbf{x})$$
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Recap: Random Forests (Breiman 2001)

- General ensemble method
 - > Idea: Create ensemble of many (50 1,000) trees.
- Empirically very good results
 - > Often as good as SVMs (and sometimes better)!
 - > Often as good as Boosting (and sometimes better)!
- Injecting randomness
 - Bootstrap sampling process
 - On average only 63% of training examples used for building the tree
 - Remaining 37% out-of-bag samples used for validation.
 - Random attribute selection
 - Randomly choose subset of K attributes to select from at each node.
 - Faster training procedure.
- Simple majority vote for tree combination

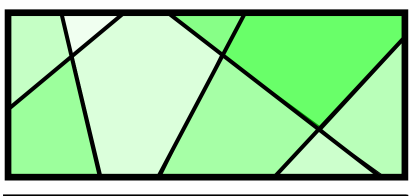
Exercise 3.5

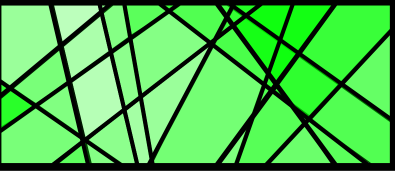


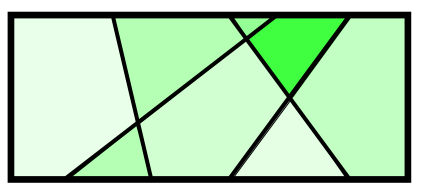
Recap: A Graphical Interpretation

Different trees induce different partitions on the data.

By combining them, we obtain a finer subdivision of the feature space...





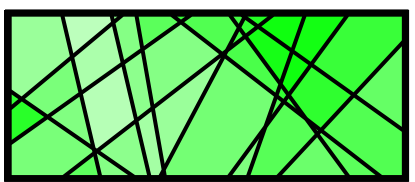




Recap: A Graphical Interpretation

Different trees induce different partitions on the data.

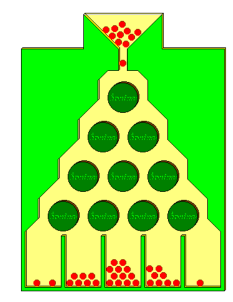
By combining them, we obtain a finer subdivision of the feature space...



...which at the same time also better reflects the uncertainty due to the bootstrapped sampling.

Recap: Extremely Randomized Decision Trees

- Random queries at each node...
 - Tree gradually develops from a classifier to a flexible container structure.
 - Node queries define (randomly selected) structure.
 - Each leaf node stores posterior probabilities
 - Learning
 - Patches are "dropped down" the trees.
 - Only pairwise pixel comparisons at each node.
 - Directly update posterior distributions at leaves
 - \Rightarrow Very fast procedure, only few pixel-wise comparisons.
 - \Rightarrow No need to store the original patches!

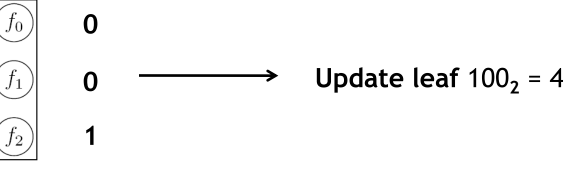


Recap: Ferns

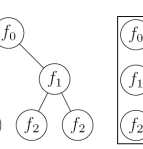
- Ferns
 - Ferns are semi-naïve Bayes classifiers.
 - They assume independence between sets of features (between the ferns)...
 - …and enumerate all possible outcomes inside each set.

Interpretation

- > Combine the tests f_l, \ldots, f_{l+S} into a binary number.
- > Update the "fern leaf" corresponding to that number.







 f_1

 f_2

 f_2



Recap: Ferns (Semi-Naïve Bayes Classifiers)

- Ferns
 - > A fern F is defined as a set of S binary features $\{f_l, \dots, f_{l+S}\}$.
 - > M: number of ferns, $N_f = S \cdot M$.
 - > This represents a compromise:

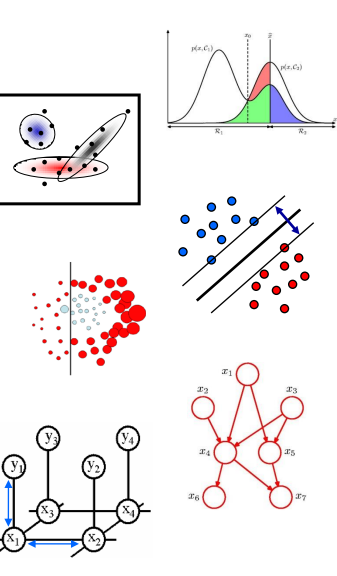
$$\begin{split} p(f_1,\ldots,f_{N_f}|\mathcal{C}_k) &\approx \prod_{j=1}^M p(F_j|\mathcal{C}_k) \\ &= \underbrace{p(f_1,\ldots,f_S|\mathcal{C}_k)}_{\text{Full joint inside fern}} \cdot \underbrace{p(f_{S+1},\ldots,f_{2S}|\mathcal{C}_k)}_{\text{Naïve Bayes between ferns}} \end{split}$$

⇒ Model with $M \cdot 2^S$ parameters ("Semi-Naïve"). ⇒ Flexible solution that allows complexity/performance tuning.

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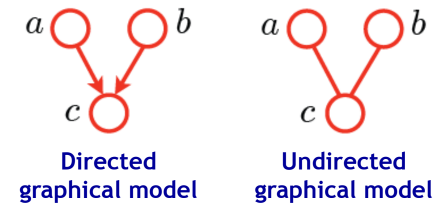


Recap: Graphical Models

- Two basic kinds of graphical models
 - » Directed graphical models or Bayesian Networks
 - > Undirected graphical models or Markov Random Fields
- Key components
 - Nodes
 - Random variables
 - > Edges

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Directed or undirected



known

The value of a random variable may be known or unknown.

