Chapter 2 (week 2)

31 Jan 2023

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"God is in every leaf of every tree"

- From Andrew Gelman (blog)
- "No problem is too small or too trivial if we really do something about it." (Dyson (2005) quoting Richard Feynman)
- (an excuse for going down rabbit holes?)

Dyson, Freeman. 2005. "Wise Man." New York Review of Books, October. https://www.nybooks.com/articles/2005/10/20/wise-man/.

feature selection

- $feature \approx a$ column of the model matrix
- termwise selection, e.g.
 - all columns associated with a categorical variable
 - all columns of a basis expansion (polynomial etc.) of a continuous variable
- columnwise selection
 - fine for prediction
 - silly for inference?
- selection maintaining the **principle of marginality** (Venables 1998)
 (i.e., don't drop lower-order effects from a model
 - (i.e., don't drop lower-order effects from a model containing interactions)
- ¿ a way to **merge categories** on the fly (based on rarity, correlation, predictive ability)?

Venables, W. N. 1998. "Exegeses on Linear Models." In. 1998 International S-PLUS User Conference. Washington, DC. http://www.stats.ox.ac.uk/pub/MASS3/Exegeses.pdf.

why select?

- save memory
- save "flops" (floating-point operations)
- optimize bias-variance tradeoff

- optimize data collection
- parsimonious/simple explanations (e.g. rms::fastbw in R)

why select (2)?

- save memory: OK
- save flops, optimize B-V
 - which is best: soft (ridge), semi-soft (lasso/SCAD), hard (stepwise/subset) penalization?

selection: filters, wrappers, embedded methods

Jović, Brkić, and Bogunović (2015)

- filters: standalone recipes
 - e.g. minimum-redundancy maximum relevance (mrMR) (Peng, Long, and Ding 2005)
 - * similar to stepwise forward, but no estimation done (compute mutual information)
 - * greedy
 - general, low-cost
- wrappers: applied around specific methods
 - e.g. stepwise regression
 - general, evaluates prediction
- embedded methods: integrate estimation and selection
 - e.g. lasso etc.
 - most efficient? can combine shrinkage and selection

Jović, A., K. Brkić, and N. Bogunović. 2015. "A Review of Feature Selection Methods with Applications." In 38th International Convention on Information and Communication Technology, Electronics and Microelectronics (MIPRO), 1200–1205. https://doi.org/10.1109/MIPRO.2015.7160458.

Peng, Hanchuan, Fuhui Long, and C. Ding. 2005. "Feature Selection Based on Mutual Information Criteria of Max-Dependency, Max-Relevance, and Min-Redundancy." *IEEE Transactions on Pattern Analysis and Machine Intelligence* 27 (8): 1226–38. https://doi.org/10.1109/TPAMI.2005.159.

stepwise abuse

- stepwise regression for **prediction** may be fine (Murtaugh 2009)
 - selection based on AIC etc. more sensible than with p-values

Murtaugh, Paul A. 2009. "Performance of Several Variable-Selection Methods Applied to Real Ecological Data." *Ecology Letters* 12 (10): 1061–68. https://doi.org/10.1111/j.1461-0248.2009.01361.x.

- note $\Delta AIC \propto p$ value, if using columnwise/1-df steps
 - * $\Delta \log(L) \leftrightarrow \Delta AIC = 0 \leftrightarrow p = 0.16$
 - * leave-one-out cross-validation (LOOCV) asymptotically equiv. to AIC (Stone (1977); but see CV)
- for **inference**, terrible if done naively (but see Blanchet, Legendre, and Borcard (2008))
 - see CrossValidated
 - unstable, biased estimates; overconfident inference ("snooping")
- ESL: stepwise as a jumping-off point/comparator for different

Stone, M. 1977. "An Asymptotic Equivalence of Choice of Model by Cross-Validation and Akaike's Criterion." J. Royal Stat. Soc. B 39 (1): 44–47. https://www.jstor.org/stable/2984877.

Blanchet, F. Guillaume, Pierre Legendre, and Daniel Borcard. 2008. "Forward Selection of Explanatory Variables." *Ecology* 89 (9): 2623–32. https://doi.org/10.1890/07-0986.1.

