HUDM 5126 Linear Models and Regression Analysis

Building the Regression Model:
Model Selection

Linear Model Selection

• Ordinary least squares (OLS) regression works well in many real-world applications. Recall, in OLS, we fit a linear model of the form:

$$Y_i = \beta_0 + \beta_1 x_{1i} + \beta_2 x_{2i} + \dots + \beta_{p-1} x_{p-1,i} + \varepsilon_i$$

• by finding estimates for the betas that minimize the sum of squared errors (SSE):

$$\sum_{i=1}^{n} \varepsilon_{i}^{2} = \sum_{i=1}^{n} (Y_{i} - \hat{Y}_{i})^{2} = \sum_{i=1}^{n} (Y_{i} - \hat{\beta}_{0} - \sum_{k=1}^{p} \hat{\beta}_{k} x_{ki})^{2} = SSE$$

• The solution to the problem can be expressed by the normal equations:

$$\hat{\boldsymbol{\beta}} = \left(\mathbf{X}^{\mathsf{T}}\mathbf{X}\right)^{-1}\mathbf{X}^{\mathsf{T}}\mathbf{Y}$$

• where the design matrix *X* is of size *n* by *p* because the first column is all ones for the intercept.

Linear Model Selection

- In some cases, though, there may be good reasons to use other fitting rules than the normal equations. Why?
 - If the sample size *n* is small compared to the number of predictors the least squares fit will have high variability.
 - If the sample size *n* is smaller than the number of predictors, the normal equations do not have a unique solution. For example, 20 variables and 20 cases yields the following output:

```
ALL 20 residuals are 0: no residual degrees of freedom!
Coefficients: (1 not defined because of singularities)
Estimate Std. Error t value Pr(>|t|)
(Intercept) -7.08494
                              NA
                                       NA
                                                 NA
            -0.00754
Х1
                              NΑ
                                       NA
                                                 NA
X2.
             0.03858
                              NA
                                       NA
                                                 NA
             0.19541
Х3
                              NA
                                       NA
                                                 NA
```

- If there are a lot of predictors (i.e., p is large), some of them may be non-informative. That is, they may not be related to the outcome variable. In that case, OLS will estimate small coefficients for the variables, but will not set the coefficients to be exactly zero.
- Thus, we might seek out a method that sets the magnitude of some coefficients to exactly zero. This would result in *feature selection* or *variable selection*.

Three Alternatives to OLS for Linear Models

- 1. Subset selection. Instead of fitting OLS on all possible predictor variables, we first eliminate some by setting their coefficients to zero and then fit OLS on the remaining predictors.
- **2. Regularization/shrinkage.** The model is fit on all *p*-1 predictors, but the magnitudes of the coefficients are shrunk towards zero. These methods work by adding a penalty to the loss function based on the magnitude of the regression coefficients. One method of regularization, called *the lasso*, uses SSE plus a regularization term as the loss function:

SSE: usual error term for OLS regression
$$\left(Y_i - \hat{\beta}_0 - \sum_{k=1}^p \hat{\beta}_k x_{ki}\right)^2 + \lambda \sum_{j=1}^p \left|\beta_j\right|$$
 Shrinkage penalty for the lasso.

3. Dimension reduction. Here we project the p predictors down into an M-dimensional space, where M < p, via linear combinations of variables. We then use the smaller set of M projections as the predictors to be fit by OLS.

Subset Selection

- Best subset selection involves fitting the OLS model for every possible combination of the p predictors and picking the one that is best.
- This is a *brute force* method where we must fit all possible models. Assume *p* predictors. The number of possible combinations (order of the predictors doesn't matter) are
- 0 variables

$$\left(\begin{array}{c} p \\ 0 \end{array}\right) = \frac{p}{0!(p)!}$$

• 1 variable:

$$\begin{pmatrix} p \\ 1 \end{pmatrix} = \frac{p}{1!(p-1)!}$$

• 2 variables:

$$\left(\begin{array}{c} p \\ 2 \end{array}\right) = \frac{p}{2!(p-2)!}$$

• 3 variables:

$$\begin{pmatrix} p \\ 3 \end{pmatrix} = \frac{p}{3!(p-3)!}$$

:

• p-1 variables:

$$\begin{pmatrix} p \\ p-1 \end{pmatrix} = \frac{p}{1!(p-1)!}$$

• p variables:

$$\left(\begin{array}{c} p \\ p \end{array} \right) = \frac{p}{0!(p)!}$$

The sum of all these combinations is 2^p .