Slide credit: Bernt Schiele

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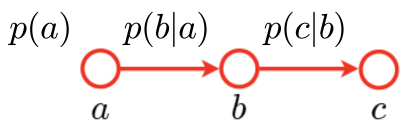
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Recap: Directed Graphical Models

• Chains of nodes:



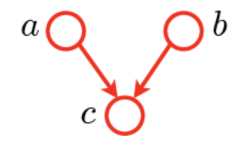
- > Knowledge about a is expressed by the prior probability: p(a)
- > Dependencies are expressed through conditional probabilities: $p(b|a), \ p(c|b)$
- Joint distribution of all three variables:

$$p(a, b, c) = p(c|a, b)p(a, b)$$
$$= p(c|b)p(b|a)p(a)$$



Recap: Directed Graphical Models

Convergent connections:



- > Here the value of c depends on both variables a and b.
- This is modeled with the conditional probability:

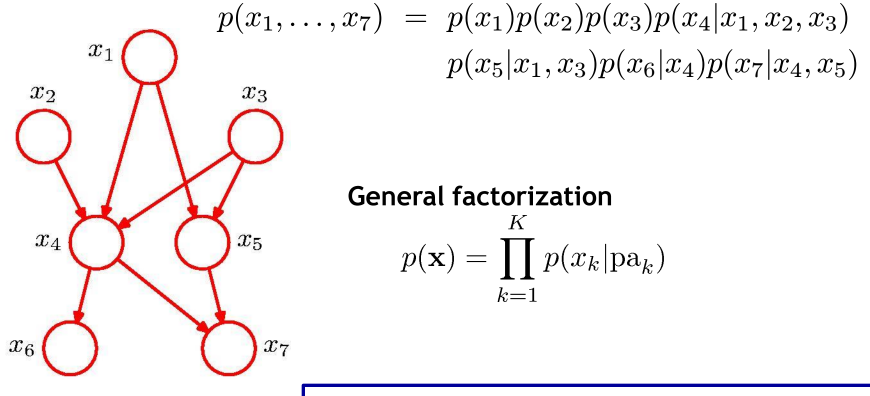
p(c|a,b)

> Therefore, the joint probability of all three variables is given as:

$$p(a, b, c) = p(c|a, b)p(a, b)$$
$$= p(c|a, b)p(a)p(b)$$

RWTHAACHEN UNIVERSITY Recap: Factorization of the Joint Probability

Computing the joint probability



We can directly read off the factorization of the joint from the network structure!



Recap: Factorized Representation

- Reduction of complexity
 - Joint probability of n binary variables requires us to represent values by brute force

 $\mathcal{O}(2^n)$ terms

The factorized form obtained from the graphical model only requires

 $\mathcal{O}(n\cdot 2^k)$ terms

-k: maximum number of parents of a node.

⇒ It's the edges that are missing in the graph that are important! They encode the simplifying assumptions we make.

Recap: Conditional Independence

- X is conditionally independent of Y given V
 - > Definition: $X \perp \!\!\!\perp Y | V \iff p(X|Y,V) = p(X|V)$
 - > Also: $X \perp\!\!\!\perp Y | V \Leftrightarrow p(X, Y | V) = p(X | V) p(Y | V)$
 - » Special case: Marginal Independence

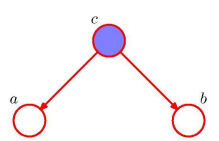
 $X \bot\!\!\!\!\perp Y \ \Leftrightarrow \ X \bot\!\!\!\!\perp Y | \emptyset \ \Leftrightarrow \ p(X,Y) = p(X) \, p(Y)$

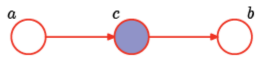
> Often, we are interested in conditional independence between sets of variables:

 $\mathcal{X} \perp \mathcal{Y} \mid \mathcal{V} \iff \{X \perp \mathcal{Y} \mid \mathcal{V}, \forall X \in \mathcal{X} \text{ and } \forall Y \in \mathcal{Y}\}$

Recap: Conditional Independence

- Three cases
 - Divergent ("Tail-to-Tail")
 - Conditional independence when c is observed.
 - > Chain ("Head-to-Tail")
 - Conditional independence when c is observed.
 - Convergent ("Head-to-Head")
 - Conditional independence when neither c, nor any of its descendants are observed.







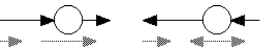
Recap: D-Separation

- Definition
 - Let A, B, and C be non-intersecting subsets of nodes in a directed graph.
 - A path from A to B is blocked if it contains a node such that either
 - The arrows on the path meet either head-to-tail or tail-to-tail at the node, and the node is in the set C, or
 - The arrows meet head-to-head at the node, and neither the node, nor any of its descendants, are in the set C.
 - > If all paths from A to B are blocked, A is said to be d-separated from B by C.
- If A is d-separated from B by C, the joint distribution over all variables in the graph satisfies $A \perp\!\!\!\perp B \mid C$.
 - > Read: "A is conditionally independent of B given C."

Exercise 4.

Recap: "Bayes Ball" Algorithm

- Graph algorithm to compute d-separation
 - Goal: Get a ball from X to Y without being blocked by \mathcal{V} . \geq
 - Depending on its direction and the previous node, the ball can
 - Pass through (from parent to all children, from child to all parents)
 - **Bounce back** (from any parent/child to all parents/children)
 - Be blocked
- Game rules
 - > An unobserved node ($W \notin \mathcal{V}$) passes through balls from parents, but also bounces back balls from children.



