Regression, function approximation

Some Common Themes Illustrated through (Generalized) MLR

Readings/Notation: I'll closely follow Bishop Ch 3.1, 3.2, which uses machine learning notation: parameters are w's (for weights), dependent variable is "t" for target, and model produces output "y".

Function Approximation / Regression/Prediction

- A predictive modeling technique
 - Given:
 - A set of input (AI) /independent (math)/ explanatory or predictor (stats) variables X
 - corresponding (set of) output/dependent/response variables Y
 - Build: a model relating X to Y
 - single value for Y given X (more common)
 - e.g. E[Y|X], the "regression of y on X.
 - Assumes Y = function of X + (zero-mean, symmetric) noise
 - Add Confidence Interval
 - Distribution of Y given X

Parametric Models

Determine functional form of model (e.g. polynomials)

- "learn" the parameters (weights) of the model using the training data.
- Example: linear regression
- Generalize: linear combination of basis functions (basis function expansion)

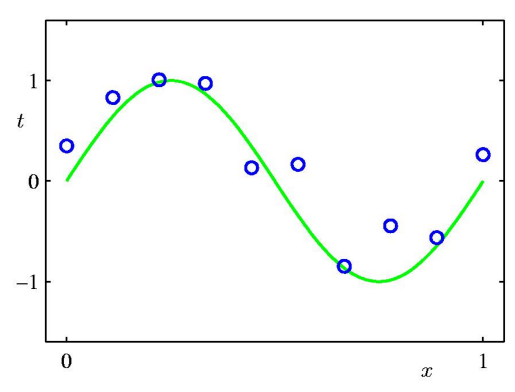
$$y(x, \mathbf{w}) = \sum_{i=0}^{M} w_i \phi_i(x) = \mathbf{w}^{\mathsf{T}} \phi(x)$$

- Special Case: linear regression.
- Special Case: polynomial: (with scalar x)

$$y(x, \mathbf{w}) = w_0 + w_1 x + \ldots + w_M x^M$$

so that the basis functions are given by $\phi_i(x) = x^i$

Polynomial Curve Fitting



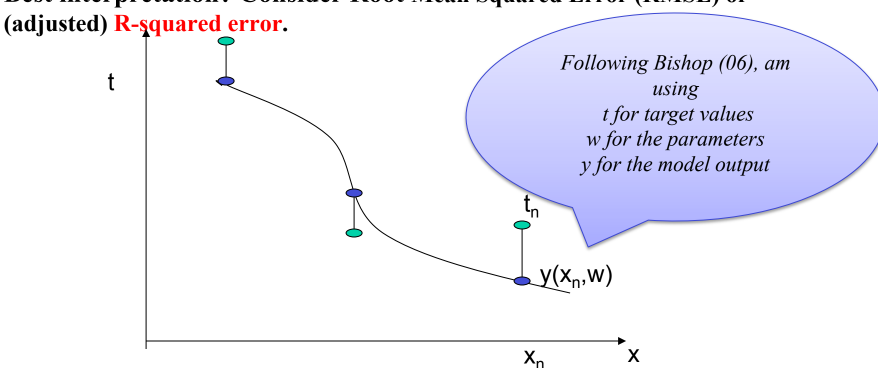
$$y(x, \mathbf{w}) = w_0 + w_1 x + w_2 x^2 + \ldots + w_M x^M = \sum_{j=0}^M w_j x^j$$

Least Squares

• Minimize sum-of-squares error (SSE) (t's are the target values)

$$E(\mathbf{w}) = \sum_{n=1}^{N} \{\mathbf{w}^{\mathsf{T}} \boldsymbol{\phi}(x_n) - t_n\}^2$$