- Algorithm for best subset selection:
 - 1. Let M_0 represent the *null model* (i.e., the model with no predictors). This model will only estimate a constant intercept which will represent the sample average.
 - 2. For k = 1, 2, ..., p-1:
 - a) Fit all p choose k models that contain exactly k predictors.
 - b) Select the best-fitting model among all the p-1 choose k models, as measured by SSE, R^2 , or some other measure of model fit, and call it M_k .
 - 3. Pick the single best model from $M_0, ..., M_{p-1}$ using cross-validation prediction error, or some other measure of model fit such as the AIC or BIC.

Model Selection Criteria – All Possible Regressions

P-1 predictors \Rightarrow 2^{P-1} potential models (each variable can be in or out of model)

 R_p^2 or SSE_p criterion (Goal: find p so that $\max(R_p^2)$ or $\min(SSE_p)$ "flattens out"):

$$R_p^2 = \frac{SSR_p}{SSTO} = 1 - \frac{SSE_p}{SSTO}$$
 $p = \# \text{ of parameters in current model}$

 $R_{a,p}^2$ or MSE_p criterion (Goal: find model that maximizes (or close to) $R_{a,p}^2$ and minimizes MSE_p):

$$R_{a,p}^{2} = 1 - \left(\frac{n-1}{n-p}\right) \frac{SSE_{p}}{SSTO} = 1 - \frac{\left(SSE_{p}/(n-p)\right)}{\left(SSTO/(n-1)\right)} = 1 - \frac{MSE_{p}}{\left(SSTO/(n-1)\right)}$$

Mallow's C_p criterion (Goal: find model with smallest p so that $C_p \le p$):

$$C_p = \frac{SSE_p}{MSE(X_1, ..., X_{p-1})} - (n-2p)$$

 AIC_p and SBC_p criteria (Goal: choose model that minimize these values):

$$AIC_p = n \ln \left(SSE_p \right) - n \ln(n) + 2p \qquad SBC_p = n \ln \left(SSE_p \right) - n \ln(n) + \left[\ln(n) \right] p$$

*PRESS*_p criterion (Goal: Small values):

$$PRESS_p = \sum_{i=1}^n \left(Y_i - \hat{Y}_{i(i)} \right)^2$$
 $\hat{Y}_{i(i)} \equiv \text{fitted value for } i^{th} \text{ case when it was not used in fitting model}$

Regression Model Building

- Setting: Possibly a large set of predictor variables (including interactions).
- Goal: Fit a parsimonious model that explains variation in *Y* with a small set of predictors
- Automated Procedures and all possible regressions:
 - Backward Elimination (Top down approach)
 - Forward Selection (Bottom up approach)
 - Stepwise Regression (Combines Forward/Backward)

Forward Selection – Traditional Approach

- Choose a significance level to enter (SLE) the model (e.g. SLE = 0.20, generally 0.05 is too low, causing too few variables to be entered);
- Fit *all* simple regression models;
- Consider the predictor with the highest *t*-statistic (equivalently, the *lowest P*-value):
 - If $P \le$ SLE, keep this variable and fit all two variable models that include this predictor;
 - If P > SLE, stop and keep previous model;
- Continue until no new predictors have P-value \leq SLE
- Note: R uses model based criteria like AIC or BIC instead.

Backward Elimination Traditional Approach

- Select a significance level to stay in the model (e.g. SLS=0.20, generally .05 is too low, causing too many variables to be removed)
- Fit the full model with all possible predictors
- Consider the predictor with lowest *t*-statistic (highest *P*-value).
 - If P > SLS, remove the predictor and fit model without this variable (must re-fit model here because partial regression coefficients change)
 - If $P \leq SLS$, stop and keep current model
- Continue until all predictors have *P*-values below SLS
- Note: R uses model based criteria: AIC, SBC instead

