## Linear Regression

## 1. The Linear Model

Assume  $Y_i = x_i^{\top} \beta + \epsilon_i$  or  $Y = X\beta + \epsilon$  with  $X \in \mathbb{R}^{n \times p}, \beta \in$  $\mathbb{R}^{p\times 1}$ ;  $(n\geqslant p)$  and  $\mathbb{E}[\epsilon_i]=0, Var(\epsilon_i)=\sigma^2$ . X is often augmented with  $(1_{N\times 1})$  to use  $\beta_1$  as bias. df = n - p[-intercept]

## 2. Least Squares Method

LS estimator is  $\hat{\beta} = \arg\min_{\beta} ||Y - X\beta||_2^2 = (X^{\top}X)^{-1}X^{\top}Y$ Estimate  $\hat{\sigma}^2 = \frac{1}{n-p} \sum_{i=1}^n (y_i - \hat{\beta}^\top x_i)^2$  with  $\mathbb{E}[\hat{\sigma}^2] = \sigma^2$ . Gauss-Markov theorem

## LS estimator $\hat{\beta}$ is the Best Linear Unbiased Estimator, i.e

for any linear unbiased estimator  $\tilde{\beta}$  of  $\beta$ ,  $var(\tilde{\beta}) - var(\hat{\beta})$  is positive semidefinite. Univormly Minimum Variance Unbiased

## Assume $x_i$ are fixed. $\hat{\beta}$ is **UMVU**, i.e. the best estimator among

all unbiased estimators. Assumptions for Linear Model

## i Linear regression equation is correct, i.e. $\mathbb{E}[\epsilon_i] = 0 \ \forall i$ .

- ii We measure  $x_i$ 's exactly. Else, need correction (?).
- iii Error is homoscedastic, i.e.  $Var(\epsilon_i) = \sigma^2 \ \forall i$ . Else, use "Weighted LS".
- iv Errors are uncorrelated, i.e.  $Cov(\epsilon_i, \epsilon_j) = 0 \ \forall i \neq j$ . Else "Generalized LS".
- v Errors are jointly normally distributed. Else "Robust Methods".

Assume  $\mathbf{Y} = X\boldsymbol{\beta} + \epsilon$ ,  $\mathbb{E}[\boldsymbol{\epsilon}] = \mathbf{0}$ ,  $Cov(\boldsymbol{\epsilon}\boldsymbol{\epsilon}^{\top}) = \sigma^2 I$  (all assumptions

## Moments of least squares estimates

### satisfied). Then

i  $\mathbb{E}[\hat{\boldsymbol{\beta}}] = \boldsymbol{\beta} \ (\hat{\boldsymbol{\beta}} \text{ is unbiased}).$ ii  $\mathbb{E}[\hat{\mathbf{Y}}] = \mathbb{E}[\mathbf{Y}] = X\boldsymbol{\beta}$  and  $\mathbb{E}[\mathbf{r}] = \mathbf{0}$ .

iii  $Cov(\hat{\beta}) = \sigma^{2}(X^{T}X)^{-1}$ .

iv  $Cov(\hat{\mathbf{Y}}) = \sigma^2 P, Cov(\mathbf{r}) = \sigma^2 (I - P), P = X(X^\top X)^{-1} X^\top$ . If additionally  $\epsilon_i, \ldots, \epsilon_n$  i.i.d.  $\sim \mathcal{N}(0, \sigma^2)$ , then

i  $\hat{\boldsymbol{\beta}} \sim \mathcal{N}_n(\boldsymbol{\beta}, \sigma^2(X^\top X)^{-1})$ 

ii  $\hat{\mathbf{Y}} \sim \mathcal{N}_n(X\boldsymbol{\beta}, \sigma^2), \ \mathbf{r} \sim \mathbb{N}_n(\mathbf{0}, \sigma^2(I - P))$ 

Even when normality assumption doesn't hold, central limit the-

orem is a justification.

3. Regression Theory

### Weakly Universal Consistency

A sequence of regression function estimates  $\{\hat{f}_n\}$  is called weakly universally consistent if it is weakly consistent, i.e.  $\lim_{n\to\infty} \mathbb{E}\left[\int_{\mathbb{R}^p} (\hat{f}_n(x) - f^*(x))^2 P(dx)\right] = 0$ . For any estimation method, we can make the rate of convergence arbitrarily slow on some distribution, hence assumptions on Pr(X, Y), minmax approach:  $\inf_{f_n} \sup_{Pr(X,Y) \in \mathcal{P}} \text{ of the expectation.}$ 

## Empirical Risk Minimization (ERM) Impose restriction on f when minimizing training error:

i Roughness penalty Add term  $\lambda J(f)$ ii Dictionary methods Use functions of the form

 $\sum_{k=1}^{K} \theta_k \phi_k(\mathbf{x}; \eta_k)$ ii Basis function methods Same as dict but no  $\eta_k$ 

v Kernel methods and local regression: f simple in local neighborhoods (e.g. kNN is constant).