Best interpretation? Consider Root Mean Squared Error (RMSE) or



Least Squares Solution*

Exact closed-form minimizer (ML solution)

$$\mathbf{w}^* = \left(\Phi^{\mathsf{T}}\Phi\right)^{-1}\Phi^{\mathsf{T}}\vec{t}$$
where $\vec{t} = (t_1, \dots, t_N)^{\mathsf{T}}$
Takeaway: direction solution

"Pseudo-inverse solution" and Φ is the *design matrix* given by

$$\Phi = \begin{pmatrix} \phi_0(\mathbf{x}_1) & \cdots & \phi_M(\mathbf{x}_1) \\ \phi_0(\mathbf{x}_2) & \cdots & \phi_M(\mathbf{x}_2) \\ \vdots & \vdots & \vdots \\ \phi_0(\mathbf{x}_N) & \cdots & \phi_M(\mathbf{x}_N) \end{pmatrix}$$

involves inversion of a matrix

Explicitly shows collinearity problem

Multiple outputs?

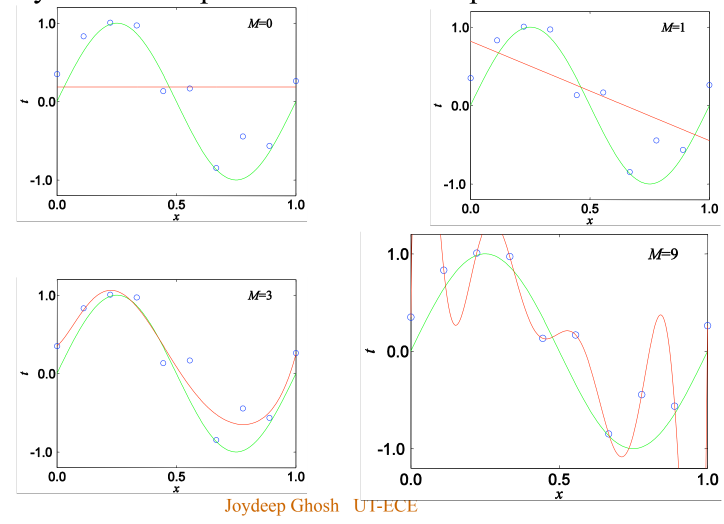
$$\mathbf{w}_{k}^{*} = \left(\mathbf{\Phi}^{\mathsf{T}}\mathbf{\Phi}\right)^{-1}\mathbf{\Phi}^{\mathsf{T}}\vec{t_{k}}$$

(psuedo-inverse portioned part shared by all outputs; rest de-coupled.)

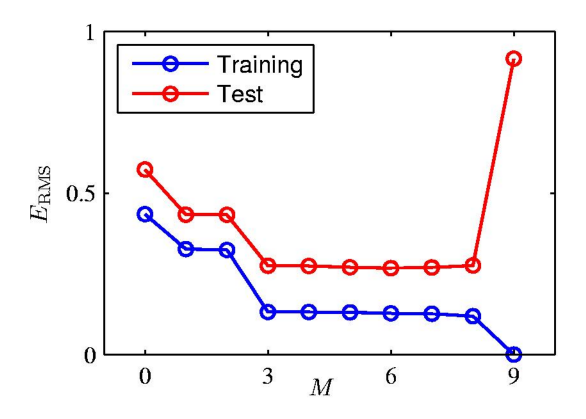
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Model Complexity and Overfitting