POLLS

- did you learn to do stepwise regression in a class? Were you warned about its limitations?
- have you used stepwise regression? were you aware of its limitations at the time?
- have you used SR "in real life"? for prediction or inference?

contrasts for categorical variables

- expanding categorical variables to dummy variables
- automatically handled by model.matrix() in R (StatsModels.jl:modelmatrix in Julia)

```
library(palmerpenguins)
library(tidyverse)
library(faux)
set.seed(101)
pp <- penguins[sample(nrow(penguins)), c("species", "island")] ## scramble
head(model.matrix(~species+island, pp))</pre>
```

```
(Intercept) speciesChinstrap speciesGentoo islandDream islandTorgersen
1
2
            1
                              1
                                             0
3
            1
                              0
                                                                          0
                                                         1
4
            1
                              0
                                             1
                                                         0
5
            1
                              1
                                             0
                                                         1
                                                                          0
6
            1
                              1
  ## faux makes nicer factors!
  ## rename variables/**idempotent** operations: f(f(x)) = f(x) x
  pp2 <- mutate(pp, across(where(is.factor), contr_code_treatment))</pre>
  head(model.matrix(~species+island, pp2))
  (Intercept) species. Chinstrap-Adelie species. Gentoo-Adelie
1
2
            1
                                                              0
3
            1
                                                              0
4
            1
                                      0
                                                              1
5
            1
                                                              0
                                                              0
  island.Dream-Biscoe island.Torgersen-Biscoe
1
2
                                              0
3
                                              0
4
                                              0
5
                                              0
                    1
6
                    1
                                              0
  colnames(model.matrix(~species*island, pp2))
[1] "(Intercept)"
[2] "species.Chinstrap-Adelie"
[3] "species.Gentoo-Adelie"
[4] "island.Dream-Biscoe"
[5] "island.Torgersen-Biscoe"
[6] "species.Chinstrap-Adelie:island.Dream-Biscoe"
```

[7] "species.Gentoo-Adelie:island.Dream-Biscoe"

```
[8] "species.Chinstrap-Adelie:island.Torgersen-Biscoe"
```

- [9] "species.Gentoo-Adelie:island.Torgersen-Biscoe"
 - identifiability constraints: leave out one category
 - post-hoc evaluation (e.g. emmeans R pkg)
 - penalized methods

regression, again

- hat matrix $(\mathbf{H} = \mathbf{X}(\mathbf{X}^{\top}\mathbf{X})^{-1}\mathbf{X}^{\top}\mathbf{y})$ as projection matrix from \mathbf{R}^{N} to \mathbf{R}^{p}
 - (what if we first transformed **X** to be orthonormal?)
- non-full-rank case $(rank(\mathbf{X}) < p)$
 - non-unique solutions
 - may break our linear algebra, depending on what we use

```
X <- matrix(c(1:3, 2*(1:3)), ncol = 2)
y <- 1:3
Matrix::rankMatrix(X)</pre>
```

```
[1] 1
attr(,"method")
[1] "tolNorm2"
attr(,"useGrad")
[1] FALSE
attr(,"tol")
[1] 6.661338e-16

try(solve(X %*% t(X)))
```

```
Error in solve.default(X \%*\% t(X)) : Lapack routine dgesv: system is exactly singular: U[2,2] = 0
```

```
try(qr.solve(qr(X),y))
```

Error in qr.solve(qr(X), y) : singular matrix 'a' in solve

```
lm.fit(X, y)$coefficients
```

x1 x2 1 NA

Q: how would we do this with SVD (svd), or Cholesky decomposition (chol)?

side note: Bessel's correction

- ESL gives $\hat{\sigma}^2 = \frac{1}{N-p-1} \cdot \text{RSS}$
 - note p doesn't include the constant term/intercept column
- note unbiased estimate of the residual variance
- MLE would give RSS/N
- unbiased estimate of resid std. error divides by N-1.5; minimum MSE (for Normal distribution) divides by N+1 (!)
- bias is scale-dependent $(E(f(x)) \neq f(E(x)))$ in general) and might not matter as much as you think

prostate cancer example

 data exploration: pairs(., gap = 0) (can be extended with panel function); corrplot::corrplot.mixed(., lower="number", upper = "ellipse"); GGally::ggpairs(). Can use faraway::prostate.

```
## a bit of data exploration
pp <- (prostate</pre>
```

train/test error

• hardly worth it for simple regression problems (measures like adjusted R^2 and AIC(c) give reasonable estimates of out-of-sample error)

Gauss-Markov theorem

- simple
- applicable as long as data are independent and homoscedastic (iid is stronger)
- MVUE (minimum-variance *unbiased* estimator)
- but **not** necessarily minimum MSE!

regression by orthogonalization (3.2.3)

- build up regression by successive orthogonalization
 - regress \mathbf{x}_j on residuals of all previous columns $(\mathbf{z}_0, \mathbf{z}_1, \dots, \mathbf{z}_j)$ to get coefficients $\hat{\gamma}_{\ell j}$, residual \mathbf{z}_j .

