Basics on generative and discriminative classification

Machine Learning and Object Recognition 2016-2017 Jakob Verbeek

Course website:

http://thoth.inrialpes.fr/~verbeek/MLOR.16.17



Practical matters

- Online course information
 - Updated schedule, links to slides and papers
 - http://thoth.inrialpes.fr/~verbeek/MLOR.16.17.php
- Grading: Final grades are determined as follows
 - 50% written exam, 50% quizes on the presented papers
 - If you present a paper: the grade for the presentation can substitute the worst grade you had for any of the quizes.
- Paper presentations:
 - each student presents once
 - each paper is presented by two or three students
- presentations last for 15~20 minutes, time yours in advance!

Classification in its simplest form

• Given training data labeled for two or more classes



Classification in its simplest form

- Given training data labeled for two or more classes
- Determine a surface that separates those classes



Classification in its simplest form

- Given training data labeled for two or more classes
- Determine a surface that separates those classes
- Use that surface to predict the class membership of new data



Classification examples in category-level recognition

- Image classification: for each of a set of labels, predict if it is relevant or not for a given image.
- For example: Person = yes, TV = yes, car = no, ...





Classification examples in category-level recognition

- Category localization: predict bounding box coordinates.
- Classify each possible bounding box as containing the category or not.
- Report most confidently classified box.



natics *mathematics*

Classification examples in category-level recognition

- Semantic segmentation: classify pixels to categories (multi-class)
- Impose spatial smoothness by Markov random field models.





Classification

- Goal is to predict for a test data input the corresponding class label.
 - Data input x, e.g. image but could be anything, format may be vector or other
 - Class label y, can take one out of at least 2 discrete values, can be more
 - In binary classification we often refer to one class as "positive", and the other as "negative"
- Classifier: function f(x) that assigns a class to x, or probabilities over the classes.
- Training data: pairs (x,y) of inputs x, and corresponding class label y.
- Learning a classifier: determine function f(x) from some family of functions based on the available training data.
- Classifier partitions the input space into regions where data is assigned to a given class
 - Specific form of these boundaries will depend on the family of classifiers used



Generative classification: principle

- Model the class conditional distribution over data x for each class y: p(x|y)
 - Data of the class can be sampled (generated) from this distribution
- Estimate the a-priori probability that a class will appear p(y)
- Infer the probability over classes using Bayes' rule of conditional probability

$$p(y|x) = \frac{p(y) p(x|y)}{p(x)}$$

• Marginal distribution on x is obtained by marginalizing the class label y $p(x) = \sum_{y} p(y) p(x|y)$



Generative classification: practice

- In order to apply Bayes' rule, we need to estimate two distributions.
- A-priori class distribution
 - In some cases the class prior probabilities are known in advance.
 - If the frequencies in the training data set are representative for the true class probabilities, then estimate the prior by these frequencies.
- Class conditional data distributions
 - Select a class of density models
 - Parametric model, e.g. Gaussian, Bernoulli, ...
 - Semi-parametric models: mixtures of Gaussian, Bernoulli, ...
 - Non-parametric models: histograms, nearest-neighbor method, ...
 - Or more structured models taking problem knowledge into account.
 - Estimate the parameters of the model using the data in the training set associated with that class.



Estimation of the class conditional model

- Given a set of n samples from a certain class, and a family of distributions $X = \{x_1, ..., x_n\}$ $P = \{p_{\theta}(x); \theta \in \Theta\}$
- How do we quantify the fit of a certain model to the data, and how do we find the best model defined in this sense?
- Maximum a-posteriori (MAP) estimation: use Bayes' rule again as follows:
 - Assume a prior distribution over the parameters of the model $p(\theta)$
 - Then the posterior likelihood of the model given the data is $p(\theta|X) = p(X|\theta)p(\theta)/p(X)$
 - Find the most likely model given the observed data $\hat{\theta} = \operatorname{argmax}_{\theta} p(\theta|X) = \operatorname{argmax}_{\theta} \{\ln p(\theta) + \ln p(X|\theta)\}$
- Maximum likelihood parameter estimation: assume prior over parameters is uniform (for bounded parameter spaces), or "near uniform" so that its effect is negligible for the posterior on the parameters.
 - In this case the MAP estimator is given by $\hat{\theta} = \operatorname{argmax}_{\theta} p(X|\theta)$
 - For i.id. samples:

$$\hat{\theta} = \operatorname{argmax}_{\theta} \prod_{i=1}^{n} p(x_i | \theta) = \operatorname{argmax}_{\theta} \sum_{i=1}^{n} \ln p(x_i | \theta)$$

Generative classification methods

- Generative probabilistic methods use Bayes' rule for prediction
 - Problem is reformulated as one of parameter/density estimation $p(y|x) = \frac{p(y) p(x|y)}{p(x)}$ $p(x) = \sum_{y} p(y) p(x|y)$
- Adding new classes to the model is easy:
 - Existing class conditional models stay as they are
 - Estimate p(x|new class) from training examples of new class
 - Re-estimate class prior probabilities



Example of generative classification

- Three-class example in 2D with parametric model
 - Single Gaussian model per class, uniform class prior
 - Exercise 1: how is this model related to the Gaussian mixture model we looked at before for clustering ?
 - Exercise 2: characterize surface of equal class probability when the covariance matrices are the same for all classes





Density estimation for class-conditional models

- Any type of data distribution may be used, preferably one that is modeling the data well, so that we can hope for accurate classification results.
- If we do not have a clear understanding of the data generating process, we can use a generic approach,
 - Gaussian distribution, or other reasonable parametric model
 - Estimation often in closed form or relatively simple process
 - Mixtures of parametric models
 - Estimation using EM algorithm, not more complicated than single parametric model
 - Non-parametric models can adapt to any data distribution given enough data for estimation. Examples: (multi-dimensional) histograms, and nearest neighbors.
 - Estimation often trivial, given a single smoothing parameter.

Histogram density estimation

- Suppose we have *N* data points use a histogram with *C* cells
- Consider maximum likelihood estimator

$$\hat{\theta} = \operatorname{argmax}_{\theta} \sum_{i=1}^{n} \ln p_{\theta}(x_i) = \operatorname{argmax}_{\theta} \sum_{c=1}^{C} n_c \ln \theta_c$$

- Take into account constraint that density should integrate to one $\theta_C := 1 - \left(\sum_{k=1}^{C-1} v_k \theta_k \right) / v_C$
- Exercise: derive maximum likelihood estimator
- Some observations:
 - Discontinuous density estimate
 - Cell size determines smoothness
 - Number of cells scales exponentially with the dimension of the data





Histogram density estimation

- Suppose we have *N* data points use a histogram with *C* cells
- Data log-likelihood

$$L(\theta) = \sum_{i=1}^{N} \ln p_{\theta}(x_i) = \sum_{c=1}^{C} n_c \ln \theta_c$$

- Take into account constraint that density should integrate to one $\theta_C := 1 - \left(\sum_{k=1}^{C-1} v_k \theta_k \right) / v_C$
- Compute derivative, and set to zero for i=1,..., C-1

$$\frac{\partial L(\theta)}{\partial \theta_i} = \frac{n_i}{\theta_i} - \frac{n_c}{\theta_c} \frac{v_i}{v_c}$$
$$\theta_i v_i = \frac{\theta_c v_c}{n_c} n_i$$

• Use fact that probability mass should integrate to one, and substitute

$$\sum_{i=1}^{C} \theta_{i} v_{i} = \frac{\theta_{C} v_{C}}{n_{C}} \sum_{i=1}^{C} n_{i} = \frac{\theta_{C} v_{C}}{n_{C}} N = 1$$
$$\theta_{i} = \frac{n_{i}}{v_{i} N}$$