## 4. Tests and Confidence Regions

sample size), s.t.  $\hat{\beta} \sim \mathcal{N}_p(\beta, \sigma^2(X^\top X)^{-1})$  is normally distributed. Then we can test the null-hypothesis  $H_{0,j}: \beta_j = 0$ 

Assume linear model with Gaussian errors (or "large enough"

$$\frac{\hat{\beta}_j}{\sqrt{\sigma^2(X^\top X)_{jj}^{-1}}} \sim \mathcal{N}(0,1) \Rightarrow T_j = \frac{\hat{\beta}_j}{\sqrt{\hat{\sigma}^2(X^\top X)_{jj}^{-1}}} \sim t_{n-p}$$
 under the null-hypothesis  $H_{0,j}$ . Unknown  $\sigma^2$  is replaced by

 $\hat{\sigma}^2$ . Note that  $t_{n-p} \approx \mathcal{N}$ . An individual t-test for  $H_{0,j}$  gives

the effect of  $\beta_j$  after subtracting the linear effect of all  $\beta_{i\neq j}$ Note that in summary.lm, the term Std. Error is  $\sqrt{\hat{\sigma}^2(X^\top X)_{ij}^{-1}}$  $\sqrt{\hat{V}ar(\hat{\beta}_j)}$ . Finally, we can also build a confidence interval using  $\left| \hat{\beta}_j \pm \sqrt{\hat{\sigma}^2(X}^\top X)_{jj}^{-1} \cdot t_{n-p;1-\alpha/2} \right|.$ Global null hypothesis and ANOVA We can also check the global null-hypothesis  $H_0: \beta_2 = \cdots =$  $\beta_p = 0$  using an analysis of variance<, which decomposes  $||\mathbf{Y} - \bar{\mathbf{Y}}||_{2}^{2} = ||\hat{\mathbf{Y}} - \bar{\mathbf{Y}}||_{2}^{2} + ||\mathbf{Y} - \hat{\mathbf{Y}}||_{2}^{2}.$ Under the global null-hypothesis  $\mathbb{E}[\mathbf{Y}]=\mathbb{E}[\bar{\mathbf{Y}}]=const.$  (no effect of predictor variables).  $\sigma^2/\hat{\sigma}^2$  yields F-statistic:

 $F = \frac{||\hat{\mathbf{Y}} - \bar{\mathbf{Y}}||^2/(p-1)}{||\mathbf{Y} - \hat{\mathbf{Y}}||^2/(n-p)} \sim F_{p-1,n-p}$ 

under the global null-hypothesis  $H_0$ . ANOVA also yields goodness of fit  $R^2 = \frac{||\hat{\mathbf{Y}} - \bar{\mathbf{Y}}||^2}{||\mathbf{Y} - \bar{\mathbf{Y}}||^2}$ , which should be close to 1.

# partial F test - sig. of predictors in .full but not .part anova(fit.part, fit.full) ANalysis Of VAriance (ANOVA) Table

### sum sq. df i msq: sum sq./df

Reg.  $\|\hat{Y} - \bar{Y}\|^2$  p-1 ii RSE:  $\sqrt{MSE}$ 

			iii Adjusted $R^2$ : $R^2 \cdot (n-1)/df$
Total	$    Y - \bar{Y}  ^2$	n-1	
5. Checking Model Assumptions			
Tukov-Anscombo Plot			

### Error should fluctuate randomly. Error increases linearly: log-

transform  $Y \mapsto \log Y$ . Error increases with  $\sqrt{Y}$ : square-roottransform  $Y \mapsto \sqrt{Y}$ . Error has parabolic shape: add quadratic term. Error is groups: they have different Variance. plot(fit, which=1) # Tukey-Anscombe plot QQ-Plot/Normal-Plot

## Plot empirical quantiles of residuals on y versus the theoreti-

cal quantiles of  $\mathcal{N}(0,1)$  on x. If assumption holds, get straight line with intercept  $\mu$  and slope  $\sigma$ . Z-shape: long-tailed distr. Curved: skewed distr. plot(fit, which=2) # QQ-plot (qqnorm, qqline)

i Scale Location (which=3): should look random ii Cook dist (which=4): impact of sample on fit

### 6. Model Selection

Assume again  $\mathbb{E}[\epsilon_i] = 0, Var(\epsilon_i = \sigma^2)$ . Address bias-variance trade-off. Bias  $\coloneqq \mathbb{E}[\hat{f}(x)] - f(x)$ , variance  $\coloneqq q/n \cdot \sigma^2 \ (q \le p)$ .

Mallows  $C_n$  statistic

Let SSE(d) be the residual sum of squares. Then  $n^{-1} \sum_{i=1}^{n} \mathbb{E} \left[ (f(x) - \hat{f}(x))^2 \right] \approx n^{-1} SSE(d) - \hat{\sigma}^2 + 2\hat{\sigma}^2 d/n.$ Thus, we search for the model that minimizes  $C_p(\mathcal{M}) =$  $\frac{SSE(d)}{2} - n + 2d$ . Alternatively, use AIC =  $2d - 2\log \hat{L}$ , where

 $\hat{L}$  = maximal value of likelihood, or BIC. AIC is equivalent to  $C_p$  for linear Gaussian models. # All subsets regression

require(leaps) fit.all = regsubsets(y~., data=data) p.regsubsets(fit.all)