• "Noisy sine" example from Chris Bishop



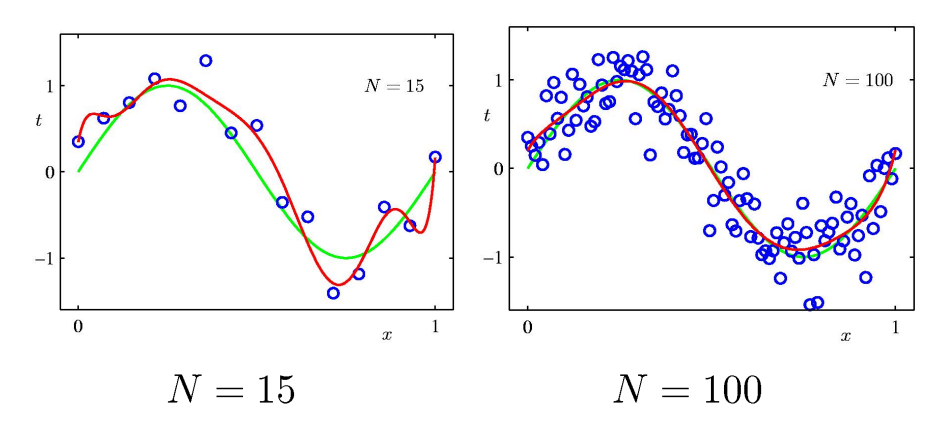
Over-fitting



Root-Mean-Square (RMS) Error vs Polynomial order

Data Set Size:

9th Order Polynomial



Regularization (to avoid overfitting)

- "regularization term" imposes penalty on less desirable solutions
 - Cost = MSE + λ Penalty (f)
 - Regularization Penalty is a functional (maps each function f onto a number)
- Popular Penalties
 - ridge regression (sum squared of weights)
 - Lasso (sum of |w|; for large λ yields sparse models)
 - Elastic net: combines both ridge and Lasso
 - number of non-zero weights
 - smoothness of function

(note: "intercept", i.e. w_0 , not included in penalties)

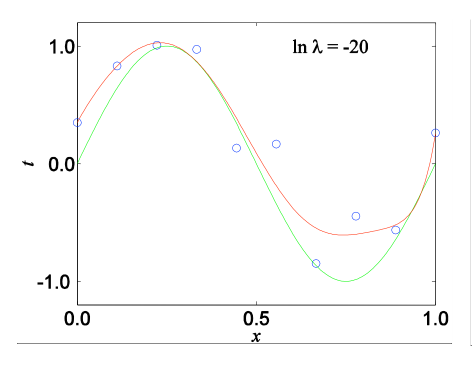
Ridge Regression Example

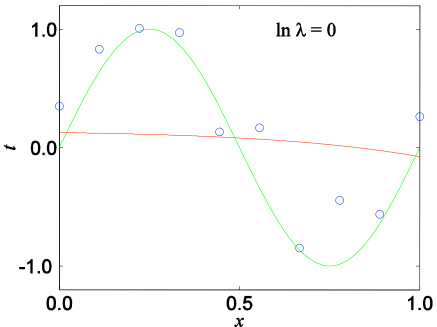
Discourage large values by adding penalty term to error

$$E(\mathbf{w}) = \sum_{n=1}^{N} \{\mathbf{w}^{\mathsf{T}} \phi(\mathbf{x}_n) - t_n\}^2 + \frac{\lambda}{2} ||\mathbf{w}||^2$$

- Also called *shrinkage* (stats) or *weight decay* (neural nets)
- The regularization coefficient λ now controls the effective model complexity
- *Closed form solution: $\mathbf{w} = (\lambda \mathbf{I} + \mathbf{\Phi}^{\mathrm{T}} \mathbf{\Phi})^{-1} \mathbf{\Phi}^{\mathrm{T}} \mathbf{t}$.
 - Leads to numerical stability as well!
- *(Related to MAP estimate in Bayesian Linear regression with isotropic covariances for both likelihood and prior, and with $\lambda = \sigma^2_{likelihood}/\sigma^2_{prior}$)

