CSC2515 – Machine Learning Sam Roweis	FEATURE SPACES 2
Socosi - Machine Learning San Koweis	• The extended representation is called a <i>feature space</i> .
Lecture 13:	 An algorithm that is linear in the feature space may be highly nonlinear in the original space if the features contain nonlinear mappings of the raw data. Thus, we can think of having "promoted" our data x into a higher-dimensional feature space z using a nonlinear mapping φ(x) and then running our original algorithm in that new space.
Kernel Machines	input space feature space
December 1, 2006	
Algorithms Based on Vector Data 1	How Should We Use the Feature Space 3
 Recall our normal approach to many classification, regression, and unsupervised learning problems: Embed the input to the problem into a vector space (e.g. Rⁿ) and then do some geometric, or linear algebraic operations. 	 The feature point z = φ(x) corresponding to an input point x is called the <i>image</i> of x; the input point x, if any, corresponding to a given feature vector z is called the <i>pre-image</i> of z. The naive way to use a feature space is to explicitly compute the image of every training point and testing point, and run our algorithm completely in feature space. Two potential problems: Problem: the feature space may be ultra-high dimensional or even infinite dimensional, so direct (explicit) calculations in feature space may not be practical or even possible.
• We can often make our algorithms more powerful by embedding the data into a richer (larger) space which includes some fixed, possibly nonlinear functions of the original measurements.	
• For example, if we measure x_1, x_2, x_3 for each datapoint, we might use the representation $z = [1, x_1, x_2, x_3, x_1^2, x_2^2, x_3^2, x_1x_2, x_2x_3]$ in a regression or classification machine.	
 We've seen this trick before: adding a bias term, quadratic regression, basis functions, generalized linear models. 	2. We may sometimes want to "bring back" an answer from feature space to input space and that involves finding pre-images which
 This trick has potential advantages (more power) and potential 	is hard and not always possible.

disadvantages (more computation, potential for overfitting).

• We could restrict ourselves to "manageable" feature spaces, but...

The Kernel Trick

• It turns out that for some special feature spaces $z = \phi(x)$, it is possible to compute the *inner product* $z_1^{\top} z_2$ between images of two input points x_1, x_2 very efficiently. This is true when the components of ϕ are the eigenfunctions of a special class of positive definite functions called *Mercer Kernels*, in which case:

$$K(x_1, x_2) = z_1^{\top} z_2 = \phi(x_1)^{\top} \phi(x_2)$$

- The key idea of kernel machines is to reduce an algorithm to one which *depends only on dot products between data vectors* and then to *replace* the dot product evaluations in feature space with kernel function evaluations in the input space.
- This "kernel trick" allows us to run algorithms in (very high dimensional) feature spaces without ever going there, provided that all we do are dot products and that we only represent feature space points that are linear combinations of known input space images.

When the Kernel Trick Works

- The kernel trick allows us to efficiently compute dot products in very high dimensional spaces using the kernel function, but it doesn't help us do addition, subtraction, outer products, etc.
- Not a big loss, since many interesting feature spaces are very high or infinite dimensional, so we couldn't even write down the results of such operations anyway.
- Because of this, in general, everything we represent in the high-dimensional feature space must be expressable as a *linear* combination of the images of training data points.
- Actually this turns out to be fine for many problems, e.g. the optimal weights in a perceptron classifier run in kernel space are guaranteed to be representable. This is true of many other algorithms, e.g. support vector machines. (There is more theory about this under (surprise surprise) the "Representer's Theorem". [originally by Kimeldorf and Wahaba, reproved by Schoelkopf et al]).

- Kernels are symmetric in their arguments: $K(x_1, x_2) = K(x_2, x_1)$.
- They are positive valued for any inputs: $K(x_1, x_2) \ge 0$.
- The Cauchy-Schwartz inequality still holds: $K^2(x_1, x_2) \leq K(x_1, x_1)K(x_2, x_2).$
- Technically, to use a function as a kernel, it must satisfy "Mercer's conditions" for a positive-definite operator.
- The intuition is easy to get for finite spaces.
- 1. Discretize x space as densely as you want into buckets x_i .
- 2. Between each two cells x_i, x_j , compute the kernel function, and write these values as a (symmetric) matrix $M_{ij} = K(x_i, x_j)$.
- 3. If the matrix is positive definite, the kernel is OK.