> An observed node ($W \in \mathcal{V}$) bounces back balls from parents, but blocks balls from children.

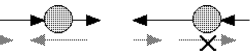
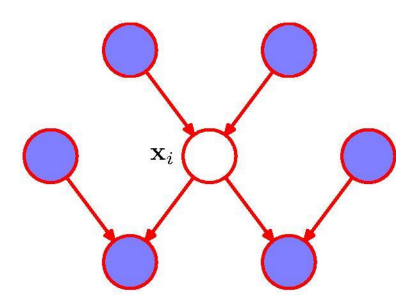






Image source: C. Bishop, 2006

Recap: The Markov Blanket



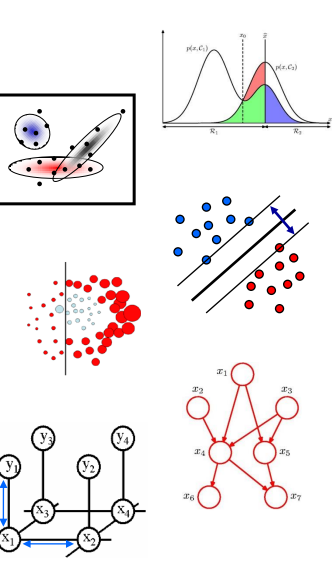
Markov blanket of a node \mathbf{x}_i

- > Minimal set of nodes that isolates \mathbf{x}_i from the rest of the graph.
- This comprises the set of
 - Parents,
 - Children, and
 - Co-parents of \mathbf{x}_i . \leftarrow This is what we have to watch out for!

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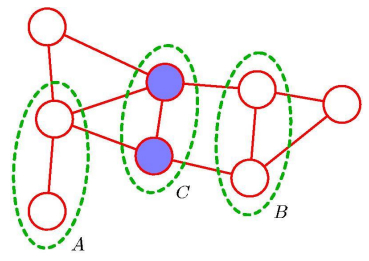
Course Outline

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 - > Exact Inference



Recap: Undirected Graphical Models

- Undirected graphical models ("Markov Random Fields")
 - Given by undirected graph



- Conditional independence for undirected graphs
 - > If every path from any node in set A to set B passes through at least one node in set C, then $A \perp B | C$.
 - Simple Markov blanket:



Recap: Factorization in MRFs

- Joint distribution
 - Written as product of potential functions over maximal cliques in the graph:

$$p(\mathbf{x}) = \frac{1}{Z} \prod_{C} \psi_C(\mathbf{x}_C)$$

> The normalization constant Z is called the partition function.

$$Z = \sum_{\mathbf{x}} \prod_{C} \psi_C(\mathbf{x}_C)$$

• Remarks

- BNs are automatically normalized. But for MRFs, we have to explicitly perform the normalization.
- Presence of normalization constant is major limitation!
 - Evaluation of Z involves summing over $\mathcal{O}(K^M)$ terms for M nodes!



Factorization in MRFs

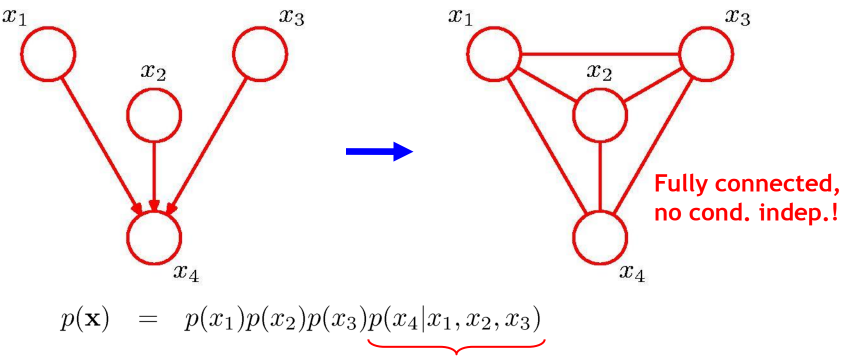
- Role of the potential functions
 - General interpretation
 - No restriction to potential functions that have a specific probabilistic interpretation as marginals or conditional distributions.
 - Convenient to express them as exponential functions ("Boltzmann distribution")

$$\psi_C(\mathbf{x}_C) = \exp\{-E(\mathbf{x}_C)\}\$$

- with an energy function E.
- Why is this convenient?
 - Joint distribution is the product of potentials \Rightarrow sum of energies.
 - We can take the log and simply work with the sums...

UNIVERSIT Recap: Converting Directed to Undirected Graphs

• Problematic case: multiple parents



Need a clique of $x_{\scriptscriptstyle 1}\text{,...,}x_{\scriptscriptstyle 4}$ to represent this factor!

Need to introduce additional links ("marry the parents").
 ⇒ This process is called moralization. It results in the moral graph.

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Recap: Conversion Algorithm

- General procedure to convert directed \rightarrow undirected
 - 1. Add undirected links to marry the parents of each node.
 - 2. Drop the arrows on the original links \Rightarrow moral graph.
 - 3. Find maximal cliques for each node and initialize all clique potentials to 1.
 - 4. Take each conditional distribution factor of the original directed graph and multiply it into one clique potential.

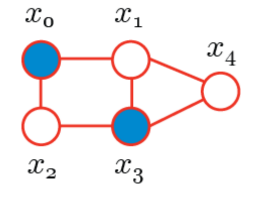
Restriction

- Conditional independence properties are often lost!
- > Moralization results in additional connections and larger cliques.