Regularized M = 9 Polynomial





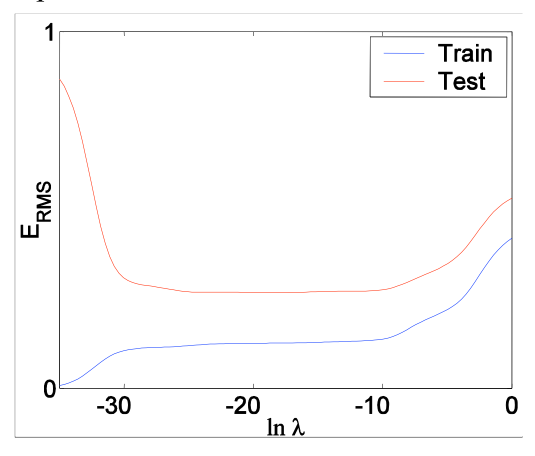
Regularized Parameters

• First col is the unregularized solution

	$\ln \lambda = -\infty$	$\ln \lambda = -20$	$\ln \lambda = 0$
w_0^{\star}	0.35	0.35	0.1273
$\mid w_{1}^{\star} \mid$	232.37	5.56	-0.0459
$w_2^{\bar{\star}}$	-5321.83	-12.27	-0.0578
$w_3^{\overline{\star}}$	48568.31	19.01	-0.0460
w_{4}^{\star}	-231639.30	-82.58	-0.0321
w_{5}^{\star}	640042.26	46.49	-0.0201
w_{6}^{\star}	-1061800.52	141.84	-0.0104
w_{7}^{\star}	1042400.18	-29.57	-0.0028
$w_{8}^{\dot{\star}}$	-557682.99	-231.55	0.0032
w_{9}^{\star}	125201.43	142.98	0.0080

Generalization

• Noisy sine problem



Evaluation

- Quality criterion for regression
 - Mean squared error (MSE) or equivalent, e.g. SSE, RMSE
 - true vs. empirical
 - normalized (R² value = % of variance explained)
 - Adjusted R²

Estimating True Performance (Formula Driven)*

- true mean squared error (MSE = SSE/N) = empirical error + complexity term
 - complexity term = f (model type, # of parameters, # of training points)
 - e.g. linear regression with N samples, P parameters
 Akaike's Final Prediction error = MSE _{empirical} (N+P) / (N P)
 - for nonlinear models, find "effective number of parameters" and plug into linear formulae

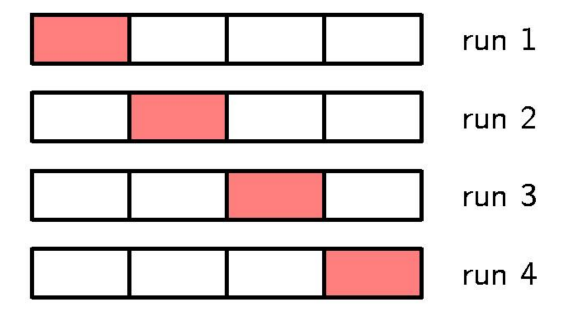
Takeaway: Formula Driven Estimates of True Performance specialized for linear models. Not so relevant in data mining context

Estimating True Performance (Data Driven)

- enough data? Use "holdout" to estimate
- Moderately large? Use k-fold cross-validation
 - extreme case (small dataset) : Leave One Out (LOO)

)

$$K = 4$$
 example



Bias-Variance Dilemma

Usually *measured* output is not a deterministic function of *given* inputs Assume: t = h(x) + zero-mean noise

• your model gives y (x). The expected squared loss,

$$\mathbb{E}[L] = \int \{y(\mathbf{x}) - h(\mathbf{x})\}^2 p(\mathbf{x}) d\mathbf{x} + \iint \{h(\mathbf{x}) - t\}^2 p(\mathbf{x}, t) d\mathbf{x} dt$$

- best predictor: $\mathbb{E}[t \mid x] = h(x)$;
- MSE_{opt} = variance of the noise inherent in the random variable t. (2nd term on RHS)
- What does the first term comprise of?