The Naive Bayes model

- Histogram estimation, and other methods, scale poorly with data dimension
 - Fine division of each dimension: many empty bins
 - Rough division of each dimension: poor density model
 - Even for one cut per dimension: 2^D cells, eg. a million cells in 20 dims.
- The number of parameters can be made linear in the data dimension by assuming independence between the dimensions

$$p(x) = \prod_{d=1}^{D} p(x(d))$$

- For example, for histogram model: we estimate a histogram per dimension
 - Still C^D cells, but only D x C parameters to estimate, instead of C^D
- Independence assumption can be unrealistic for high dimensional data
 - But classification performance may still be good using the derived p(y|x)
 - Partial independence, e.g. using graphical models, relaxes this problem.
- Principle can be applied to estimation with any type of density estimate

Example of a naïve Bayes model

- Hand-written digit classification
 - Input: binary 28x28 scanned digit images



- Desired output: class label of image
- Generative model over 28 x 28 pixel images: 2⁷⁸⁴ possible images
 - Independent Bernoulli model for each class
 - Probability per pixel per class
 - Maximum likelihood estimator is average value per pixel/bit per class

$$p(x|y=c) = \prod_{d} p(x^{d}|y=c)$$

$$p(x^{d}=1|y=c) = \theta_{cd}$$

$$p(x^{d}=0|y=c) = 1 - \theta_{cd}$$



• Classify using Bayes' rule: $p(y|x) = \frac{p(y) p(x|y)}{p(x)}$



k-nearest-neighbor density estimation: principle

- Instead of having fixed cells as in histogram method,
 - Center cell on the test sample for which we evaluate the density.
 - Fix number of samples in the cell, find the corresponding **cell size**.
- Probability to find a point in a sphere **A** centered on x_0 with volume **v** is

$$P(x \in A) = \int_{A} p(x) dx$$

- A smooth density is approximately constant in small region, and thus $P(x \in A) = \int_{A} p(x) dx \approx \int_{A} p(x_0) dx = p(x_0) v_A$
- Alternatively: estimate **P** from the fraction of training data in **A**: $P(x \in A) \approx \frac{K}{N}$

 $p(x_0) \approx \frac{\kappa}{Nv}$

- Total N data points, k in the sphere A
- Combine the above to obtain estimate
 - Same per-cell density estimate as in histogram estimator
- Note: density estimates not guaranteed to integrate to one!



k-nearest-neighbor density estimation: practice

- Procedure in practice:
 - Choose k
 - For given \boldsymbol{x} , compute the volume \boldsymbol{v} which contain \boldsymbol{k} samples.
 - Estimate density with $p(x) \approx \frac{k}{Nv}$
- Volume of a sphere with radius *r* in *d* dimensions is

$$v(r,d) = \frac{2r^d \pi^{d/2}}{\Gamma(d/2+1)}$$

- What effect does k have?
 - Data sampled from mixture of Gaussians plotted in green
 - Larger k, larger region, smoother estimate
 - Similar role as cell size for histogram estimation



K-nearest-neighbors for classification

- Use Bayes' rule with kNN density estimation for p(x|y)
 - Find sphere volume v to capture **k** data points for estimate $p(x) = \frac{k}{Nv}$
 - Use the same sphere for each class for estimates $p(x|y=c) = \frac{k_c}{N_v}$
 - Estimate class prior probabilities $p(y=c) = \frac{N_c}{N}$
 - Calculate class posterior distribution as fraction of k neighbors in class c



Smoothing effects for large values of k: data set











Summary generative classification methods

- (Semi-) Parametric models, e.g. p(x|y) is Gaussian, or mixture of ...
 - Pros: no need to store training data, just the class conditional models
 - Cons: may fit the data poorly, and might therefore lead to poor classification result
- Non-parametric models:
 - Pros:
 - flexibility, no assumptions distribution shape, learning is trivial
 - KNN can be used for anything that comes with a distance.
 - Cons of histograms:
 - Only practical in low dimensional data (<5 or so), application in high dimensional data leads to exponentially many and mostly empty cells
 - Naïve Bayes modeling in higher dimensional cases
 - Cons of k-nearest neighbors
 - Need to store all training data (memory cost)
 - Computing nearest neighbors (computational cost)