Examples of Kernels

$$(\phi(x) = x)$$

- Affine: $K(\mathbf{x}_1, \mathbf{x}_2) = \mathbf{x}_1^\top \mathbf{Q} \mathbf{x}_2$ (Q symmetric pos.def)
- Gaussian: $K(\mathbf{x}_1, \mathbf{x}_2) = \exp[-.5 \|\mathbf{x}_1 \mathbf{x}_2\|^2]$

• Linear: $K(\mathbf{x}_1, \mathbf{x}_2) = \mathbf{x}_1^\top \mathbf{x}_2$

- (watch scaling!)
- Sigmoid: $K(\mathbf{x}_1, \mathbf{x}_2) = \tanh(a \mathbf{x}_1^\top \mathbf{x}_2 + b)$

• Polynomial: $K(\mathbf{x}_1, \mathbf{x}_2) = (1 + \mathbf{x}_1^{\mathsf{T}} \mathbf{x}_2)^k$

- Closure rules:
- The sum of any two kernels is a kernel.
- The product of any two kernels is a kernel.
- -A kernel plus a constant is a kernel.
- -A scalar times a kernel is a kernel.

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GEOMETRY OF FEATURE SPACE

- Dot product between two points $= K(x_i, x_j) > 0$, and so all points lie in a single orthant in feature space.
- Length of a point in feature space:

$$||z_i||^2 = K(x_i, x_i)$$

(so for Gaussian kernels, everybody lies on surface of unit sphere)

• Distance between two points in feature space:

$$|z_1 - z_2||^2 = K(x_1, x_1) + K(x_2, x_2) - 2K(x_1, x_2)|$$

• Distance between a point and the mean of all others:

$$||z_k - \bar{z}||^2 = K(x_k, x_k) + \frac{1}{N^2} \sum_{ij} K(x_i, x_j) - 2 \sum_i K(x_k, x_i)$$

• Zero mean calculations in feature space:

$$< z_i - \bar{z}, z_j - \bar{z} >= K(x_i, x_j) + \frac{1}{N^2} \sum_{k\ell} K(x_k, x_\ell) - \sum_k K(x_k, x_i) \sum_k K(x_k, x_i) \sum_{k\ell} K(x_k, x_\ell) - \sum_k K(x_k, x_\ell) \sum_{k\ell} K(x_k, x_\ell) - \sum_{k\ell} K(x_k, x_\ell) \sum_{k\ell} K(x_k, x_\ell) - \sum_{k\ell} K(x_k, x_\ell) \sum_{k\ell} K(x_k, x_\ell$$

GRAM MATRIX

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- The "Gram Matrix" is the N by N symmetric matrix of all pairwise kernel evaluations: $G_{ij} = K(x_i, x_j)$.
- If you successfully "kernelize" an algorithm, then your algorithm will only need to consult/compute entries of the Gram matrix as it runs, because it depends only on dot products between the data points.
- An equivalent characterization (due to Saitoh) of Mercer's conditions is that a valid kernel generates symmetric positive definite Gram matrices for any finite sample of raw data $\{x_i\}$.

- The art of designing a kernel machine is to take a standard algorithm and massage it so that all references to the original data vectors x appear only in dot products $\langle x_i, x_j \rangle$.
- Often you can do this and obtain an exactly equivalent algorithm to the one you started with.

Sometimes you need to make small modifications.

