

## LECTURE 4:

## REGRESSION I

October 3, 2006

ERROR FUNCTION IS CRUCIAL (EG CONSTANT MODEL) 2

- Constant model says  $y = a$ , independent of  $x$ .  
(What is the constant model in classification?)
- Q: What should we use for  $a$ ?  
The mean? The median? The mode (for quantized data)?
- A: Depends on your error function (noise model).
- For squared error, the mean is best:

$$e = \sum_{n=1}^N (y_n - a)^2$$

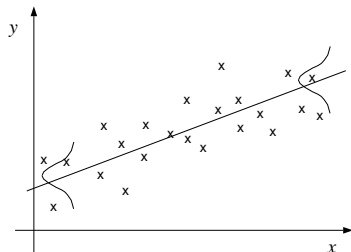
$$\frac{de}{da} = 2 \sum_n (y_n - a)$$

$$a^* = \frac{1}{N} \sum_n y_n$$

REMINDER: REGRESSION 1

1

- Multiple inputs  $x$ , mixed cts. and discrete.
- Continuous output(s)  $y$ . (Consider each separately.)
- Goal: predict output on future unseen inputs.
- Still conditional density estimation:  $p(y|x)$  (c.f. classification)
- For now, consider continuous inputs and a single output...

ABSOLUTE ERROR 3

3

- For abs error, we get the median:

$$e = \sum_{n=1}^N |y_n - a|$$

$$\frac{de}{da} = \sum_n \text{sign}[a - y_n]$$

$$= (\#y_n \text{ smaller than } a) - (\#y_n \text{ bigger than } a)$$

$$a^* = \text{median}[y_1 \dots y_N]$$

(if there are an even number of datapoints or exact duplicates, any value between the middle two is optimal)

- Moral: Even for constant model life is not so simple...

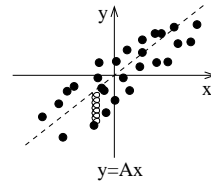
- Linear model:

$$y = \sum w_i x_i + w_0 = \mathbf{w}^\top \mathbf{x} + w_0 = \mathbf{w}^\top \tilde{\mathbf{x}}$$

Geometry: a line or hyperplane.

The bias term  $w_0$  offsets the line (hyperplane) from the origin.

Think of vertical springs connecting  $y_n$  to line  $\mathbf{w}^\top \mathbf{x}$ .



- Goal: minimize energy.
- Usually augment  $\mathbf{x}$  with constant and absorb bias into  $\mathbf{w}$ .
- Q: What's the best  $\mathbf{w}$ ?
- A: As you know, it depends on your cost function!

(The constant model corresponds to  $w_0 = a, w_i = 0$ .)

- What probabilistic model corresponds to squared error? Gaussian:

$$p(y|\mathbf{x}, \mathbf{w}) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{1}{2\sigma^2}(y - \mathbf{w}^\top \mathbf{x})^2}$$

$$\log p(y|\mathbf{x}, \mathbf{w}) = -\frac{1}{2}(y - \mathbf{w}^\top \mathbf{x})^2 + \text{const}$$

$$\log p(\mathbf{y}|\mathbf{X}, \mathbf{w}) = \sum_n \log p(y_n|\mathbf{x}_n, \mathbf{w}) \quad \text{for iid data}$$

- So minimizing squared error  $\equiv$  maximum Gaussian noise likelihood
- What if we have multiple outputs ( $y$  is a vector)?  
Turns out we can just treat each one as a separate regression problem, even if the output noise is correlated.  
(However, if the output noises are correlated *and* the noise changes from case to case, then the solutions become coupled.)

- For squared error, the problem is *linear least squares*, or *ordinary least squares*, and we can get a direct solution:

$$y = \mathbf{w}^\top \mathbf{x}$$

$$e = \sum_n (y_n - \mathbf{w}^\top \mathbf{x}_n)^2$$

$$\mathbf{w}^* = (\mathbf{X}\mathbf{X}^\top)^{-1} \mathbf{X}\mathbf{y}^\top$$

- $\mathbf{X}$  is matrix of inputs (one per column);  $\mathbf{y}$  is output row vector.
- This is one of the most famous equations in all of linear algebra: the *discrete Weiner filter*.
- It says: take the correlation between inputs and outputs, but don't be fooled by large input-input correlations.
- Predicted values are  $\hat{y}_n = \mathbf{y}\mathbf{X}^\top(\mathbf{X}\mathbf{X}^\top)^{-1}\mathbf{x}_n$ .