Forwards and backwards selection Forward selection: (i) Start with empty model. (ii) (Greedily)

fit.bw = step(fit.full, direction="backward")

fit.fw = step(fit.empty, direction="forward",

Keep adding variable that reduces the residual sum of squares cal polynomial, i.e. we replace  $m_x$  with  $\beta_1 + \sum_{i=2}^p \beta_i (x_i - x)^{i-1}$ the most. (iii) When done, pick submodel which minimizes  $C_p$ . (usually p = 2 or p = 4). Often better at edges and yields first Backward selection: (i) Start with full model. (ii) (Greedily) Keep excluding predictor that increases the residual sum of fit.loess <- loess(y ~ x, data=data.frame(x=x, y=y\_pert), squares the least. (iii) When done, pick submodel which mini-→ span=0.2971339, surface='direct') mizes  $C_p$ . Backwards selection typically better but more expenfit.loess.pred <- predict(fit.loess, newdata=x) sive. When  $p \ge n$ , use forward selection. Both methods prone to overfitting — p-values (and similar values) are not valid anymore The hat matrix Sand effects look too significant. We want to construct S with  $\hat{\mathbf{Y}} = S\mathbf{Y}$ , i.e. the linear op-# Backward / forward selection fit.empty = lm(y~1, data=data)fit.full = lm(y~., data=data)

predict.lm(fit, newdata, interval=..) poly(formula, degree=d) # polynomial fits of the data in the → predictive variables up to degree d

## Density Estimation Kernel estimator

Estimate density  $\hat{f}(x)=\frac{1}{nh}\sum_{i=1}^n w((x-X_i)/h).$  Kernels include (i) rectangular  $(w(x)=0.5\cdot 1_{|x|<1}),$  (ii) triangular, or (iii) Gaussian. We require  $\int_{\mathbb{R}} K(x)dx = 1$ . The bandwidth parameter h is crucial and determines the "smoothness" of the density Choosing a bandwidth h

A simple approach is using k-nearest neighbors, i.e. h(x) = $\max_{x_i \in KNN_k(x)} ||x - x_i||_2$  with tuning parameter k. Note that  $\int_{\mathbb{D}} K(x)dx = 1$  might be violated. Naturally, the bandwidth also induces a bias-variance trade-off. Note that MSE(x) $\mathbb{E}\left[\left(\hat{f}(x) - f(x)\right)^2\right] = \left(\mathbb{E}[\hat{f}(x)] - f(x)\right)^2 + Var(\hat{f}(x)), \text{ so we can}$ try to minimize the integrated MSE over all points to find the best bandwidth.

## Density estimation in higher dimensions

Basically use  $\hat{f}(\mathbf{x}) = \frac{1}{nh^d} \sum_{i=1}^n K((\mathbf{x} - \mathbf{X}_i)/h)$  with a Kernel that supports vectors. The Gaussian kernel is the only one that is radially symmetric. Note that in higher dimensions, density estimation becomes very hard, due to data points becoming very

## Nonparametric Regression

Nonparametric regression with one predictor variable, i.e.  $Y_i$  $m(x_i) + \epsilon_i$  with  $\epsilon_{1:n}$  i.i.d and  $\mathbb{E}[\epsilon_i] = 0$ . We want  $m(x) = \mathbb{E}[Y|x]$ and "some" smoothness. Kernel regression estimator

$$\hat{m}(x) = \frac{\sum_{i=1}^{n} \omega_i Y_i}{\sum_i \omega_i} = \underset{m_x \in \mathbb{R}}{\arg \min} \sum_{i=1}^{n} \omega_i (Y_i - m_x)^2$$
(3.1)

with  $\omega_i = K\left(\frac{x_i - x}{h}\right)$  a kernel centered at  $x_i$  and bandwidth h. As h small  $\rightarrow$  large then (high variance)  $\rightarrow$  (high bias). For  $x_i$  equidistant there exists  $h_{opt} = f(\sigma_\epsilon^2, m''(x))$  which can be ksmooth(x, y, kernel="normal", bandwidth=0.2, x.points=x)\$y

fit.lo<-lokerns(X, Y, x.out=X, hetero=TRUE, is.rand=TRUE) fit.gl<-glkerns(X, Y, x.out=X, hetero=TRUE, is.rand=TRUE)</pre>

i Bias  $(mean(\hat{y} - y_{true}))$ : high  $\rightarrow$  under fitting ii Variance (measure of fluctuation): high → over fitting

erator mapping the labels to the predictions. Given the regression (smoothing) function s, we compute  $S_{ij} = s(\mathbf{x}, \mathbf{e}_i, h)$ 

with  $\mathbf{e}_j$  the j-th unit vector. Then  $Cov(\hat{m}(\mathbf{x})) = Cov(S\mathbf{Y}) =$  $SCov(\mathbf{Y})S^{\top} = \sigma_{\epsilon}^{2}SS^{\top}$ . Set  $df = \mathbf{tr}S$  and estimate  $\hat{\sigma}_{\epsilon}^{2} =$  $\sum_{i=1}^{n} (Y_i - \hat{m}(x_i))^2 / (n - df)$ . Then •  $\widehat{s.e.}(\hat{m}(x_i)) = \sqrt{\widehat{Var}(\hat{m}(x_i))} = \hat{\sigma}_{\epsilon \lambda} \sqrt{(SS^\top)_{i,i}}$ 

Local polynomial regression estimator

Instead of finding a local constant  $m_x$  we can also find a lo

•  $\hat{m}(x_i) \approx \mathcal{N}\left(\mathbb{E}[\hat{m}(x_i)], Var(\hat{m}(x_i))\right)$ 

•  $I = \hat{m}(x_i) \pm 1.96 \cdot \widehat{s.e.}(\hat{m}(x_i)) \rightarrow \text{(pointwise) CI}$ # Construct S matrix (or hatMatrix() from sfsmics)