Discriminative classification methods

- Generative classification models
 - Model the density of inputs x from each class p(x|y)
 - Estimate class prior probability p(y)
 - Use Bayes' rule to infer distribution over class given input
- In discriminative classification methods we directly estimate class probability given input: p(y|x)
 - Choose class of decision functions in feature space
 - Estimate function that maximizes performance on the training set
 - Classify a new pattern on the basis of this decision rule.



Binary linear classifier

• Decision function is linear in the features:

$$f(x) = w^T x + b = b + \sum_{i=1}^d w_i x_i$$

- Classification based on the sign of f(x)
- Orientation is determined by **w**
- Offset from origin is determined by *b*
- Decision surface is (d-1) dimensional hyper-plane orthogonal to w, given by

 $f(x) = w^T x + b = 0$





Common loss functions for classification

- Assign class label using y = sign(f(x))
- Measure how model quality on a test sample using loss function
 - ► Zero-One loss: $L(y_i, f(x_i)) = [y_i f(x_i) \le 0]$
 - Hinge loss: $L(y_i, f(x_i)) = max(0, 1 y_i f(x_i))$ Logistic loss: $L(y_i, f(x_i)) = \log_2(1 + e^{-y_i f(x_i)})$







Common loss functions for classification

- Assign class label using y = sign(f(x))
 - ► Zero-One loss: $L(y_i, f(x_i)) = [y_i f(x_i) \le 0]$
 - Hinge loss: L(y)
 - Logistic loss:

$$L(y_{i}, f(x_{i})) = max (0, 1 - y_{i}f(x_{i}))$$

$$L(y_{i}, f(x_{i})) = \log_{2}(1 + e^{-y_{i}f(x_{i})})$$

- The zero-one loss counts the number of misclassifications, which is the "ideal" empirical loss
 - Discontinuity at zero makes optimization intractable
- Hinge and logistic loss provide continuous and convex upperbounds, which allow for continuous optimization



Dealing with more than two classes

- First idea: construction from multiple binary classifiers
 - Learn binary "base" classifiers independently
- One vs rest approach: 1 vs (2 & 3) 2 vs (1 & 3) 7 3 vs (1 & 2) ► \mathcal{R}_1 Problem: Region claimed by several classes \mathcal{R}_2 \mathcal{C}_1 , \mathcal{R}_3 \mathcal{C}_2 • not \mathcal{C}_1 not \mathcal{C}_2



Dealing with more than two classes

- First idea: construction from multiple binary classifiers
 - Learn binary "base" classifiers independently
- One vs one approach:
 - 1 vs 2
 - 1 vs 3
 - 2 vs 3
- Problem: conflicts in some regions





Dealing with more than two classes

• Instead: define a separate linear score function for each class

 $f_k(x) = w_k^T x + b_k$

• Assign sample to the class of the function with maximum value

$$y = arg max_k f_k(x)$$

• Exercise 1: give the expression for points where two classes have equal score

- Exercise 2: show that the set of points assigned to a class is convex
 - If two points fall in the region, then also all points on connecting line



Logistic discriminant for two classes

- Map linear score function to class probabilities with sigmoid function $p(y=+1|x)=\sigma(w^Tx+b)$
 - For binary classification problem, we have by definition

$$p(y=-1|x)=1-p(y=+1|x)$$

Exercise: show that

$$p(y=-1|x)=\sigma(-(w^{T}x+b))$$

and thus





Logistic discriminant for two classes

- Map linear score function to class probabilities with sigmoid function
- The class boundary is obtained for p(y|x)=1/2, thus by setting linear function in exponent to zero