 regress \mathbf{y} on \mathbf{z}_p to get $\hat{\beta}_p$ order???
- Gram-Schmidt orthogonalization (successive projection)
- if **Z** is the residual columns and Γ is the (upper-triangular) matrix of $\gamma_{\ell j}$, then $\mathbf{X} = \mathbf{Z}\Gamma$
- if $\mathbf{D} = \text{Diag}(||\mathbf{z}_i||)$
- and $\mathbf{X} = \mathbf{Z}\mathbf{D}^{-1}\mathbf{D}\Gamma = \mathbf{Q}\mathbf{R}$ with \mathbf{Q} orthonormal, \mathbf{R} upper triangular
- \rightarrow standard decomposition!

multiple outputs

- somewhat niche problem ...
- changing y to Y, β to B, the algebra mostly stays the same
- separate coefficients for each problem
- if homoscedastic, no need to consider correlation of observations!

return to subset/stepwise selection

- still not sure it's worth it
- can update efficiently based on QR decomp
- forward-stagewise: less efficient
- digression: inefficiency as a virtue
 - improve bias-var tradeoff by worsening fit
 - early stopping, dropout, etc. etc.

shrinkage methods

ridge

- L2 penalty on coefficients
- predictors must be normalized! (scale of β_j depends on
- scale of x_j) equivalence between penalty $(+\lambda \sum \beta^2)$ and constraint $(\sum \beta^2 \le t)$
 - ("one-to-one correspondence" between λ and t, but not simple!)
- add $\lambda \mathbf{I}$ in the normal equations
- works for non-full-rank problems

Bayesian analogue

• analogous to setting iid Gaussian prior on individual β parameters

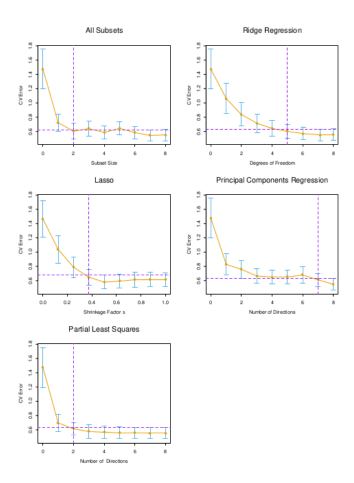


Figure 1: ESL fig 3.7

- log-posterior = log-likelihood + log-prior $\propto \sigma^2 RSS + \lambda \sum \beta^2$
- MAP (maximum *a posteriori*) estimate, **not** "proper" Bayesian est (mode, not mean, of posterior)

solving ridge by QR

- note that we can solve ridge regression by introducing pseudo-observations (data augmentation)
- set

$$\mathbf{B} = \left(\begin{array}{c} \mathbf{X} \\ \sqrt{\lambda} \mathbf{I} \end{array}\right)$$

- and $\mathbf{y}^* = (\mathbf{y} \ 0)$
- so that $\mathbf{B}^{\top}\mathbf{B} = \mathbf{X}^{\top}\mathbf{X} + \lambda I$ and the residual sum of squares is unchanged
- and solving $(\mathbf{B}^{\top}\mathbf{B})\beta = \mathbf{B}\mathbf{y}^*$ by QR decomposition (Atlas 2013)
- $\ensuremath{\mathcal{C}}$ a trick for solving for successive λ values faster ... ?

Atlas. 2013. "QR Factorization for Ridge Regression." *Mathematics Stack Exchange*. https://math.stackexchange.com/questions/299481/qr-factorization-for-ridge-regression.

singular value decomposition

• if $\mathbf{X} = \mathbf{U}\mathbf{D}\mathbf{V}^{\top}$ then

$$\begin{split} \mathbf{X}(\mathbf{X}^{\top}\mathbf{X})^{-1}\mathbf{X}^{\top}\mathbf{y} &= \mathbf{U}\mathbf{D}\mathbf{V}^{\top}(\mathbf{V}\mathbf{D}\mathbf{U}^{\top}\cdot\mathbf{U}\mathbf{D}\ \mathbf{V}^{\top})^{-1}\mathbf{V}\mathbf{D}\mathbf{U}^{\top}\mathbf{y} \\ &= \mathbf{U}\mathbf{D}\mathbf{V}^{\top}(\mathbf{V}\mathbf{D}^{2}\mathbf{V}^{\top})^{-1}\mathbf{V}\mathbf{D}\mathbf{U}^{\top}\mathbf{y} \\ &= \mathbf{U}\mathbf{U}^{\top}\mathbf{v} \end{split}$$