- Can we estimate the noise level? Yes. An unbiased estimate is:

$$\sigma^2 \approx \frac{1}{N - d - 1} \sum_n (y_n - \mathbf{w}^\top \mathbf{x}_n)^2$$

- What about the variance of parameters? Yes, also:

$$\text{var}[\mathbf{w}] = (\mathbf{X}\mathbf{X}^\top)^{-1} \sigma^2$$

- This allows us to put some crude error bars on our predictions:

$p(\hat{y}|\mathbf{x})$  is Gaussian with

$$\text{mean} = \mathbf{w}^\top \mathbf{x}$$

$$\text{variance} = \mathbf{w}^\top (\mathbf{X}\mathbf{X}^\top)^{-1} \mathbf{w} + \sigma^2$$

- There is uncertainty in the prediction both from the output noise and from our uncertainty about our estimated parameters.

- What if we use absolute error with the linear model?  
What's the equivalent of the median estimator?

$$\min_{\mathbf{w}} \sum_n |y_n - \mathbf{w}^\top \mathbf{x}_n|$$

- We need to solve a *linear programming problem*:

$$\begin{aligned} \min \sum_n t_n \\ \text{subject to} \quad -t_n \leq y_n - \mathbf{w}^\top \mathbf{x}_n \leq t_n \end{aligned}$$

- What probabilistic model corresponds to absolute error? Laplacian:

$$\begin{aligned} p(y|\mathbf{x}, \mathbf{w}) &= ae^{-a|y - \mathbf{w}^\top \mathbf{x}|} \\ \log p(y|\mathbf{x}, \mathbf{w}) &= -a|y - \mathbf{w}^\top \mathbf{x}| + \text{const} \end{aligned}$$

- What? You thought the linear model was simple enough that we don't need to regularize it? Everything needs regularization!
- Example 1: you have fewer training cases than input dimensions. Now  $\mathbf{X}\mathbf{X}^\top$  will not be invertible.
- Example 2: certain input dimensions are useless (on average) at predicting the output. But because of noise or small samples, you can always reduce the training error a tiny bit by putting huge weights on these dimensions. At test time you get killed.
- Two common solutions:
  - input subset selection
  - parameter shrinkage

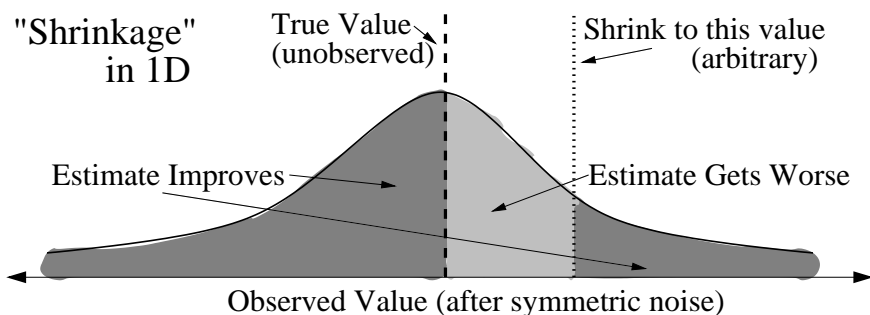
- Use only a few  $x_i$  as inputs, discard the rest.  
Advantages: introduces inductive bias, produces small models.  
Disadvantage: high variability because of binary choices.
- Forward stepwise selection: start with constant and iteratively add the single  $x_i$  which most decreases error.
- Backward stepwise selection: start with all inputs and iteratively remove the single  $x_i$  which least increases error.
- Leaps-and-Bounds: Furnival and Wilson (74) came up with a very clever branch-and-bound trick for efficiently trying *all possible* subsets. Works for up to  $\approx 40$  variables.
- Choose subset size with cross validation or F-statistic tests.

- Idea: pull ("shrink") estimated parameters towards some fixed values that do not depend on the data. ("Stein's paradox".)
- Usually we shrink towards zero (but sometimes towards the mean of some other set of weights).
- Shrinking to zero: penalize coefficients based on their size.
- For a penalty which is the sum of the squares of the weights, this is known as "weight decay" or "ridge regression":

$$\begin{aligned} y &= \mathbf{w}^\top \mathbf{x} \\ e &= \sum_n (y_n - \mathbf{w}^\top \mathbf{x})^2 + \lambda \sum_i w_i^2 \\ \mathbf{w}^* &= (\mathbf{X}\mathbf{X}^\top + \lambda I)^{-1} \mathbf{X}\mathbf{y}^\top \end{aligned}$$

where  $I$  is the identity matrix.

- We used the same trick when we were training Gaussian class-conditional classifiers.