Multi-class logistic discriminant

- Map score function of each class to class probabilities with "soft-max" function
 - Absorb bias into w and x

$$p(y=c|x) = \frac{\exp(f_c(x))}{\sum_{k=1}^{K} \exp(f_k(x))}$$

- The class probability estimates are non-negative, and sum to one.
- Relative probability of most likely class increases exponentially with the difference in the linear score functions

$$\frac{p(y=c|x)}{p(y=k|x)} = \frac{\exp(f_c(x))}{\exp(f_k(x))} = \exp(f_c(x) - f_k(x))$$

• For any given pair of classes we find that they are equally likely on a hyperplane in the feature space





(a ())

Maximum likelihood parameter estimation

- Maximize the log-likelihood of predicting the correct class label for training data
 - Predictions are made independently, so sum log-likelihood of all training data $L = \sum_{n=1}^{N} \log p(y_n | x_n)$
- Derivative of log-likelihood as intuitive interpretation



- No closed-form solution, but log-likelihood is concave in parameters
 - no local optima, use general purpose convex optimization methods
 - For example: gradient started from w=0
 - w is linear combination of data points
 - Sign of coefficients depends on class labels

Maximum a-posteriori (MAP) parameter estimation

- Let us assume a zero-mean Gaussian prior distribution on w
 - We expect weight vectors with a small norm
- Find w that maximizes posterior likelihood $\hat{w} = \operatorname{argmax}_{w} \sum_{n=1}^{N} \ln p(y_{n}|x_{n}, w) + \ln p(w)$
- Can be rewritten as following "penalized" maximum likelihood estimator: $\hat{w} = \operatorname{argmax}_{w} \sum_{n=1}^{N} \ln p(y_n | x_n, w) - \lambda ||w||_2^2$
 - With lambda non-negative
- Penalty for "large" w, bounds the scale of w in case of separable data
- Exercise: show that for separable data the norm of the optimal w's would be infinite without using the penalty term.



Support Vector Machines

- Find linear function to separate positive and negative examples
- Which function best separates the samples ?
 - Function inducing the largest margin



Support vector machines

- Without loss of generality, let function value at the margin be +/- 1
- Now constrain w to that all points fall on correct side of the margin:

 $y_i(w^T x_i + b) \ge 1$

• By construction we have that the "support vectors", the ones that define the margin, have function values

$$w^T x_i + b = y_i$$

• Express the size of the margin in terms of w.





Support vector machines

Let's consider a support vector x from the positive class $f(x) = w^T x + b = 1$ Let z be its projection on the decision plane Since w is normal vector to the decision plane, we have $z = x - \alpha w$ and since z is on the decision plane $f(z) = w^T(x - \alpha w) + b = 0$ Solve for alpha $w^{T}(x-\alpha w)+b=0$ $w^T x + b - \alpha w^T w = 0$ $\alpha w^T w = 1$ $\alpha = \frac{1}{\|\boldsymbol{w}\|_2^2}$ Margin is twice distance from x to z $||x-z||_2 = ||x-(x-\alpha w)||_2$ $\|\alpha w\|_2 = \alpha \|w\|_2$ $\frac{\|w\|_2}{\|w\|_2^2} = \frac{1}{\|w\|_2}$ Support vectors Margin

Support vector machines

- To find the maximum-margin separating hyperplane, we
 - Maximize the margin, while ensuring correct classification
 - Minimize the norm of w, s.t. $\forall_i : y_i(w^T x_i + b) \ge 1$
- Solve using quadratic program with linear inequality constraints over p+1 variables

$$argmin_{w,b} \frac{1}{2} w^{T} w$$

subject to $y_{i}(w^{T} x_{i} + b) \ge 1$





Support vector machines: inseperable classes

- For non-separable classes we incorporate hinge-loss $L(y_i, f(x_i)) = max(0, 1 - y_i f(x_i))$
- Recall: convex and piece-wise linear upper bound on zero/one loss.
 - Zero if point on the correct side of the margin
 - Otherwise given by absolute difference from score at margin



Support vector machines: inseperable classes

Minimize penalized loss function

$$min_{w,b} \quad \lambda \frac{1}{2} w^T w + \sum_i max(0, 1 - y_i(w^T x_i + b))$$

Quadratic function, plus piece-wise linear functions.