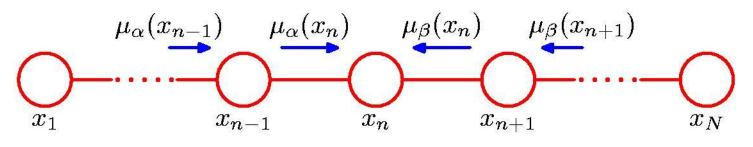
Recap: Computing Marginals

- How do we apply graphical models?
 - Given some observed variables, we want to compute distributions of the unobserved variables.
 - > In particular, we want to compute marginal distributions, for example $p(x_4)$.



- How can we compute marginals?
 - Classical technique: sum-product algorithm by Judea Pearl.
 - In the context of (loopy) undirected models, this is also called (loopy) belief propagation [Weiss, 1997].
 - Basic idea: message-passing.

Recap: Message Passing on a Chain



- > Idea
 - Pass messages from the two ends towards the query node x_n .
- > Define the messages recursively:

$$\mu_{\alpha}(x_n) = \sum_{x_{n-1}} \psi_{n-1,n}(x_{n-1}, x_n) \mu_{\alpha}(x_{n-1})$$
$$\mu_{\beta}(x_n) = \sum_{x_{n+1}} \psi_{n,n+1}(x_n, x_{n+1}) \mu_{\beta}(x_{n+1})$$

 \succ Compute the normalization constant Z at any node x_m .

$$Z = \sum_{x_n} \mu_{\alpha}(x_n) \mu_{\beta}(x_n)$$

Slide adapted from Chris Bishop

B. Leibe

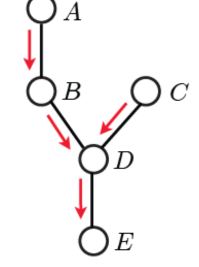
103 Image source: C. Bishop, 2006

Recap: Message Passing on Trees

- General procedure for all tree graphs.
 - Root the tree at the variable that we want to compute the marginal of.
 - Start computing messages at the leaves.
 - Compute the messages for all nodes for which all incoming messages have already been computed.
 - Repeat until we reach the root.
- If we want to compute the marginals for all possible nodes (roots), we can reuse some of the messages.
 - Computational expense linear in the number of nodes.
- We already motivated message passing for inference.
 - How can we formalize this into a general algorithm?

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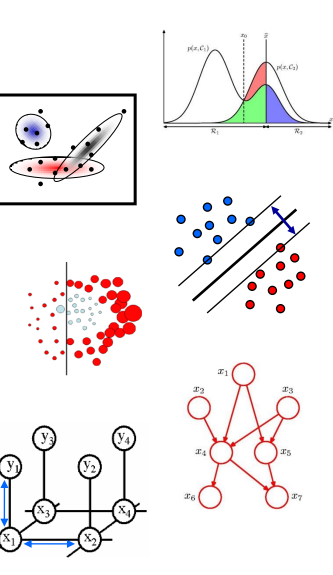
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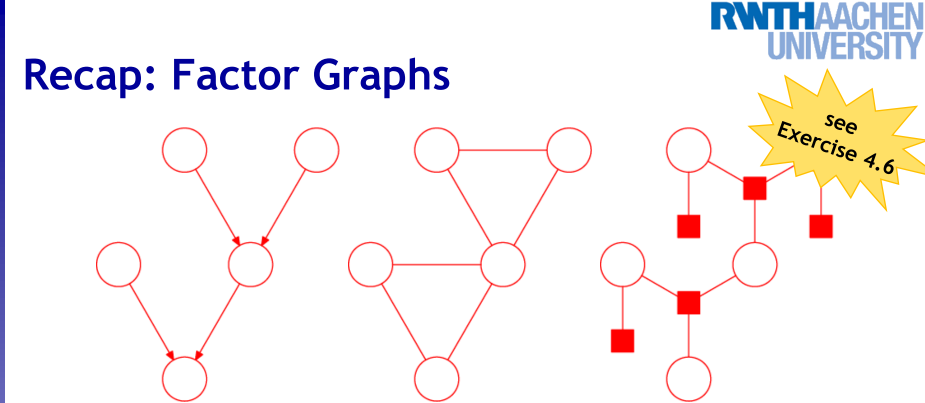


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Course Outline

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- Joint probability
 - > Can be expressed as product of factors: $p(\mathbf{x}) = \frac{1}{Z} \prod f_s(\mathbf{x}_s)$
 - > Factor graphs make this explicit through separate factor nodes.
- Converting a directed polytree
 - Conversion to undirected tree creates loops due to moralization!
 - > Conversion to a factor graph again results in a tree!



Recap: Sum-Product Algorithm

- Objectives
 - > Efficient, exact inference algorithm for finding marginals.

• Procedure:

- > Pick an arbitrary node as root.
- Compute and propagate messages from the leaf nodes to the root, storing received messages at every node.
- Compute and propagate messages from the root to the leaf nodes, storing received messages at every node.
- Compute the product of received messages at each node for which the marginal is required, and normalize if necessary.