- Thus, a kernel machine contains two modules: the algorithm and the kernel function. Choosing the kernel function is a hard problem which we won't discuss today.
- "Kernelizing" an algorithm can actually be pretty easy. How about a few examples...
- Example: K-NN Classification: Compute the distance between two points in feature space:

 $\|\mathbf{z}_1 - \mathbf{z}_2\|^2 = K(\mathbf{x}_1, \mathbf{x}_1) + K(\mathbf{x}_2, \mathbf{x}_2) - 2K(\mathbf{x}_1, \mathbf{x}_2)$

- EXAMPLE: K-MEANS CLUSTERING
- Distance between two points in feature space:

$$\mathbf{z}_1 - \mathbf{z}_2 \|^2 = K(\mathbf{x}_1, \mathbf{x}_1) + K(\mathbf{x}_2, \mathbf{x}_2) - 2K(\mathbf{x}_1, \mathbf{x}_2)$$

• Represent cluster centres as linear combinations of data points:

$$\mathbf{c}_k = \sum_i \alpha_{ik} \mathbf{z}_i$$

• Distance between a new point and a cluster centre in feature space:

$$\|\mathbf{z} - \mathbf{c}_k\|^2 = K(\mathbf{x}, \mathbf{x}) + \sum_{ij} \alpha_{ik} \alpha_{jk} K(\mathbf{x}_i, \mathbf{x}_j) - 2 \sum_i \alpha_{ik} K(\mathbf{x}, \mathbf{x}_i)$$

• True or false?

In regular K-Means (running directly in the input space), the cluster centres are always linear combinations of the data points?

• Is Kernel K-Means a good idea or a dumb idea?

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All eigenvectors with nonzero eigenvalues must lie in the span o the data, and thus can be written as linear combinations of the data (think about why...):

$$\mathbf{v} = \sum_{i} \alpha_i \mathbf{x}_i$$

• Now we can rewrite the eigenvector condition:

$$\lambda \sum_{i} \alpha_i \mathbf{x}_i = \frac{1}{N} \sum_{ij} \alpha_i \mathbf{x}_j \mathbf{x}_i^\top \mathbf{x}_j$$

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• Eigenvector condition:

$$\lambda \sum_{i} \alpha_{i} \mathbf{x}_{i} = \frac{1}{N} \sum_{ij} \alpha_{i} \mathbf{x}_{j} \mathbf{x}_{i}^{\top} \mathbf{x}_{j}$$

• Take the dot product with \mathbf{x}_k on left and right:

$$\lambda \sum_{i} \alpha_{i} \mathbf{x}_{k}^{\top} \mathbf{x}_{i} = \frac{1}{N} \sum_{ij} \alpha_{i} \mathbf{x}_{k}^{\top} \mathbf{x}_{j} \mathbf{x}_{i}^{\top} \mathbf{x}_{j}$$

• The above is true for all k, so write it as a vector equation in α :

$$N\lambda G\alpha = G^2\alpha$$
$$N\lambda\alpha = G\alpha$$

 Result: Form G, find its eigenvectors α, and use these to construct linear combinations of original data points which are the eigenvectors of the original covariance matrix.
 (Careful! Zero mean and normalization of eigenvectors.)

Example: Fisher Discriminant

- See if you can figure this one out on your own...(or look it up)
- Hint: The optimal Fisher discriminant weight can be written as the eigenvector corresponding to largest eigenvalue of a particular matrix, which is the inverse of the average within-class covariance times the average between-class covariance.
- Express the optimal weight as a linear combination of the examples and the within-class and betwee-class covariances in terms of those linear combination coefficients and the Gram matrix.
- You get a new eigenvector problem of size equal to the number of datapoints as opposed to the dimension of the inputs.

The Desire for Sparsity

- In kernel machines, the principal trick is to convert the problem into a *dual* form, which usually involves representing everything in the feature space as a linear combination of images of the training points: $\mathbf{z} = \sum_{i} \alpha_i \phi(\mathbf{x}_i)$
- \bullet Then we do all our calculations with the dual variables α_i and we never have to "touch" feature space directly.
- For very large datasets, it is desirable to have many of the coefficients α_i be exactly zero (sparsity) to reduce computational load, especially at test time.
- As a *separate trick*, different from the kernel trick, we can look for ways to make things sparse.
- These tricks are often confused, because the most famous kernel machine (the SVM) used them both.