Stepwise Regression – Traditional Approach

- Select SLS and SLE (SLE<SLS)
- Starts like Forward Selection (Bottom up process)
- New variables must have $P \leq SLE$ to enter
- Re-tests all "old variables" that have already been entered, must have $P \le SLS$ to stay in model
- Continues until no new variables can be entered and no old variables need to be removed
- Note: R uses model based criteria: AIC, SBC instead

Model Validation

- When we have a lot of data, we would like to see how well a model fit on one set of data (training sample) compares to one fit on a new set of data (validation sample), and how the training model fits the new data.
- Want data sets to be similar wrt levels of the predictors
- Training set should have at least 6-10 times as many observations than potential predictors
- Models should give similar model fits based on SSE_p , $PRESS_p$, C_p , and MSE_p and regression coefficients
- Mean Square Prediction Error when training model is applied to validation sample: $\sum_{MSPR = \frac{\sum_{i=1}^{n^*} \left(Y_i \hat{Y}_i\right)^2}{n^*}} \hat{Y}_i = b_0^T + b_1^T X_{i1}^V + ... + b_{p-1}^T X_{i,p-1}^V$

Simulating Data for an Example

- Generate data with 10 correlated predictors
- It needs to be multivariate normal, so we need the mvtnorm package.
- We also need the clusterGeneration package for the covariance matrix.
- Set the coefficients for 5 of the predictors to 0, that is, creating non-significant predictors.
- Fit the model
- Check if the 5 unnecessary predictors were eliminated by the various model selection methods

$summary(lm(Y \sim X))$

True coefficient values.

-0.25

0.00

0.08

-0.03

0.00
0.31

0.00

-0.27

-0.16

0.00

0.00

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)	
(Intercept)	-1.086839	1.810540	-0.600	0.549842	
X1	-0.020904	0.048126	-0.434	0.665084	
X2	0.063468	0.043645	1.454	0.149413	
Х3	-0.073515	0.043913	-1.674	0.097621	•
X4	0.009611	0.033618	0.286	0.775629	
X5	0.252679	0.046435	5.442	4.61e-07	***
Х6	-0.011763	0.082320	-0.143	0.886700	
X7	-0.262061	0.070673	-3.708	0.000363	***
X8	-0.120030	0.065162	-1.842	0.068802	•
Х9	-0.051375	0.043551	-1.180	0.241288	
X10	0.002091	0.045053	0.046	0.963087	

Residual standard error: 1.044 on 89 degrees of freedom Multiple R-squared: 0.417, Adjusted R-squared: 0.3515 F-statistic: 6.367 on 10 and 89 DF, p-value: 2.598e-07

The outcome variable *Y* was generated as a linear combination of the *X* variables plus some random normal error.

Five of the *X* variables were assigned to have a coefficient of exactly 0. That is, they are uninformative, noisy variables.

The uninformative variables (i.e., those with no linear relationship with the outcome *Y*) are colored in red.