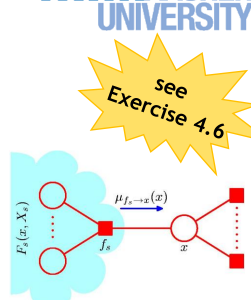
$$p(x) \propto \prod_{s \in \operatorname{ne}(x)} \mu_{f_s \to x}(x)$$

- Computational effort
 - > Total number of messages = $2 \cdot \text{number of graph edges.}$

Recap: Sum-Product Algorithm

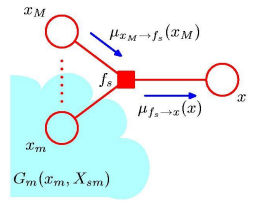
- Two kinds of messages
 - Message from factor node to variable nodes:
 - Sum of factor contributions

$$\mu_{f_s \to x}(x) \equiv \sum_{X_s} F_s(x, X_s)$$
$$= \sum_{X_s} f_s(\mathbf{x}_s) \prod_{m \in \operatorname{ne}(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$$

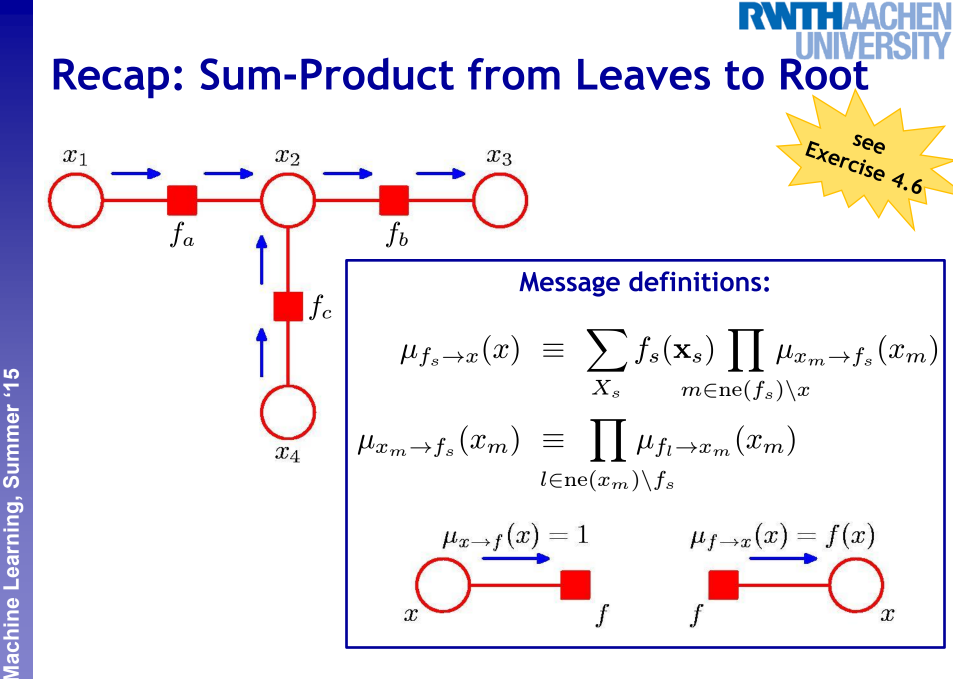


- Message from variable node to factor node:
 - Product of incoming messages

$$\mu_{x_m \to f_s}(x_m) \equiv \prod_{l \in \operatorname{ne}(x_m) \setminus f_s} \mu_{f_l \to x_m}(x_m)$$

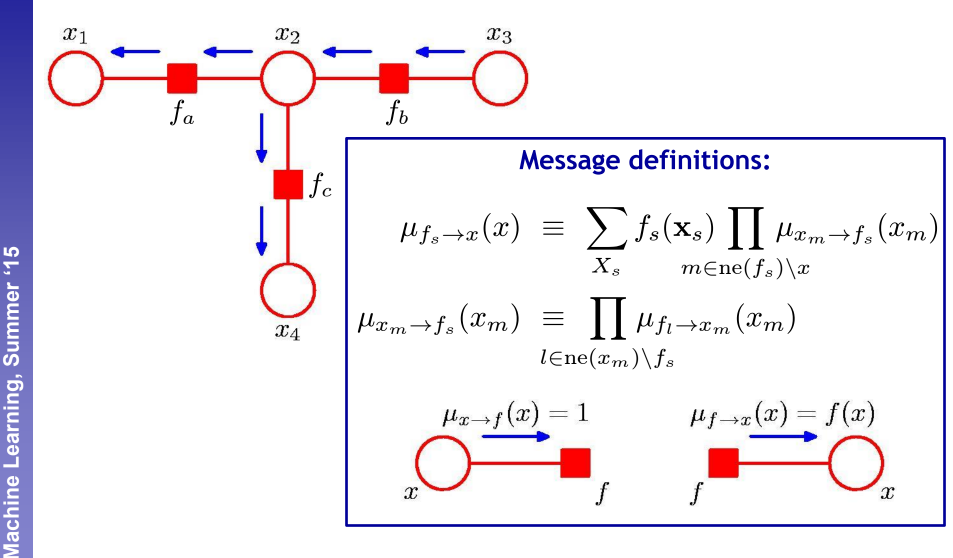


 \Rightarrow Simple propagation scheme.



109 Image source: C. Bishop, 2006

RWTHAACHEN UNIVERSITY Recap: Sum-Product from Root to Leaves



110 Image source: C. Bishop, 2006



Recap: Max-Sum Algorithm

- Objective: an efficient algorithm for finding
 - \succ Value $\mathbf{x}^{ ext{max}}$ that maximises $p(\mathbf{x})$;
 - > Value of $p(\mathbf{x}^{\max})$.

 \Rightarrow Application of dynamic programming in graphical models.

• Key ideas

> We are interested in the maximum value of the joint distribution

$$p(\mathbf{x}^{\max}) = \max_{\mathbf{x}} p(\mathbf{x})$$

- \Rightarrow Maximize the product $p(\mathbf{x})$.
- For numerical reasons, use the logarithm.

$$\ln\left(\max_{\mathbf{x}} p(\mathbf{x})\right) = \max_{\mathbf{x}} \ln p(\mathbf{x}).$$

 \Rightarrow Maximize the sum (of log-probabilities).