N <- length(x); Eye <- diag(N) S.nw <- matrix(0, nrow=N, ncol=N) for (j in 1:N) { S.nw[, j] <- ksmooth(x, y\_, kernel="normal", bandwidth=0.2, x.points=x)\$y # Compute standard error est.nw <- ksmooth(x=x, y=y, kernel="normal", bandwidth=0.2, → x.points=x)\$y sig\_sq.nw <- sum((y - est.nw)^2) / (N - sum(diag(S.nw)))

# Splines without having to specify knots: Find $\arg\min_{m\in C^0(\mathbb{R})}\sum_{i=1}^n(Y_i-m(x_i))^2+\lambda\int_{\mathbb{R}}m''(z)^2dz$ . Minimizer

se.nw <- sqrt(sig\_sq.nw \* diag(S.nw \%\*\% t(S.nw)))

Smoothing splines and penalized regression

is finite dimensional — it is a cubic spline that can be computed using a set of basis functions  $m_{\lambda}(x) = \sum_{j=1}^{n} \beta_{j} B_{j}(x)$  or  $||\mathbf{Y} - B\boldsymbol{\beta}||^2 + \lambda \boldsymbol{\beta}^{\mathsf{T}} \Omega \boldsymbol{\beta} \Rightarrow \hat{\boldsymbol{\beta}} = (B^{\mathsf{T}} B + \lambda \Omega)^{-1} B^{\mathsf{T}} \mathbf{Y}$ . Choose  $\lambda$ on the scale of  $df = \mathbf{tr}(S_{\lambda})$ . fit.ss = smooth.spline(x, y, df=df) ss.preds = predict(fit.ss, newdata=x)\$y

## Cross Validation Let $(X_1, Y_1), \ldots, (X_n, Y_n)$ i.i.d $\sim P$ . We would like to compute

 $\mathbb{E}_{(X_{\text{new}}, Y_{\text{new}})}[\rho(\mathbf{Y}_{\text{new}}, \hat{m}_{\text{train}}(X_{\text{new}}))]$ Constructing cross-validation datasets

## Approaches include

Leave-one-out CV:  $n^{-1} \sum_{i=1}^{n} \rho\left(Y_i, \hat{m}_{n-1}^{(-i)}(X_i)\right)$  ca. unbiased. k-fold CV:  $K^{-1}\sum_{i=1}^{K}|\mathcal{B}_k|^{-1}\sum_{i\in\mathcal{B}_k}\rho\left(Y_i,\hat{m}_{n-|\mathcal{B}_k|}^{(-\mathcal{B}_k)}(X_i)\right)$ 

Smaller variance than LOOCV. Random division: Like k-fold, but build  $\mathcal{B}_k$  by sampling without

replacement (≈ 10%). Usually fastest. Tricks using hat matrix

For linear fitting operators and the loss  $\rho(y,x) = (y-x)^2$  we

can exploit the hat matrix and get the full LOOCV result in a

$$n^{-1}\sum_{i=1}^{\hat{n}}\left(Y_i-\hat{m}_{n-1}^{(-i)}(X_i)\right)^2=n^{-1}\sum_{i=1}^{n}\left(\frac{Y_i-\hat{m}(X_i)}{1-S_{ii}}\right)^2.$$
 It can be cheaper to just compute  $\operatorname{tr}(\mathcal{S})$  (instead of all  $S_{ii}$ ),

 $GCV = \frac{n^{-1} \sum_{i=1}^{n} (Y_i - \hat{m}(X_i))^2}{(1 - n^{-1} \mathbf{tr}(S))^2}.$ 

$$GCV = \frac{1}{(1 - n^{-1}\mathbf{tr}(S))^2}$$
The two equations coincide if  $S_{ii} = c \ \forall i$ .

One-standard-error rule

Choose the most parsimonious model whose error is no more than one standard error above the error of the best model.

## the predictive power of a model or algorithm. Nonparametric Bootstrap Let $Z_{1:n}$ i.i.d $\sim P$ with $Z_i = (X_i, Y_i), X_i \in \mathbb{R}^p, Y_i \in \mathbb{R}$ , and let $\hat{\theta}_n = g(Z_{1:n})$ be an estimator. We would like to know the dis-Tribution of $\hat{\theta}_n$ . We approximate **P** by the *empirical distribution* $\hat{P}_n$ . Then we can repeatedly sample $Z_{1:n}^* \sim \hat{P}_n$ independently and compute $\hat{\theta}_n^* = g(Z_{1:n}^*)$ . The histogram (or any density estimator) then describes the distribution of $\hat{\theta}_n^*$ . The algorithm a) Sample (with replacement) $Z_{1:n}^*$ i.i.d $\sim \hat{P}_n$ . b) Compute the bootstrapped estimator $\hat{\theta}_n^* = g(Z_{1:n}^*)$ . c) Repeat B times to obtain $\hat{\theta}_{\pi}^{*1:B}$ . d) Approximate $\mathbb{E}^*[\hat{\theta}_n^*] \approx B^{-1} \sum_{i=1}^B \hat{\theta}_n^{*i}$ and $Var^*(\hat{\theta}_n^*) \approx$ $\hat{\theta}_n^* \approx \text{empirical } \alpha\text{-quantile of } \hat{\theta}_n^{*1:B}$ Central limit theorem Let $X_i$ be a random variable with $\mathbb{E}[X_i] = \mu$ and $Var(X_i) = \sigma^2$ Then $n^{-1} \sum_{i=1}^{n} X_i \xrightarrow{n \to \infty} \mathcal{N}(\mu, \sigma^2/n)$ . Bootstrap consistence Consistency of the bootstrap typically holds if the limiting distribution of $\hat{\theta}_n$ is Normal and if $Z_{1:n}$ are i.i.d. Mathematically, for an increasing sequence $a_n$ and $\forall x$ , $\mathbb{P}[a_n(\hat{\theta}_n - \theta) \leq x]$ $\mathbb{P}^*[a_n(\hat{\theta}_n^* - \hat{\theta}_n) \leqslant x] \xrightarrow{P} 0 \text{ as } n \to \infty. \text{ Then } Op^*(\hat{\theta}_n^*)/Op(\hat{\theta}_n) \xrightarrow{P} 0$ 1 with $Op \in \{Var, \mathbb{E}\}$ . Bootstrap confidence intervals Given bootstrap consistence, we can compute confidence interi quantile / quantile: $[q_{\hat{\Theta}} * (\alpha/2), q_{\hat{\Theta}} * (1 - \alpha/2)]$ iii normal: $2\hat{\Theta} - \overline{\hat{\Theta}^*} \pm q_X(1 - \alpha/2) \cdot \hat{sd}(\hat{\Theta})$ $(X \sim \mathcal{N}(0, 1); \text{ corrects for bias } \hat{\Theta} - \hat{\Theta}^*)$ require("boot") tm = function(x, ind) {mean(x[ind], trim = 0.1)} res.boot = boot(data=sample, statistic=tm, R=1000)