Controlling Overfitting

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- A *third* aspect to kernel machines is how to control overfitting.
- For example, in the perceptron, if we use a very nonlinear kernel, we might always be able to separate our data exactly. Then we could be seriously overfitting.
- We can use *weight decay* to prevent this, by penalizing $||\mathbf{w}||^2$ in addition to trying to separate our training sample in the feature space. This is equivalent to *maximum margin*.
- A deep motivation for weight decay in this context comes from minimizing error bounds based on the "VC dimension" theory.
- This idea is often confused with the kernel & sparsity tricks because the most famous kernel machine (the SVM) also used weight decay to control overfitting and discussed the VC motivation.

MAXIMUM MARGIN HYPERPLANE

- \bullet Margin = minimum distance to the plane of any point.
- Principle: of all the hyperplanes that separate the data perfectly, pick the one which maximizes the margin
- Since the scale is arbitrary, we will set the numerical value of the margin to be 1.
- Now maximizing the margin is equivalent to picking the separating hyperplane that minimizes the norm of the weight vector:

 $\min \|\mathbf{w}\|^2 \qquad \text{subject to} \quad y_i[\mathbf{w}^\top \mathbf{x}_i + b] \ge 1$

x x x $(w.x) + w_0 > 0$

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Maximum Margin = Minimum Norm

- Maximizing the margin is equivalent to picking the separating hyperplane that minimizes the norm of the weight vector: $\min \|\mathbf{w}\|^2$ subject to $y_i[\mathbf{w}^\top \mathbf{x}_i + b] \ge 1$
- Use Lagrange multipliers to enforce the constraint: $\min \|\mathbf{w}\|^2 \sum_i \alpha_i (y_i [\mathbf{w}^\top \mathbf{x}_i + b] 1) \qquad \alpha_i \ge 0$
- We can convert it to *dual form* by setting partial derivatives to zero and substituting.
- This is just like ridge-regression or weight decay.
- We can also allow some points to voilate the margin by being inside it or even by being on the wrong size of it. This is achieved by adding non-negative "slack variables" ξ_i and modifying the constraints to the form:

$$y_i[\mathbf{w}^{\top}\mathbf{x}_i + b] \ge 1 - \xi_i$$

 $PRIMAL \rightarrow DUAL$

- Use Lagrange multipliers to enforce the constraint: $\min \|\mathbf{w}\|^2 - \sum_i \alpha_i (y_i [\mathbf{w}^\top \mathbf{x}_i + b] - 1) \qquad \alpha_i \ge 0$
- set $\partial/\partial \mathbf{w} = 0$ and $\partial/\partial b = 0$: $\mathbf{w}^* = \sum_i y_i \alpha_i \mathbf{x}_i$ $\sum_i y_i \alpha_i = 0$
- The dual problem is now: $\min \sum_{i} \alpha_{i} \frac{1}{2} \sum_{ij} \alpha_{i} \alpha_{j} y_{i} y_{j} \mathbf{x}_{i}^{\top} \mathbf{x}_{j}$ $\alpha_{i} \geq 0 \qquad \sum_{i} y_{i} \alpha_{i} = 0$
- This is a *quadratic programming* problem.
- It is convex. Unique solution!
- If we are allowing slack, we must also penalize the total amount of slack by adding $\sum_i \xi_i$ to the (primal) objective function.

Sparsity of Solution

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- Not only is the solution unique, but it is also sparse.
- Only the training points nearest to the separating hyperplane (ie with margin exactly 1) have $\alpha_i > 0$. These points are called the "active" points, or the *support vectors* since the final weight vector depends only on them:

$$\mathbf{w}^* = \sum_i y_i \alpha_i \mathbf{x}_i$$

- This is a lucky coincidence that has confused many people: in the case of SVM classification the two goals of controlling overfitting and inducing sparsity can both be achieved simultaneously with only a single trick: maximum margin (minimum weight norm).
- But it is not always like this.

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THANKS! IT'S BEEN FUN!

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• Last class.

Projects due Dec15, noon, by email to csc2515@cs (attachment/url) Postscript or PDF ONLY, in NIPS format, max 5 pages.

Readings must be also completed by Dec15, noon, (use the web form)

- Thanks for sticking with it. Hope you learned something, and had fun also. Sorry about all the math and about A2.
- Please send me comments/corrections for the notes so I can improve next year's course.