Recap: Max-Sum Algorithm

Initialization (leaf nodes) \bullet

$$\mu_{x \to f}(x) = 0 \qquad \qquad \mu_{f \to x}(x) = \ln f(x)$$

Г

- Recursion
 - Messages

$$\begin{array}{ll} \text{Messages} \\ \mu_{f \to x}(x) &= \max_{x_1, \dots, x_M} \left[\ln f(x, x_1, \dots, x_M) + \sum_{m \in \operatorname{ne}(f_s) \setminus x} \mu_{x_m \to f}(x_m) \right] \\ \\ \mu_{x \to f}(x) &= \sum_{l \in \operatorname{ne}(x) \setminus f} \mu_{f_l \to x}(x) \end{array}$$

> For each node, keep a record of which values of the variables gave rise to the maximum state:

$$\phi(x) = \arg \max_{x_1, \dots, x_M} \left[\ln f(x, x_1, \dots, x_M) + \sum_{m \in \operatorname{ne}(f_s) \setminus x} \mu_{x_m \to f}(x_m) \right]$$

Slide adapted from Chris Bishop

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Recap: Max-Sum Algorithm

- Termination (root node)
 - Score of maximal configuration

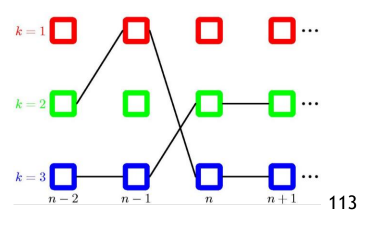
$$p^{\max} = \max_{x} \left[\sum_{s \in \operatorname{ne}(x)} \mu_{f_s \to x}(x) \right]$$

> Value of root node variable giving rise to that maximum

$$x^{\max} = \arg \max_{x} \left[\sum_{s \in \operatorname{ne}(x)} \mu_{f_s \to x}(x) \right]$$

 Back-track to get the remaining variable values

$$x_{n-1}^{\max} = \phi(x_n^{\max})$$



Recap: Junction Tree Algorithm

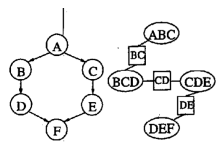
- **Motivation**
 - **Exact** inference on general graphs. \geq
 - Works by turning the initial graph into a junction tree and then running a sum-product-like algorithm.
 - Intractable on graphs with large cliques.

Main steps

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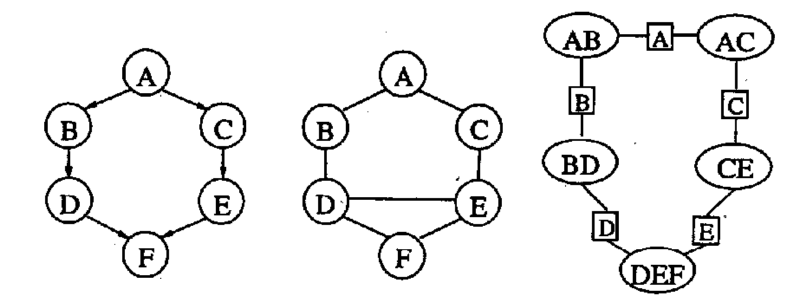
- 1. If starting from directed graph, first convert it to an undirected graph by moralization.
- 2. Introduce additional links by triangulation in order to reduce the size of cycles.
- 3. Find cliques of the moralized, triangulated graph.
- 4. Construct a new graph from the maximal cliques.
- 5. Remove minimal links to break cycles and get a junction tree.
- \Rightarrow Apply regular message passing to perform inference.



See Exercise 5.



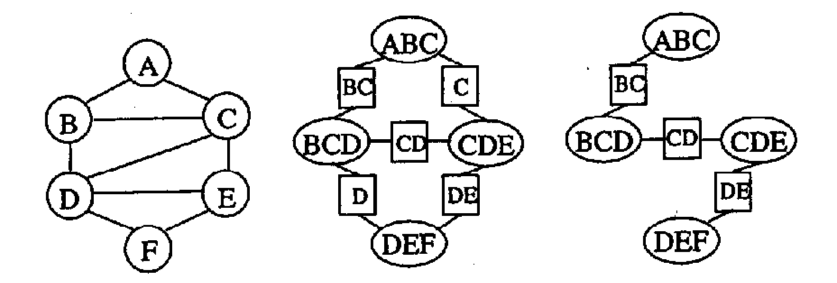
Recap: Junction Tree Example



- Without triangulation step
 - The final graph will contain cycles that we cannot break without losing the running intersection property!



Recap: Junction Tree Example



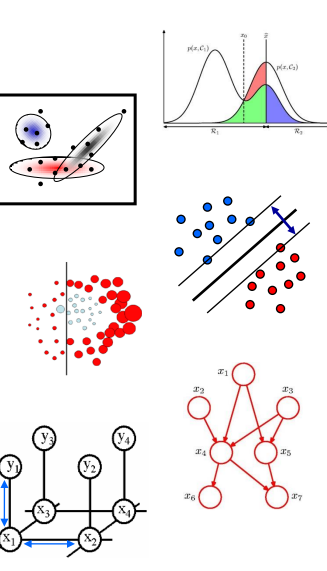
When applying the triangulation

- > Only small cycles remain that are easy to break.
- Running intersection property is maintained.

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Course Outline

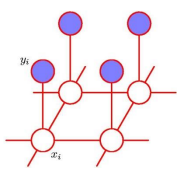
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Recap: MRF Structure for Images

• Basic structure



Noisy observations

"True" image content

Two components

- > Observation model
 - How likely is it that node x_i has label L_i given observation y_i ?
 - This relationship is usually learned from training data.
- Neighborhood relations
 - Simplest case: 4-neighborhood
 - Serve as smoothing terms.
 - \Rightarrow Discourage neighboring pixels to have different labels.
 - This can either be learned or be set to fixed "penalties".