Bootstrap

## ii rev. quantile: $[\hat{\Theta} - q_{\hat{\Theta} - \hat{\Phi}} (1 - \alpha/2), \hat{\Theta} - q_{\hat{\Theta} - \hat{\Phi}} (\alpha/2)]$ Note $\hat{q}_{\alpha} = q_{\alpha}^* - \hat{\theta}_n$ with $q_{\alpha}^* = \alpha$ -bootstrap quantile of $\hat{\theta}_n^*$ . Thus $[\hat{\theta}_n - \hat{q}_{1-\alpha/2}, \hat{\theta}_n - \hat{q}_{\alpha/2}] = [2\hat{\theta}_n - q_{1-\alpha/2}^*, 2\hat{\theta}_n - q_{\alpha/2}^*].$ boot.ci(res.boot, conf=0.95, type=c("basic","norm","perc")) # "basic"=reverse quantile, "norm"=normal, "perc"=quantile require(MASS); mle = fitdistr(boogg, "gamma")\$estimate boot.est = matrix(NA, nrow=R, ncol=1) for (i in 1:R) { boogg.s = rgamma(N, shape=mle[1], rate=mle[2]) # boogg.s = sample(boogg, N, replace=T) # NP boot.est[i] = quantile(boogg.s, probs=0.75) # Quantile / Percentile quantile(boot.est, probs=c(a/2, 1-a/2)) mean.est = mean(boot.est) sd.hat = sqrt(1/(R-1)\*sum((boot.est - mean.est)^2)) 2\*est-mean.est + c(-1,1)\*qnorm(1-a/2)\*sd.hat# Reverse quantile / Basic quantile est - quantile(boot.est-est, probs=c(1-a/2, a/2)) Double bootstrap Idea: Find $\alpha'$ s.t. actual coverage of bootstrap CI $I^*(1-\alpha')$ is i Draw BS sample $Z^*$ . Sample from $Z^*$ to obtain $Z^{**}$ . Compute CI $I^{**}(1-\alpha)$ for $\hat{\Theta}^*$ based on B draws $Z^{**}$ . Compute coverage of $\hat{\Theta}$ by $I^{**}$ (1 or 0). ii Repeat i) M times to obtain M coverage values. Compute mean to obtain actual coverage of $I^{**}$ .

Vary  $\alpha'$  and repeat previous steps to find  $\alpha'$  with

coverage $(I^{**}(1-\alpha')) = 1 - \alpha$ . Use CI  $I^*(1-\alpha')$ .

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Efron's parametric and nonparametric bootstrap can be described as "simulating from an estimated model" and can be used for states inference (confidence intervals and testing) and estimating version when P_{\hat{\Theta}} is a good fit (e.g. known model structure P)
                                                                                               and few data points available.
                                                                                                require(MASS); fit.gamma = fitdistr(boogg, "gamma")
                                                                                                fun.theta = function(x) {quantile(x, probs=0.75)}
                                                                                                fun.gen = function(x, mle) {rgamma(length(x), shape=mle[1],
                                                                                                 \rightarrow rate=mle[2])}
                                                                                                res.boot = boot(data, fun.theta, R=1000, sim="parametric",
                                                                                                 → ran.gen=fun.gen, mle=fit.gamma$estimate);
                                                                                               Bootstrap error estimate
                                                                                               Generalization error (loss \rho(y, m(x))) of model m (fitted to full
                                                                                               data set) can be estimated by fitting models m^{*,i} to bootstrap
                                                                                               samples and computing
                                                                                                  • errors on full data set e^{*,i} = n^{-1} \sum_{i=1}^{n} \rho(y, m^{*,i}(x_i))
                                                                                                  • OOB errors e_{ob}^{*,i} = n_{ob,i}^{-1} \sum_{i=1}^{n_{ob,i}} \rho(y_{ob,i}, m^{*,i}(x_{ob,i}))
   (B-1)^{-1}\sum_{i=1}^{B}\left(\hat{\theta}_{n}^{*i}-B^{-1}\sum_{i=1}^{B}\hat{\theta}_{n}^{*j}\right)^{2}. Then \alpha-quantile of The error of m is then approximated by R^{-1}\sum_{i=1}^{R}e^{*,i}.
                                                                                                 Classification
                                                                                             Given (X_1, Y_1), \ldots, (X_n, Y_n) i.i.d. with Y_i \in \{0, \ldots, J-1\}, determine \pi_j(x) = \mathbb{P}[Y = j \mid X = x] \quad \forall j = 0, 1, \ldots, J-1. The optimal classifier is C_{Bayes}(x) = \arg\max_{0 \le j \le J-1} \min_{T \in \mathcal{T}} (x). Then so equivalent to the finite-dimensional problem
                                                                                              Bayes risk for the 0-1-loss is \mathbb{P}[\mathcal{C}_{Bayes}(X_{new}) \neq Y_{new}].
                                                                                              Discriminant analysis
                                                                                               Bayes formula
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Parametric Bootstrap