The Bias-Variance Decomposition

• Suppose we were given multiple data sets, each of size N. Any particular data set, \mathcal{D} , will give a particular function $y(x; \mathcal{D})$. We then have

$$\{y(\mathbf{x}; \mathcal{D}) - h(\mathbf{x})\}^{2}$$

$$= \{y(\mathbf{x}; \mathcal{D}) - \mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})] + \mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})] - h(\mathbf{x})\}^{2}$$

$$= \{y(\mathbf{x}; \mathcal{D}) - \mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})]\}^{2} + \{\mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})] - h(\mathbf{x})\}^{2}$$

$$+2\{y(\mathbf{x}; \mathcal{D}) - \mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})]\}\{\mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})] - h(\mathbf{x})\}.$$

The Bias-Variance Decomposition (2)

• Taking the expectation over D yields

$$\mathbb{E}_{\mathcal{D}} \left[\{ y(\mathbf{x}; \mathcal{D}) - h(\mathbf{x}) \}^{2} \right]$$

$$= \underbrace{\{ \mathbb{E}_{\mathcal{D}} [y(\mathbf{x}; \mathcal{D})] - h(\mathbf{x}) \}^{2}}_{\text{(bias)}^{2}} + \underbrace{\mathbb{E}_{\mathcal{D}} \left[\{ y(\mathbf{x}; \mathcal{D}) - \mathbb{E}_{\mathcal{D}} [y(\mathbf{x}; \mathcal{D})] \}^{2} \right]}_{\text{variance}}.$$

• (try to express both terms in words)

The Bias-Variance Decomposition (3)

Thus we can write

where
$$\operatorname{expected\ loss} = (\operatorname{bias})^2 + \operatorname{variance} + \operatorname{noise}$$

$$(\operatorname{bias})^2 = \int \{\mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})] - h(\mathbf{x})\}^2 p(\mathbf{x}) \, d\mathbf{x}$$

$$\operatorname{variance} = \int \mathbb{E}_{\mathcal{D}}\left[\{y(\mathbf{x}; \mathcal{D}) - \mathbb{E}_{\mathcal{D}}[y(\mathbf{x}; \mathcal{D})]\}^2\right] p(\mathbf{x}) \, d\mathbf{x}$$

$$\operatorname{noise} = \iint \{h(\mathbf{x}) - t\}^2 p(\mathbf{x}, t) \, d\mathbf{x} \, dt$$

Bias: how good the average model is;

Variance: how sensitive the model is to variations in data.

NOTE: the bias and variance concepts here apply to a predictive model, rather than to an estimator of a specific value.

Bias-Variance Tradeoff

- Change model type? Affect bias
- More training data: decrease variance
 - "consistent estimators" converge to ideal solution as $|D| \rightarrow$ infinity
 - For small data sets, lower complexity models may be preferred.
- Ideal solution: suitable model type & complexity

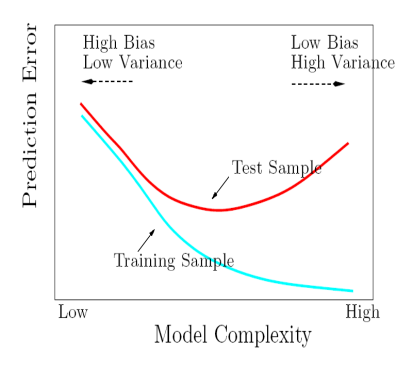
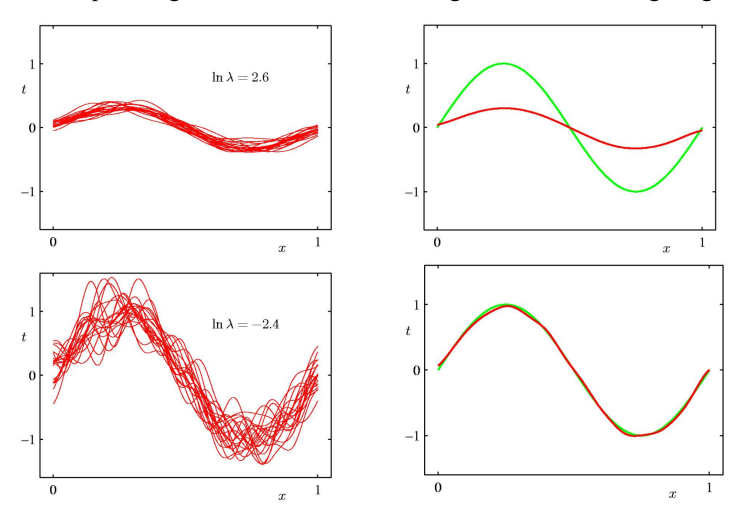