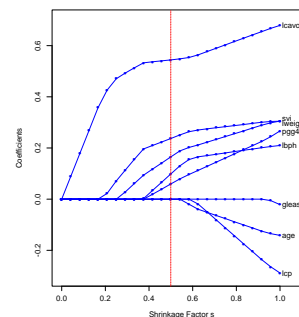
[This explanation was originally shown to me by Geoff Hinton.]

- Shrinkage has less variance but doesn't give sparse models like subset selection does. Can we get the best of both worlds?

- Lasso: squared error with absolute weight penalty.

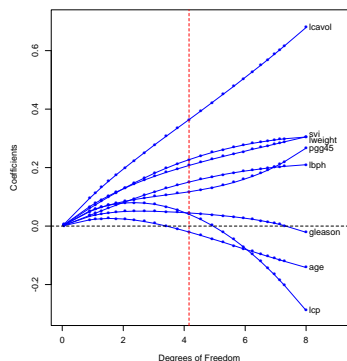
$$e = \sum_n (y_n - \mathbf{w}^T \mathbf{x})^2 + \lambda \sum_i |w_i|$$

- Requires quadratic programming to solve, but still unique optimum.



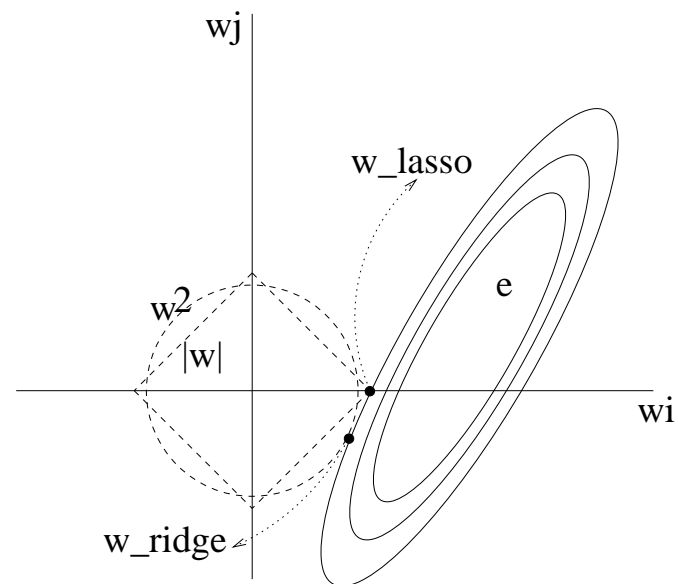
A very cool thing happens. As you increase the lasso penalty, many coefficients go exactly to zero. Why is that? Geometry!

- Set  $\lambda$  with cross-validation. (There is a trick which lets you compute the leave-one-out error very efficiently without refitting  $N$  times. See “generalized cross-validation”.)
- Don't shrink the constant (bias) term!



Warning: ridge regression is *not* invariant to input rescaling.

Often we want to “whiten/sphere” inputs first (i.e. rescale them so their sample covariance is a multiple of the identity matrix).



- We can augment the inputs, not just with a constant to get a bias term, but with lots of other things.
- If we decide beforehand on how to augment the input, this is still linear regression:

$$y = \sum_j w_j h_j(\mathbf{x})$$

- For linear regression, just use  $h_0 = 1, h_j = x_j$ .
- Optimal weights are still easy to find:

$$e = \sum_n (y_n - \mathbf{w}^T \mathbf{h}(\mathbf{x}_n))^2$$

$$\mathbf{w}^* = (\mathbf{H}\mathbf{H}^T)^{-1} \mathbf{H}\mathbf{y}^T$$

where  $\mathbf{h}(\mathbf{x})$  is a vector of basis function outputs and  $\mathbf{H}$  is a matrix with columns  $\mathbf{h}(\mathbf{x}_n)$  (sometimes called the “design matrix”).

- You can construct a special basis set that gives piecewise constant functions between pre-specified split points (“knots”)  $a_i$ :

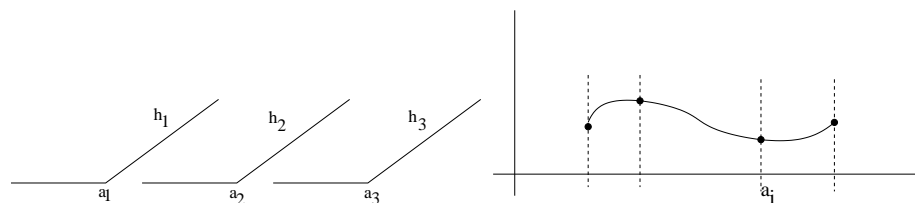
$$h_1 = (x - a_1)_+ \quad h_2 = (x - a_2)_+ \dots h_k = (x - a_k)_+ \quad h_{k+1} = x \quad h_{k+2} = 1$$

where  $(x - a_i)_+$  is the positive part of  $(x - a_i)$ .