- Transformation into a quadratic program
 - Define "slack variables" that measure the loss for each data point
 - Should be non-negative, and at least as large as the loss

$$\min_{w,b,\{\xi_i\}} \quad \lambda \frac{1}{2} w^T w + \sum_i \xi_i$$

subject to $\forall_i: \xi_i \ge 0$ and $\xi_i \ge 1 - y_i (w^T x_i + b)$

 Solution of the quadratic program has the property that w is a linear combination of the data points.



SVM solution properties

• Optimal w is a linear combination of data points

$$w = \sum_{n=1}^{N} \alpha_n y_n x_n$$

- Alpha weights are zero for all points on the correct side of the margin
- Points on the margin, or on the wrong side, have non-zero weight
 - Called support vectors
- Classification function thus has form $f(x) = w^T x + b = \sum_{n=1}^N \alpha_n y_n x_n^T x + b$
 - relies only on inner products between the test point x and data points with non-zero alpha's
- Solving the optimization problem also requires access to the data only in terms of inner products between pairs of training points



Relation SVM and logistic regression

- A classification error occurs when sign of the function does not match the sign of the class label: the zero-one loss $z = y_i f(x_i) \le 0$
- Consider error minimized when training classifier:
 - Hinge loss: $\xi_i = max(0, 1 y_i f(x_i)) = max(0, 1 z)$
 - Logistic loss: $-\log p(y_i|x_i) = -\log \sigma(y_i f(x_i)) = \log(1 + \exp(-z))$



- L2 penalty for SVM motivated by margin between the classes
- For Logistic discriminant we find it via MAP estimation with a Gaussian prior
- Both lead to efficient optimization
 - Hinge-loss is piece-wise linear: quadratic programming
 - Logistic loss is smooth : smooth convex optimization methods

Summary of discriminative linear classification

- Two most widely used linear classifiers in practice:
 - Logistic discriminant (supports more than 2 classes directly)
 - Support vector machines (multi-class extensions possible)
- For both, in the case of binary classification
 - Criterion that is minimized is a convex bound on zero-one loss
 - weight vector **w** is a linear combination of the data points $w = \sum_{n=1}^{N} \alpha_n x_n$

• This means that we only need the inner-products between data points to calculate the linear functions $f(x)=w^Tx+b$

$$= \sum_{n=1}^{N} \alpha_n x_n^T x + b$$
$$= \sum_{n=1}^{N} \alpha_n k(x_n, x) + b$$

► The "kernel" function k(,) computes the inner products

Nonlinear Classification

• 1 dimensional data that is linearly separable



- But what if the data is not linearly seperable? 0
- We can map it to a higher-dimensional space:



- General idea: map the original input space to some higher-dimensional feature space where the training set is separable
- Exercise: find features that could separate the 2d data linearly



Nonlinear classification with kernels

• The kernel trick: instead of explicitly computing the feature transformation $\varphi(\mathbf{x})$, define a kernel function K such that

 $K(\mathbf{x}_i, \mathbf{x}_j) = \boldsymbol{\varphi}(\mathbf{x}_i) \cdot \boldsymbol{\varphi}(\mathbf{x}_j)$

- Conversely, if a kernel satisfies Mercer's condition then it computes an inner product in some feature space, possibly with large or infinite number of dimensions
 - Mercer's Condition: The square N x N matrix with kernel evaluations for any arbitrary N data points should always be a positive definite matrix.
- This gives a **nonlinear decision boundary** in the original space:

$$f(x) = b + w^{T} \varphi(x)$$

= $b + \sum_{i} \alpha_{i} \varphi(x_{i})^{T} \varphi(x)$
= $b + \sum_{i} \alpha_{i} k(x_{i}, x)$



What is the kernel function that corresponds to this feature mapping ?