Figure 2.11: Test and training error as a function of model complexity.

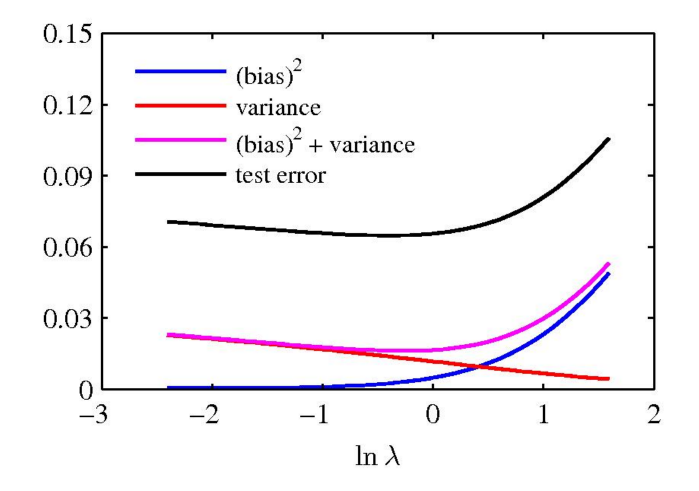
Effect of Regularization on Bias-Variance

• Bishop 06, fig 3.5. Model is sum of 24 gaussians, with ridge regression



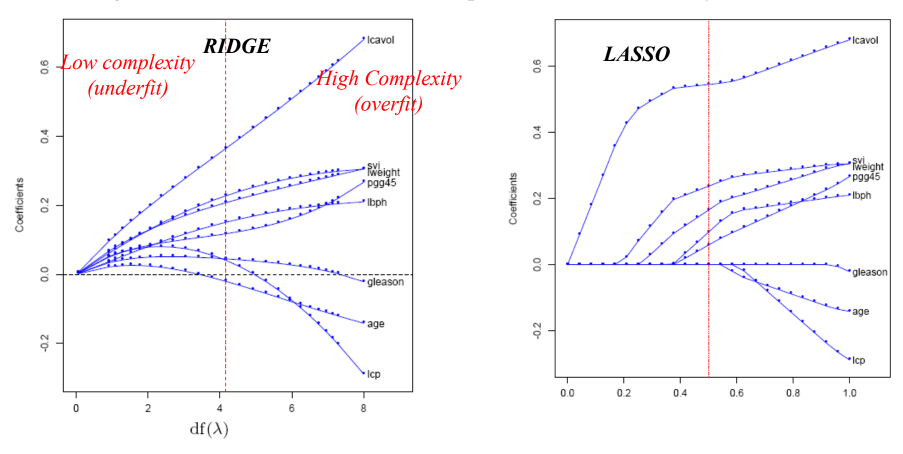
Bias-Variance vs. Regularization Amount

• What happens to the curves as amount of training data increases?



Ridge vs. Lasso

• HTF figs 3.7, 3.9: Prostate Cancer example. Red line chosen by Cross-validation



Effect on values of coefficients as "effective degrees of freedom (DOF)" is increased for (a) Ridge regression (left) and (b) Lasso (Right).

High λ translates to low DOF, so λ is being progressively decreased from left to right along the x-axis.