- To enforce continuity up to the  $(r - 1)^{st}$  derivative, use

$$h_1 = (x - a_1)_+^r \dots h_k = (x - a_k)_+^r \quad h_{k+1} = x^r \dots h_{k+r+1} = 1$$

- Most common: *cubic splines*, corresponding to  $r = 3$ .
- Can also enforce linearity beyond edges: *natural cubic spline*.



- Any fixed, generalized basis set that depends on the inputs can be used to make a *generalized linear model*.
- Elements of basis are called *dictionary functions*.
- Examples include splines, radial basis functions, wavelets, etc.
- Common things to add: quadratic or other polynomial terms, sinusoidal terms, exponentials, square roots, logarithms, etc.
- Terms can depend on more than one input e.g.

$$h_j(\mathbf{x}) = x_2 x_8 x_9$$

$$h_j(\mathbf{x}) = \|\mathbf{x}\|^2$$

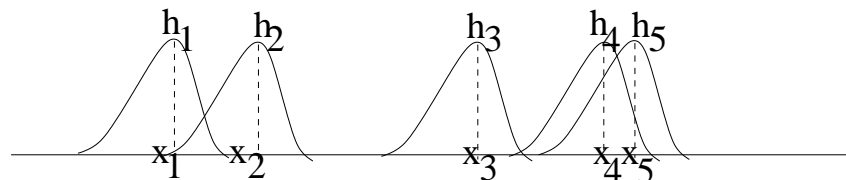
$$h_j(\mathbf{x}) = [a \leq x_i \leq b][c \leq x_j \leq d]$$

- These models can also be used for classification: as inputs to logistic/softmax regression, as a space for Fisher discriminants/Gaussians class-conditionals, KNN, etc.
- This is a simple version of the “kernel” idea (more later).

- One way to generate a nice automatic basis is to place a dictionary element on each input datapoint, whose value depends on the distance of the input from the point it is on top of:

$$h_n(\mathbf{x}) = \exp\left[-\frac{1}{2\sigma^2}\|\mathbf{x} - \mathbf{x}_n\|^2\right]$$

- Tricky part is setting  $\sigma^2$ .



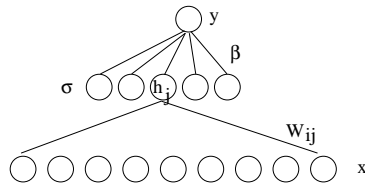
- Another generalized linear model, this time with the basis set:

$$h_j = g \left( \sum_i w_{ij} x_i \right)$$

with  $g(\cdot)$  a “squashing” function with limited outputs, e.g.

$$g(z) = \frac{1}{1 + e^{-z}} \quad g(z) = \tanh(z)$$

- The outputs  $h_j$  are known as the “hidden layer”.



- Piecewise constant 1D splines with knots at each data point value in each dimension. Also the “reflected pairs”.

$$h_{ni}(\mathbf{x}) = (x_i - x_{ni})_+ \quad h_{2ni}(\mathbf{x}) = (x_{ni} - x_i)_+$$

- Now use forward stepwise regression, choosing from these basic elements *and* any product between them and an existing dictionary element.
- Then do backwards deletion.
- Another Stanford masterpiece. (What’s in the water in Palo Alto?)

- In all the examples above, the basis functions were fixed.
- Ideally, we’d like to adjust the basis set also.  
E.g. where are the knots for splines, the centres for RBF’s, what are the input-to-hidden weights for neural networks?
- Three strategies: (1) shrinkage, (2) subset selection, (3) adaptive.

1. We can do *ridge regression* on a large generalized basis set, by penalizing the coefficients. For splines, this technique gives *smoothing splines*.
2. We can also start with a huge dictionary and try to pick a few elements. This is *subset selection* from a broader choice set.
3. Lastly, we can have a fixed number of *adaptive* elements. Next lecture we can see how to do this, for certain error functions, using *gradient descent*. But the solutions are no longer optimal.

- Regression trees (very similar to MARS but worse).
- Partial Least Squares
- Empirical Bayes (ML-II)  
Automatic Relevance Determination (ARD)
- Canonical Correlation Analysis  
(regression with a low-rank constraint)

More reading: Hastie et al. Ch4,5,9.4