Recap: How to Set the Potentials?

- Unary potentials
 - > E.g. color model, modeled with a Mixture of Gaussians

$$\phi(x_i, y_i; \theta_{\phi}) = \log \sum_k \theta_{\phi}(x_i, k) p(k|x_i) \mathcal{N}(y_i; \bar{y}_k, \Sigma_k)$$

 \Rightarrow Learn color distributions for each label

$$\phi(x_p = 1, y_p) \phi(x_p = 0, y$$

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Recap: How to Set the Potentials?

- Pairwise potentials
 - Potts Model

$$\psi(x_i, x_j; \theta_{\psi}) = \theta_{\psi} \delta(x_i \neq x_j)$$

- Simplest discontinuity preserving model.
- Discontinuities between any pair of labels are penalized equally.
- Useful when labels are unordered or number of labels is small.
- Extension: "contrast sensitive Potts model"

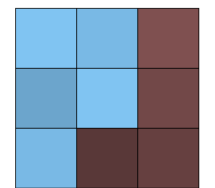
$$\psi(x_i, x_j, g_{ij}(y); \theta_{\psi}) = \theta_{\psi} g_{ij}(y) \delta(x_i \neq x_j)$$

where

$$g_{ij}(y) = e^{-\beta \|y_i - y_j\|^2}$$
 $\beta = 2 / avg(\|y_i - y_j\|^2)$

- Discourages label changes except in places where there is also a large change in the observations.

Recap: Graph Cuts for Binary Problems



"expected" intensities of object and background I^s and I^t can be re-estimated $D_p(t)$ $T_p(t)$ $T_p(t)$ $T_p($

$$D_p(s) \propto \exp\left(-\|I_p - I^s\|^2 / 2\sigma^2\right)$$
$$D_p(t) \propto \exp\left(-\|I_p - I^t\|^2 / 2\sigma^2\right)$$

EM-style optimization

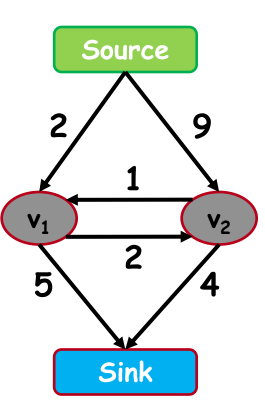
121 [Boykov & Jolly, ICCV'01]

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Recap: s-t-Mincut Equivalent to Maxflow

Flow = 0



Augmenting Path Based Algorithms

- 1. Find path from source to sink with positive capacity
- 2. Push maximum possible flow through this path
- 3. Repeat until no path can be found

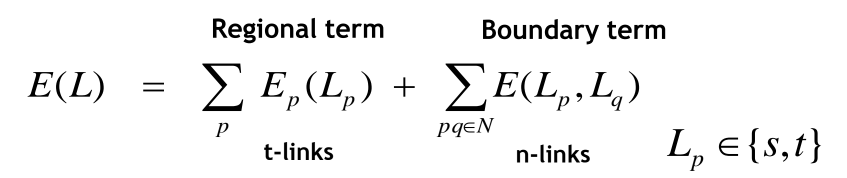
Algorithms assume non-negative capacity

Slide credit: Pushmeet Kohli

B. Leibe

See Exercise 5.2

RWTHAACHEN UNIVERSITY Recap: When Can s-t Graph Cuts Be Applied?



• s-t graph cuts can only globally minimize binary energies that are submodular. [Boros & Hummer, 2002, Kolmogorov & Zabih, 2004]

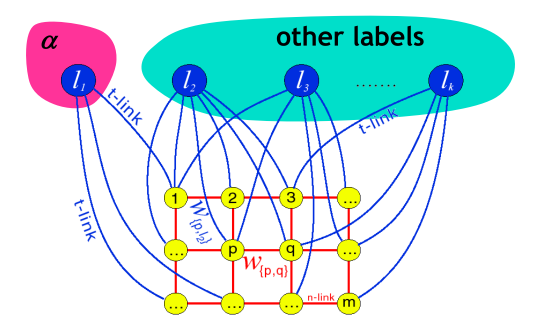
$$E(L)$$
 can be minimized
by s-t graph cuts $\longleftrightarrow E(s,s) + E(t,t) \le E(s,t) + E(t,s)$ Submodularity ("convexity")

- Submodularity is the discrete equivalent to convexity.
 - Implies that every local energy minimum is a global minimum.
 - \Rightarrow Solution will be globally optimal.



Recap: α-Expansion Move

- Basic idea:
 - Break multi-way cut computation into a sequence of binary s-t cuts.



No longer globally optimal result, but guaranteed approximation quality and typically converges in few iterations.