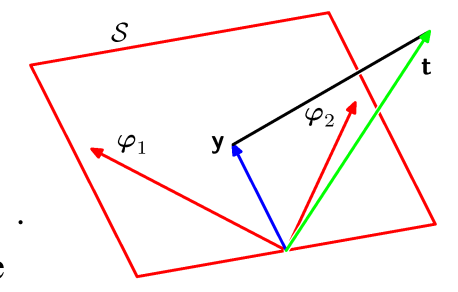
Extras

Geometry of Least Squares*

•Consider

$$\mathbf{y} = \mathbf{\Phi}\mathbf{w}_{\mathrm{ML}} = [oldsymbol{arphi}_1, \ldots, oldsymbol{arphi}_M] \, \mathbf{w}_{\mathrm{ML}}.$$
 $\mathbf{y} \in \mathcal{S} \subseteq \mathcal{T} \qquad \mathbf{t} \in \mathcal{T}$ $oldsymbol{h}$ -dimensional M-dimensional

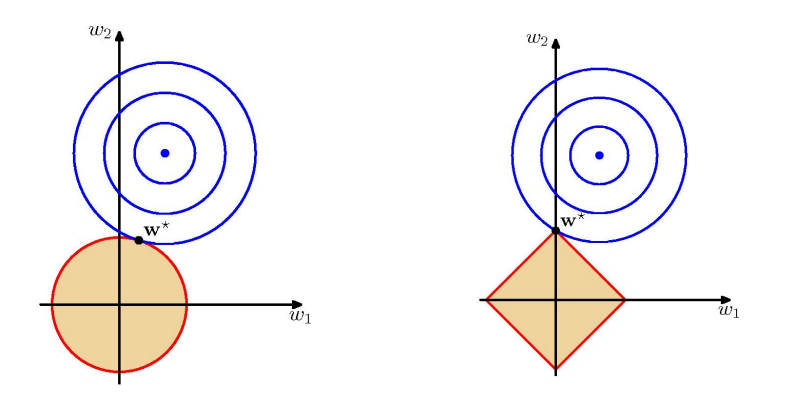
- •S is spanned by $\varphi_1, \dots, \varphi_M$
- •W_{ML} minimizes the distance between t and its orthogonal projection on S, i.e. y.



Takeaway: You are restricted by your choice of the features

Comparing Shrinkage Methods B06: fig 3.4

- ridge regression (Regularization Penalty = sum squared of weights) vs
- Lasso ((Regularization Penalty = sum of |w|)
 red: constant penalty contour; blue: unregularized error contours



Least Angle Regression (LAR; HTF 3.4.4)*

- Takeaway: Efficient procedure for fitting an entire lasso sequence with the cost of a single least squares fit.
- R code: lar Algorithm 3.2 Least Angle Regression.
 - Standardize the predictors to have mean zero and unit norm. Start with the residual r = y − y, β₁,β₂,...,β_p = 0.
 - Find the predictor x_i most correlated with r.
 - Move β_j from 0 towards its least-squares coefficient ⟨x_j, r⟩, until some other competitor x_k has as much correlation with the current residual as does x_j.
 - Move β_j and β_k in the direction defined by their joint least squares coefficient of the current residual on (x_j, x_k), until some other competitor x_l has as much correlation with the current residual.
 - Continue in this way until all p predictors have been entered. After min(N − 1, p) steps, we arrive at the full least-squares solution.

Group Lasso for Sparse Learning*

- SLEP package
 - http://www.public.asu.edu/~jye02/Software/SLEP/overview.htm
- ℓ_1 -Regularized (Constrained) Sparse Learning
- ℓ_1/ℓ_q -Regularized Sparse Learning (q>1)
- Fused Lasso
- Sparse Inverse Covariance Estimation
- Sparse Group Lasso
- Tree Structured Group Lasso
- Overlapping Group Lasso
- Takeaway: A variety of methods exist to shrink parameters in different ways