- and ridge translates to $\sum \mathbf{u}_j \frac{d_j^2}{d_i^2 + \lambda} \mathbf{u}_j^{\top} \mathbf{y}$
- i.e. shrinking the $j^{ ext{th}}$ principal component by $\frac{d_j^2}{d_i^2 + \lambda}$
- (if inputs are orthonormal all coefficients are shrunk equally)

effective df

- this also shows that effective df = trace of hat matrix = $\sum \frac{d_j^2}{d_j^2 + \lambda}$
- see also Hastie (2020)

ridge projection

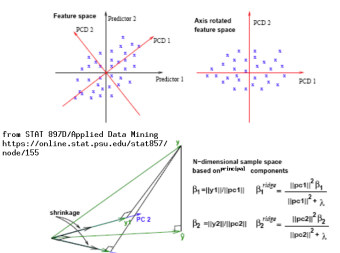


Figure 2: The Geometric interpretation of principal components and shrinkage by ridge regression.

lasso

- L1 regularization
- sparsity-inducing
- least-angle regression (LARS): nice, but superseded (also, doesn't work for GLMs)
- glmnet et al. use cyclic/pathwise coordinate descent (Friedman, Hastie, and Tibshirani 2010) (also in Julia analogue)
 - plus "warm-start" algorithm

Hastie, Trevor. 2020. "Ridge Regularization: An Essential Concept in Data Science." *Technometrics* 62 (4): 426–33. https://doi.org/10.1080/00401706.2020.1791959.

Friedman, Jerome, Trevor Hastie, and Rob Tibshirani. 2010. "Regularization Paths for Generalized Linear Models via Coordinate Descent." *Journal of Statistical Software* 33 (1): 1–22. https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2929880/.

pathwise coordinate descent

- ESL § 3.8.6
- $\tilde{\beta}_k(\lambda)$ is **current** estimate of $\beta_k(\lambda)$. Then

$$R = \frac{1}{2} \sum_{i=1}^{N} \left(y_i - \sum_{k \neq j} x_{ik} \tilde{\beta}_k(\lambda) - x_{ij} \beta_j \right)^2 + \lambda \sum_{k \neq j} |\tilde{\beta}_k(\lambda)| + \lambda |\beta_j|$$

- i.e. univariate lasso on j with k parameters fixed
- or lasso on $partial\ residual\ (y_i \tilde{y}_i^{(j)}) = y_i \sum_{k \neq j} \tilde{\beta}_k(\lambda)$
- solution:

$$\tilde{\beta}_j(\lambda) \leftarrow S\left(\sum_{i=1}^N x_{ij}(y_i - \tilde{y}_i^{(j)}), \lambda\right)$$

- where $S(t, \lambda) = \text{sign}(t)(|t| \lambda)$
- can't do all λ automatically, but warm start algorithm works quickly
 - start with large λ such that all coefficients $\rightarrow 0$
 - reduce in small steps, using values from previous λ to initialize
- ¿how much worse does this get for other loss functions (e.g. GLMs)?

other penalties

- could use L_p penalization with $1 (equivalent to a generalized normal or exponential power prior: <math>\propto \exp\left(|(x-\mu)/s|^p\right)$ (gnorm package)
- elastic-net (penalty $\propto \alpha \sum \beta^2 + (1 \alpha) \sum |\beta|$)
 - computationally nicer and sparsity-inducing

ridge vs lasso vs best-subset vs elastic net

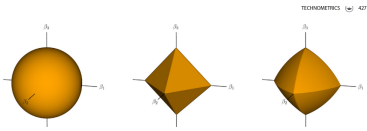


Figure 1. Constraint balls for ridge, lasso, and elastic-net regularization. The sharp edges and corners of the latter two allow for variable selection as well as shrinkage

and more penalties

- fit unrestricted (linear regression or other) model on lassoselected variables (why??) (Zhao, Witten, and Shojaie 2021)
- relaxed lasso: re-fit lasso on selected variables (why??)
- smoothly clipped absolute deviation (SCAD): $\lambda |\beta| \to J_{\alpha}(\beta, \lambda)$, with

$$\frac{dJ_a(\beta,\lambda)}{d\beta} = \lambda \cdot \mathrm{sign}(\beta) \left[I(|\beta| \leq \lambda) + \frac{(a\lambda - |\beta|)_+}{(a-1)\lambda} I(||\beta| > \lambda) \right]$$

for $a \geq 2$

• adaptive lasso $\approx |\beta|^{1-\nu}$

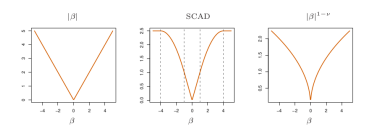


FIGURE 3.20. The lasso and two alternative non-convex penalties designed to penaltize large coefficients less. For SCAD we use $\lambda=1$ and a=4, and $\nu=\frac{1}{2}$ in the last panel.

grouped lasso

• ESL § 3.8.4; Yuan and Lin (2006)

Zhao, Sen, Daniela Witten, and Ali Shojaie. 2021. "In Defense of the Indefensible: A Very Naïve Approach to High-Dimensional Inference." *Statistical Science* 36 (4): 562–77. https://doi.org/10.1214/20-STS815.