Recap: Converting an MRF to an s-t Graph

Graph *g;

For all pixels p

/* Add a node to the graph */
nodeID(p) = g->add_node();

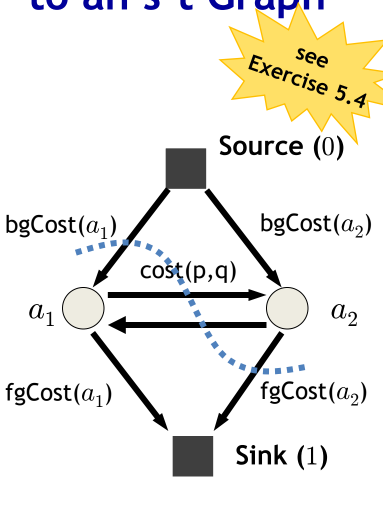
/* Set cost of terminal edges */
set_weights(nodeID(p), fgCost(p), bgCost(p));

end

```
g->compute_maxflow();
```

```
label_p = g->is_connected_to_source(nodeID(p));
```

```
// is the label of pixel p (0 or 1)
```



$$a_1 = bg a_2 = fg$$

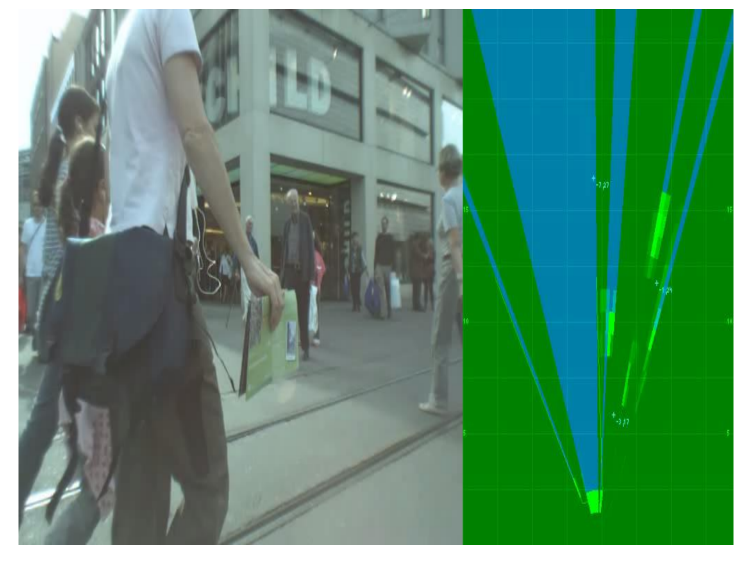


Any Questions?

So what can you do with all of this?

Mobile Object Detection & Tracking







[Ess, Leibe, Schindler, Van Gool, CVPR'08]

Learning Person-Object Interactions



128 B. Leibe [T. Baumgartner, D. Mitzel, B. Leibe, CVPR'13]

ΓΗΔΔ(;)

NTHAACH

RF (HOG)

Semantic Segmentation

image







ground truth





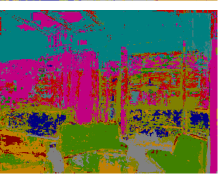
Baseline







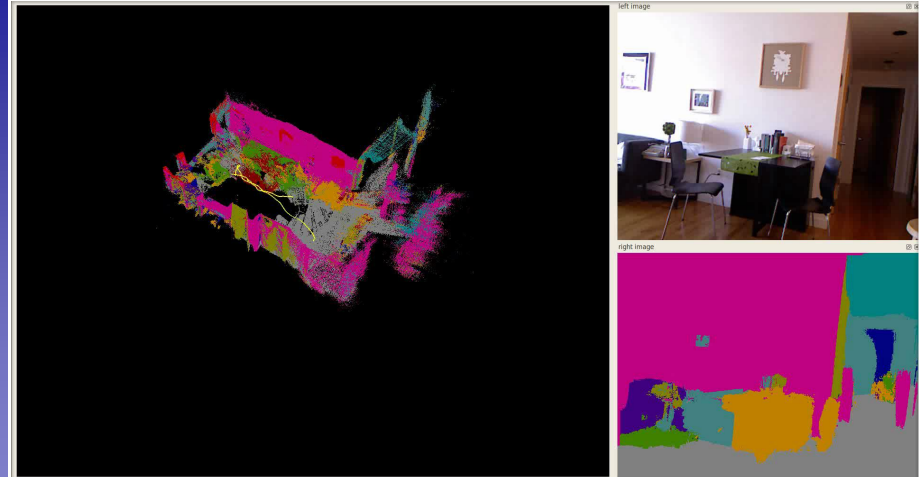






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3D Labeling Results - Living Room

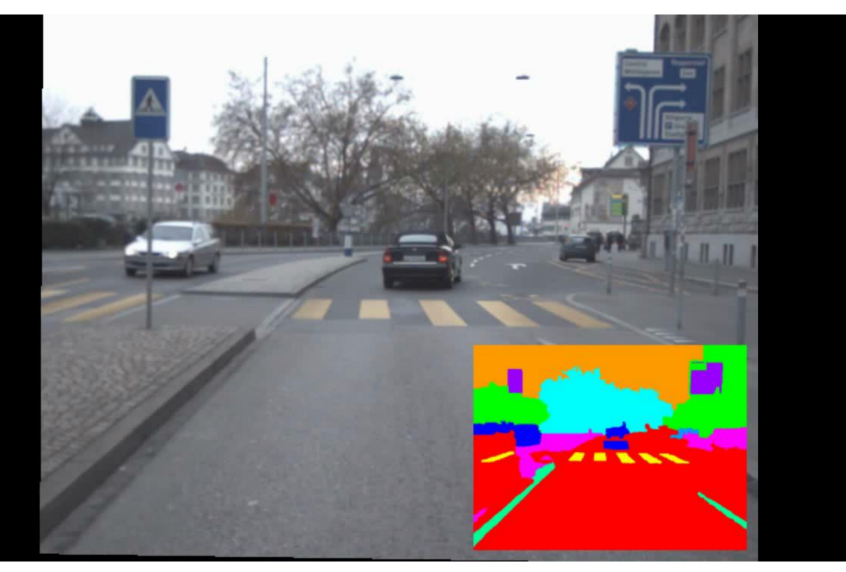


<u>play video</u>

130 [Hermans, Floros, Leibe, submission to ICCV'13]



Semantic Scene Segmentation





Any More Questions?

Good luck for the exam!