## $\min_{f \in \mathcal{H}_K} \frac{1}{n} \sum_{i=1}^n L(y_i, \sum_{j=1}^n \alpha_j K_{x_j}(\mathbf{x}_i)) + \lambda \alpha^\top \mathbf{K} \alpha,$ where $\sum_{j=1}^{n} \alpha_j K_{x_j}(\mathbf{x}_i) = (\mathbf{K}\alpha)_i$ E.g. squared loss regression: $\arg\min_{\alpha} \frac{1}{n} ||\mathbf{y} - \mathbf{K}\alpha||^2 + \lambda \alpha^{\top} \mathbf{K}\alpha = (\mathbf{K} + n\lambda \mathbf{I})^{-1} \mathbf{y}$ **LDA:** Assume $X \mid Y = j \sim \mathcal{N}(\mu_j, \Sigma), \mathbb{P}[Y = j] = p_j$ . Then by $\pi_{j}(x) = \frac{f_{X|Y=j}(x) \cdot p_{j}}{\sum_{k=0}^{J-1} f_{X|Y=k}(x) \cdot p_{k}}$ with each $f_{X|Y=j}$ a Gaussian $\mathcal{N}(\mu_j, \Sigma)$ . We can estimate $\mu_j$ and $\Sigma$ by the (closed-form) MLEs, and we also need a prior for Y, which often is fixed as $\hat{p}_i = n_i/n$ . This results in $\hat{\delta}_i(x) = (x - \hat{\mu}_i/2)^{\top} \Sigma^{-1} \hat{\mu}_i + \log(\hat{p}_i)$ with linear (in x) decision boundaries $\hat{\delta}_{i}(x) = \hat{\delta}_{i}(x)$ and $C(x) = \arg\max_{i} \hat{\delta}_{i}(x)$ . **QDA:** Now we assume different $\Sigma_i$ for each class and obtain quadratic decision boundaries $\hat{\delta}_{i}(x) = -\log(\det(\hat{\Sigma}_{i}))/2 - (x - \log(\det(\hat{\Sigma}_{i}))/2)$ $(\hat{\mu}_i)^{\top} \hat{\Sigma}_i^{-1} (x - \hat{\mu}_i)/2 + \log(\hat{p}_i)$ . The price: $J \cdot p(p+1)/2$ parameters (for all $\Sigma_i$ ) vs. p(p+1)/2 (symmetry of $\Sigma$ ) for a single $\Sigma$ $(\mu \ (J \cdot p) \text{ and priors } (J-1) \text{ are the same}),$

## class\_lda = lda(x=df[, c("x1", "x2")], grouping=df[, "y"]) Logistic regression for binary classification Given some model $g: \mathbb{R}^p \to \mathbb{R}$ (e.g. a linear model) we can use the logistic transform $\pi \mapsto \log(\pi/(1-\pi))$ to get probabilities: $\log(\pi(x)/(1-\pi(x))) = g(x)$ and $\pi(x) = 1/(1+\exp(-g(x)))$ This implies $Y_i \sim \text{Bernoulli}(\pi(x_i))$ (e.g. weighted coin flip) The likelihood is thus $\prod_{i=1}^n \pi(x_i)^{Y_i} (1 - \pi(x_i))^{1-Y_i}$ . We typically estimate $\beta$ using gradient descent. As $n \to \infty$ we can asymptotically compute the standard errors $\widehat{s.e.}(\hat{\beta}_i)$ and t-test statistics $\hat{\beta}_i/\widehat{s.e.}(\hat{\beta}_i) \sim \mathcal{N}(0,1)$ (under $H_{0,j}: \beta_j = 0$ ). fit = glm(Y~., data=data, family="binomial") mean((predict(fit, type="response") > 0.5) == data\$Y) Linear predictors Note that both LDA and Logistic regression are linear in the prediction variables. For LDA that comes from the Gaussian assumption (i.e. "linearization" of the true distribution), for Logistic regression it comes from the linear log-odds function.