- What happens if we use the same kernel for higher dimensional data
 - Which feature vector $\varphi(x)$ corresponds to it ?

$$k(x, y) = (x^{T} y + 1)^{2} = 1 + 2x^{T} y + (x^{T} y)^{2}$$

- First term, encodes an additional 1 in each feature vector
- Second term, encodes scaling of the original features by sqrt(2)
- Let's consider the third term $(x^T y)^2 = (x_1 y_1 + ... + x_D y_D)^2$

$$= \sum_{d=1}^{D} (x_d y_d)^2 + 2 \sum_{d=1}^{D} \sum_{i=d+1}^{D} (x_d y_d) (x_i y_i)$$
$$= \sum_{d=1}^{D} x_d^2 y_d^2 + 2 \sum_{d=1}^{D} \sum_{i=d+1}^{D} (x_d x_i) (y_d y_i)$$

Products of two distinct elements

In total we have 1 + 2D + D(D-1)/2 features !

Original features

But the kernel is computed as efficiently as dot-product in original space

Squares

$$\varphi(x) = \left(1, \sqrt{2} x_1, \sqrt{2} x_2, \dots, \sqrt{2} x_D, x_1^2, x_2^2, \dots, x_D^2, \sqrt{2} x_1 x_2, \dots, \sqrt{2} x_1 x_D, \dots, \sqrt{2} x_{D-1} x_D\right)^T$$

Common kernels for bag-of-word histograms

• Hellinger kernel:

 $k(h_1,h_2) = \sum_d \sqrt{h_1(i)} \times \sqrt{h_2(i)}$

• Histogram intersection kernel:

 $k(h_1,h_2) = \sum_d \min(h_1(d),h_2(d))$

- Exercise: find the feature transformation, when h(d) is a bounded integer
- Generalized Gaussian kernel: $k(h_1, h_2) = \exp\left(-\frac{1}{A}d(h_1, h_2)\right)$
 - *d* can be Euclidean distance, χ^2 distance, Earth Mover's Distance, etc. See also:

J. Zhang, M. Marszalek, S. Lazebnik, and C. Schmid, Local features and kernels for classification of texture and object categories: a

comprehensive study. Int. Journal of Computer Vision, 2007

Logistic discriminant with kernels

- Let us assume a given kernel, and weight vectors $w_c = \sum_{i=1}^n \alpha_{ic} \varphi(x_i)$
 - Express the score functions using the kernel

$$f_{c}(\mathbf{x}_{j}) = b_{c} + \sum_{i=1}^{n} \alpha_{ic} \langle \varphi(\mathbf{x}_{i}), \varphi(\mathbf{x}_{j}) \rangle = b_{c} + \sum_{i=1}^{n} \alpha_{ic} k(\mathbf{x}_{i}, \mathbf{x}_{j}) = b_{c} + \alpha_{c}^{T} \mathbf{k}_{j}$$

• Where
$$\mathbf{k}_j = [k(\mathbf{x}_j, \mathbf{x}_1), \dots, k(\mathbf{x}_j, \mathbf{x}_n)]^T$$

 $\mathbf{\alpha}_c = [\alpha_{1c}, \dots, \alpha_{nc}]^T$

Express the L2 penalty on the weight vectors using the kernel

$$\langle \boldsymbol{w}_{c}, \boldsymbol{w}_{c} \rangle = \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_{ic} \alpha_{jc} k(\boldsymbol{x}_{i}, \boldsymbol{x}_{j}) = \boldsymbol{\alpha}_{c}^{T} \boldsymbol{K} \boldsymbol{\alpha}_{c}$$

- Where $[\mathbf{K}]_{ij} = k(\mathbf{x}_i, \mathbf{x}_j)$
- MAP estimation of the alpha's and b's amounts to maximize

$$\sum_{i=1}^{n} \ln p(y_i | \mathbf{x}_i) - \lambda \frac{1}{2} \sum_{c=1}^{C} \boldsymbol{\alpha}_c^T \boldsymbol{K} \boldsymbol{\alpha}_c$$