Yuan, Ming, and Yi Lin. 2006. "Model Selection and Estimation in Regression with Grouped Variables." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)* 68 (1): 49–67. https://doi.org/10.1111/j.1467-9868.2005.00532.x.

- lasso on groups of parameters: compute $||\beta_\ell||_2$ by group $(\beta_\ell$ is the sub-vector of parameters in group ℓ , of length $p_\ell)$
- RSS criterion plus penalty

$$\lambda \sum_{\ell=1}^L \sqrt{p_\ell} ||\beta_\ell||_2$$

* reduces to lasso if every parameter is in a separate group $(||c||_2 = |c|)$ if c is a scalar) * ESL: "encourages sparsity at both the group and individual levels" * ¿ridge-like within groups, lasso-like between groups;

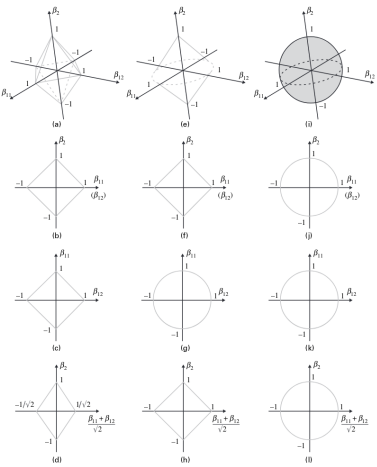


Fig. 1. (a)–(d) l_1 -penalty, (e)–(h) group lasso penalty and (i)–(l) l_2 -penalty

- ¿has someone written a formula-to-groupedlasso interface;
- sparse grouped lasso: like elastic net (convex combination) but for regular lasso + grouped lasso

finding packages

```
a1 <- available.packages()
grep("lasso", rownames(a1), ignore.case = TRUE, value = TRUE)</pre>
```

[1]	"abglasso"	"ALassoSurvIC"	"BayesianGLasso"
[4]	"biglasso"	"bolasso"	"BTdecayLasso"
[7]	"BTLLasso"	"CARlasso"	"CDLasso"
[10]	"cglasso"	"clogitLasso"	"covglasso"
[13]	"CVglasso"	"DIFlasso"	"DLASSO"
[16]	"DWLasso"	"elasso"	"extlasso"
[19]	"gamlss.lasso"	"genlasso"	"gglasso"
[22]	"glamlasso"	"glasso"	"glassoFast"
[25]	"glmmLasso"	"GPCMlasso"	"grplasso"
[28]	"grplassocat"	"hglasso"	"higlasso"
[31]	"ipflasso"	"islasso"	"LassoBacktracking"
[34]	"LassoGEE"	"LassoNet"	"lassopv"
[37]	"lassoshooting"	"LassoSIR"	"lglasso"
[40]	"mglasso"	"MSGLasso"	"MWLasso"
[43]	"nnlasso"	"PabonLasso"	"PACLasso"
[46]	"palasso"	"pcLasso"	"PCLassoReg"
[49]	"ppmlasso"	"prioritylasso"	"sealasso"
[52]	"sglasso"	"slasso"	"smoothedLasso"
[55]	"SSLASSO"	"SummaryLasso"	"Tlasso"
[58]	"vennLasso"	"VSOLassoBag"	

• also see sos package

arm-waving

- optimization: scaling/robustness vs speed
- how do we decide on a 'best' model?

- run everything and compare on a test set? (Do we need another level of nested cross-validation?)
- appropriate metrics: fit quality? fit quality/time or within a time threshold?
- interpretability?
- analogue of no free lunch theorem: "any two optimization algorithms are equivalent when their performance is averaged across all possible problems" (Wolpert and Macready 1997; Giraud-Carrier and Provost 2005)

Wolpert, D. H., and W. G. Macready. 1997. "No Free Lunch Theorems for Optimization." *IEEE Transactions on Evolutionary Computation* 1 (1): 67–82. https://doi.org/10.1109/4235.585893.

Giraud-Carrier, Christophe, and Foster Provost. 2005. "Toward a Justification of Meta-Learning: Is the No Free Lunch Theorem a Show-Stopper?" Proceedings of the ICML-2005 Workshop on Meta-Learning, January.