# Multiclass case (J > 2)a) J classes $\to J$ binary variables: $\tilde{\pi}_j(x) = \frac{p^i j^{(x)}}{\sum_{j=0}^{J-1} \hat{\pi}_j(x)}$ Using multinomial distribution (parametric linear logistic) (see c) "Reference class" $\log(\pi_j(x)/\pi_0(x)) = g_j(x)$ d) Pairwise 1-vs-1, fitting $\binom{J}{2} \cdot p$ parameters Let $g_{tree}(\mathbf{x}) = \sum_{r=1}^{M} \beta_r \mathbf{1}_{[\mathbf{x} \in \mathcal{R}_r]}$ , where $\{\mathcal{R}_1, \dots, \mathcal{R}_M\}$ is a partition of $\mathbb{R}^p$ . The function is piecewise cone) Exploiting "ordered" classes with proportional odds

## $\min_{\mathbf{w}} \sum_{i=1}^n L(y_i, f(x_i)) + \frac{1}{2C} ||\mathbf{w}||^2 \text{ with } f(\mathbf{x}) = \mathbf{w}^\top \mathbf{x} + b \text{ and hinge loss } L(y, f) = (1 - yf)_+ \text{ with } (u)_+ = max(u, 0). \text{ Kernel}$ SVM solution is of the form $\hat{\beta}_0 + \hat{f}(\mathbf{x}) = \hat{\beta}_0 + \sum_{i=1}^n \hat{\alpha}_i K(\mathbf{x}, \mathbf{x}_i)$ require(e1071) tune.out <- tune(svm, Y~., data=dat, kernel="radial", $\mathbb{R}^p \to \mathbb{R}$ and ensembles them (which comes at the loss of inter-→ ranges=list(cost=c(...), gamma=c(...))) summary (tune.out) Flexible regression and classification methods 1. Additive models Decompose multivariate function as $g_{add}$ : $\mathbb{R}^p \to \mathbb{R}, x \mapsto$ $\mu + \sum_{j=1}^{p} g_j(x_j)$ with $g_j(\cdot) : \mathbb{R} \to \mathbb{R}$ , $\mathbb{E}[g_j(X_j)] = 0$ . The zeromean requirement for each $g_i(\cdot)$ makes the problem well posed. This approach is a generalization of linear models, and similarly can not model interaction terms $g_{j,k}(x_j,x_k)$ . Avoid the curse of dimensionality by construction. To fit a model, let $S_i$ be a smoothing technique (e.g. Nadaraya-Watson kernel estimators). Then, the backfitting algorithm Compute $\hat{\mu} = n^{-1} \sum_{i=1}^{n} Y_i$ and initialize $\hat{g}_j(\cdot) := 0$ . Cycle through the indices j = 1, 2, ..., p, 1, 2, ... and update $\hat{g}_j = \mathcal{S}_j(\mathbf{Y} - \hat{\mu}\mathbf{1} - \sum_{k \neq j} \hat{g}_k)$ . Stop each function at conver-• Normalize the functions: $\tilde{g}_j(\cdot) = \hat{g}_j(\cdot) - n^{-1} \sum_{i=1}^n \hat{g}_j(X_{ij})$ . This basically makes the algorithm repeatedly solve the 1dimensional fitting problem. The algorithm may be slow but often works and can use any 1-dimensional fitting technique. When fitting Additive models in R with the function gam, the smoothers $S_i$ are penalized regression splines, and the degrees of freedom for each spline (i.e. each variable) will be determined through cross-validation. fit <-gam(Y~s(x1)+s(x2)+..., data=data) # wrapFormula() plot(fit, pages=1, shade=TRUE) # show effect predictor sfsmisc::TA.plot(fit, labels="o") Classification and Regression Trees

stant. When a partition is given, we can estimate  $\hat{\beta}_r$  =

 $\sum_{i=1}^{n} Y_i 1_{[\mathbf{x} \in \mathcal{R}_r]} / \sum_{i=1}^{n} 1_{[\mathbf{x} \in \mathcal{R}_r]}$ . For multiclass classification