Logistic discriminant with kernels

• Recall that $p(y_i | \mathbf{x}_i) = \frac{\exp(f_{y_i}(\mathbf{x}_i))}{\sum_c \exp f_c(\mathbf{x}_i)}$ and

and
$$f_c(\mathbf{x}_i) = b_c + \boldsymbol{\alpha}_c^T \boldsymbol{k}_i$$

- Therefore we want to maximize $E(\{\alpha_c\}, \{b_c\}) = \sum_{i=1}^{n} \left(f_{y_i}(\mathbf{x}_i) - \ln \sum_c \exp f_{y_i}(\mathbf{x}_i) \right) - \lambda \frac{1}{2} \sum_c \alpha_c^T \mathbf{K} \alpha_c$
- Consider the partial derivative of this function with respect to the b's, and the gradient with respect to the alpha vectors

$$\frac{\partial E}{\partial b_c} = \sum_{i=1}^{n} \left([y_i = c] - p(c | \mathbf{x}_i) \right)$$
$$\nabla_{\mathbf{\alpha}_c} E = \sum_{i=1}^{n} \left([y_i = c] - p(c | \mathbf{x}_i) \right) \mathbf{k}_i - \lambda \mathbf{K} \mathbf{\alpha}_c$$

Essentially the same gradients as in the linear case, feature vector is replaced with a column of the kernel matrix



Support vector machines with kernels

• Minimize quadratic program

$$\min_{\mathbf{w}, \mathbf{b}, \{\xi_i\}} \quad \lambda \frac{1}{2} \mathbf{w}^T \mathbf{w} + \sum_i \xi_i$$

subject to \forall_i : $\xi_i \ge 0$ and $\xi_i \ge 1 - y_i f(\mathbf{x}_i)$

- Let us again define the classification function in terms of kernel evaluations $f(\mathbf{x}_i) = b + \boldsymbol{\alpha}^T \boldsymbol{k}_i$
 - Then we obtain a quadratic program in b, alpha, and the slack variables

$$\min_{\boldsymbol{\alpha}, b, [\xi_i]} \quad \lambda \frac{1}{2} \boldsymbol{\alpha}^T \boldsymbol{K} \boldsymbol{\alpha} + \sum_i \xi_i$$

subject to $\boldsymbol{\forall}_i: \ \xi_i \ge 0$ and $\xi_i \ge 1 - y_i (b + \boldsymbol{\alpha}^T \boldsymbol{k}_i)$



Summary linear classification & kernels

- Linear classifiers learned by minimizing convex cost functions
 - Logistic discriminant: smooth objective, minimized using gradient-based methods
 - Support vector machines: piecewise linear objective, quadratic programming
 - Both require only computing inner product between data points
- Non-linear classification can be done with linear classifiers over new features that are non-linear functions of the original features
 - Kernel functions efficiently compute inner products in (very) high-dimensional spaces, can even be infinite dimensional in some cases.
- Using kernel functions non-linear classification has drawbacks
 - Requires storing the support vectors, may cost lots of memory in practice
 - Computing kernel between new data point and support vectors may be computationally expensive (at least more expensive than linear classifier)
- The "kernel trick" also applies for other linear data analysis techniques
 - Principle component analysis, k-means clustering, regression, ...



Reading material

- A good book that covers all machine learning aspects of the course is
 - Pattern recognition & machine learning
 Chris Bishop, Springer, 2006
- For clustering with k-means & mixture of Gaussians read
 - Section 2.3.9
 - Chapter 9, except 9.3.4
 - Optionally, Section 1.6 on information theory
- For classification read
 - Section 2.5, except 2.5.1
 - Section 4.1.1 & 4.1.2
 - Section 4.2.1 & 4.2.2
 - Section 4.3.2 & 4.3.4
 - Section 6.2
 - Section 7.1 start + 7.1.1 & 7.1.2
- (Much) more on kernels: course "Advanced Learning Models" in MSIAM program