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compute \hat{g}_{i=1}^{*}(\cdot). Repeat B times.
ii Aggregate bootstrap estimates with \hat{g}_{\mathrm{B}ag}(\cdot)
   B^{-1} \sum_{i=1}^{B} \hat{g}_{i}^{*}(\cdot) \approx \mathbb{E}^{*}[\hat{g}^{*}(\cdot)].
Note that \hat{g}_{\mathrm{B}ag}(\cdot) = \hat{g}(\cdot) + (\mathbb{E}^*[\hat{g}^*(\cdot)] - \hat{g}(\cdot)). We can re-
                                        bootstrap bias estimate
duce variance at price of higher bias (at least for trees). In fact,
for many x, Var(\hat{g}_{Bag}(x)) < Var(\hat{g}(x)). We can use larger trees
(higher variance) to balance the bias-variance trade-off.
For Subsample aggregating (Subbagging), we
(X_1^*, Y_1^*), \ldots, (X_m^*, Y_m^*) without replacement (e.g. with
m = \lfloor n/2 \rfloor, which can be cheaper overall and is equivalent to
Bagging in some simple settings.
AdaBoost.M1
 i Initialize the weights w_i = 1/n, i = 1, \ldots, n.
ii For m = 1, \ldots, M:
  a) Fit a weak classifier G_m(x) to training data using w_i's.
  b) Compute \operatorname{err}_m = \frac{\sum_{i=1}^n w_i I\{y_i \neq G_m(\mathbf{x}_i)\}}{\sum_i n}
  c) Compute \alpha_m = \log \{(1 - \operatorname{err}_m) / \operatorname{err}_m \}.
  d) Set w_i \leftarrow w_i \times \exp\left[\alpha_m I\left\{y_i \neq G_m\left(\boldsymbol{x}_i\right)\right\}\right], i = 1, \dots, n.
 ii Output G(\mathbf{x}) = \text{sign} \left\{ \sum_{m=1}^{M} \alpha_m G_m(\mathbf{x}) \right\}.
 i RStudio: Tools \rightarrow Global Options \rightarrow Code \rightarrow Display
ii ROC: FPR (x) vs TPR (y)
iii P[> |t|]: large (> 0.05) \rightarrow \beta_i \approx 0 \rightarrow \beta_i not relevant; small
   \rightarrow \beta_i! = 0 \rightarrow \beta_i relevant & reject H_{i,0} (R: pt(...))
 print(.., digits=..); table(df$Class, y.pred); residuals(..)
 factor(Class) # for classification; optim(..); termplot(...)
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### RKHS are Hilbert spaces generated by positive definite kernels, iii Apply ii) to one cell of the current partition (select like above). called Mercer Kernels. E.g. Mercel kernel: $K(\mathbf{x}_i, \mathbf{x}_i)$ v Iterate iii) until until specified max, partition size is achieved. $\phi(\mathbf{x}_i)^{\top}\phi(\mathbf{x}_i)$ v Prune tree by removing leaves resulting in smalles increase in Mercer theorem some (CV) metric. The size of a tree T is the number of leaves (=1 + cuts). For Any mercer kernel K admits eigen decomposition of the form some goodness-of-fit measure R(T) (e.g. SSE, NLL), the cost- $K(\mathbf{x}, \mathbf{x'}) = \sum_{j=1}^{\infty} \gamma_j \phi_j(\mathbf{x}) \phi_j(\mathbf{x'})$ , where $\gamma_j$ are decreasingly complexity measure is $\mathcal{R}_{\alpha}(\mathcal{T}) = \mathcal{R}(\mathcal{T}) + \alpha \cdot \text{size}(\mathcal{T})$ . For some $\alpha$ , ordered, $\sum_{j\geqslant 1} |\gamma_j|^2 < \infty$ and $\phi_{jj}$ is a set of orthonormal we thus choose $\mathcal{T}(\alpha) = \arg\min_{\mathcal{T} \subset \mathcal{T}_M} \mathcal{R}_{\alpha}(\mathcal{T})$ . The parameter functions in $L^2$ . The RKHS induced by kernel is $\mathcal{H}_K$ $\alpha$ is chosen by CV. The 1 s.e. rule says: Choose smallest tree $\left\{f: \mathbb{R}^p \to \mathbb{R}: f(\mathbf{x}) = \sum_{i=1}^{\infty} c_i \phi_i(\mathbf{x}), \ ||f||_{\mathcal{H}_K}^2 = \sum_{i=1}^{\infty} \frac{c_i^2}{\gamma_i} < \infty \right\}$ such that its performance is at most one standard error larger than the minimal one The norm corresponds to the inner product $\langle f, h \rangle_{\mathcal{H}_{K}}$ num\_leafs<-length(unique(tree\$where)) depth <- trunc(log2(max(as.integer(row.names(tree\$frame))))) $\sum_{i=1}^{\infty} \frac{c_i d_i}{\gamma_i}$ ., where $f = \sum_{i=1}^{\infty} c_i \phi_i(\mathbf{x})$ and $g = \sum_{i=1}^{\infty} d_i \phi_i(\mathbf{x})$

Manually Calculating Predicted Class

**RKHS** 

Representer Theorem

i Predict class for x:  $\arg\min_{k\in[1,J]}\sum_{j=1}^{J}L(j,k)\cdot\pi_{j}(x)$ 

ii E.g loss for predicting class  $c: \sum_{j=1}^J L(j,c) \cdot \pi_j(x)$ 

library(rpart); require(rpart.plot); tree = rpart(y~., data=data, control=rpart.control(cp=0.0, plotcp(tree); cps = tree\$cptable # cp =  $\alpha/\mathcal{R}_{\alpha}(\mathcal{T}_{\alpha})$ idx.min = which.min(cps[, "xerror"]) std.min = cps[idx.min, "xstd"] cp.1se = cps[abs(cps[,"xerror"]-cps[idx.min,"xerror"])<std.min,] pruned.tree = prune.rpart(tree, cp=cp.1se[1, "CP"]) rf <- randomForest(Boston.train, y.train, xtest=Boston.test, → ytest=y.test, ntree=ntree, mtry=ncol(Boston.train)) rf\$mse[ntree] # 00B error; rftestmse[ntree] tail(rf\$err.rate) # cumulative error rate

 $\hat{\pi}_j(\mathbf{x}) = \sum_{i=1}^n \mathbf{1}_{[Y_i = j]} \mathbf{1}_{[\mathbf{x} \in \mathcal{R}_r]} / \sum_{i=1}^n \mathbf{1}_{[\mathbf{x} \in \mathcal{R}_r]} \text{ for } \mathbf{x} \in \mathcal{R}_r.$ 

ii Split  $\mathcal{R}$  at d in dimension j, where d is from the set of midpoints of observed values. Select j, d s.t. neg. log-likelihood

Add the resulting two cells and remove the refined one.

Greddy algorithm to find axes parallel partition:

i Initialize M=1 subset  $\mathcal{P}=\{\mathcal{R}=\mathbb{R}^p\}$ 

decrease is maximized by refinement.

# Bagging and Boosting

## Bagging and Subbagging Bootstrap $\mathbf{agg}\mathrm{regating}$ (bagging) (mostly on trees), uses $\hat{g}(\cdot)$

i Generate bootstrap sample  $(X_1^*, Y_1^*), \dots, (X_n^*, Y_n^*)$  and