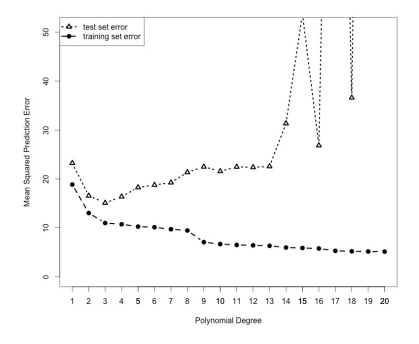
CHOSEN WITH BIC:													
	(Intercept)	X1	X2	Х3	X4	Х5	Х6	х7	X8	Х9	X10	logLikelihood	BIC
0	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	-25.4208998	50.84180
1	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	FALSE	FALSE	FALSE	FALSE	-18.1910797	40.98733
2	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	TRUE	FALSE	FALSE	FALSE	-6.4467824	22.10391
3	TRUE	FALSE	FALSE	FALSE	FALSE	TRUE	FALSE	TRUE	TRUE	FALSE	FALSE	-3.4853247	20.78616
4*	TRUE	FALSE	TRUE	FALSE	FALSE	TRUE	FALSE	TRUE	TRUE	FALSE	FALSE	-1.0415533	20.50379
5	TRUE	FALSE	TRUE	TRUE	FALSE	TRUE	FALSE	TRUE	TRUE	FALSE	FALSE	0.5539902	21.91787
6	TRUE	FALSE	TRUE	TRUE	FALSE	TRUE	FALSE	TRUE	TRUE	TRUE	FALSE	1.2836776	25.06367
7	TRUE	TRUE	TRUE	TRUE	FALSE	TRUE	FALSE	TRUE	TRUE	TRUE	FALSE	1.5073631	29.22147
8	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	FALSE	TRUE	TRUE	TRUE	FALSE	1.5469240	33.74751
9	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	FALSE	1.5584188	38.32969
10	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	TRUE	1.5596289	42.93244
СНО	SEN WITH AIC	:											
	(Intercept)	X1	X2	Х3	X4	X5	Х6	х7	X8	Х9	X10	logLikelihood	AIC
0	TRUE	FALSE											
1			FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	FALSE	-25.4208998	50.841800
	TRUE	_	_	FALSE	_	TRUE	_	_	_		_	-25.4208998 -18.1910797	
2		FALSE	FALSE	_	FALSE	TRUE	_	FALSE	FALSE		FALSE	-18.1910797 -6.4467824	38.382159 16.893565
2	TRUE	FALSE FALSE	FALSE FALSE	FALSE	FALSE FALSE	TRUE TRUE	FALSE	FALSE	FALSE FALSE	FALSE	FALSE FALSE	-18.1910797	38.382159 16.893565
	TRUE TRUE	FALSE FALSE	FALSE FALSE	FALSE FALSE	FALSE FALSE	TRUE TRUE TRUE	FALSE FALSE	FALSE TRUE	FALSE FALSE TRUE	FALSE FALSE	FALSE FALSE	-18.1910797 -6.4467824	38.382159 16.893565 12.970649
3	TRUE TRUE TRUE	FALSE FALSE FALSE	FALSE FALSE	FALSE FALSE FALSE	FALSE FALSE	TRUE TRUE TRUE TRUE	FALSE FALSE	FALSE TRUE TRUE	FALSE FALSE TRUE TRUE	FALSE FALSE FALSE	FALSE FALSE FALSE	-18.1910797 -6.4467824 -3.4853247	38.382159 16.893565 12.970649
3	TRUE TRUE TRUE	FALSE FALSE FALSE	FALSE FALSE FALSE TRUE	FALSE FALSE FALSE TRUE	FALSE FALSE FALSE	TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE	FALSE TRUE TRUE TRUE	FALSE FALSE TRUE TRUE	FALSE FALSE FALSE FALSE	FALSE FALSE FALSE	-18.1910797 -6.4467824 -3.4853247 -1.0415533	38.382159 16.893565 12.970649 10.083107
3 4 5*	TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE	FALSE FALSE FALSE TRUE	FALSE FALSE FALSE TRUE TRUE	FALSE FALSE FALSE FALSE	TRUE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE	FALSE TRUE TRUE TRUE	FALSE FALSE TRUE TRUE	FALSE FALSE FALSE FALSE TRUE	FALSE FALSE FALSE FALSE	-18.1910797 -6.4467824 -3.4853247 -1.0415533 0.5539902	38.382159 16.893565 12.970649 10.083107 8.892020 9.432645
3 4 5* 6	TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE	FALSE FALSE FALSE TRUE TRUE	FALSE FALSE FALSE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE	TRUE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE	FALSE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE	-18.1910797 -6.4467824 -3.4853247 -1.0415533 0.5539902 1.2836776	38.382159 16.893565 12.970649 10.083107 8.892020 9.432645 10.985274
3 4 5* 6 7	TRUE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE TRUE	FALSE FALSE TRUE TRUE TRUE	FALSE FALSE FALSE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE FALSE	TRUE TRUE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE FALSE	FALSE TRUE TRUE TRUE TRUE TRUE	FALSE FALSE TRUE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE TRUE TRUE TRUE	FALSE FALSE FALSE FALSE FALSE FALSE	-18.1910797 -6.4467824 -3.4853247 -1.0415533 0.5539902 1.2836776 1.5073631	38.382159 16.893565 12.970649 10.083107 8.892020 9.432645 10.985274 12.906152

- Best subset selection with the BIC dropped all five non-informative variables but failed to retain X_2 , one of the true linear predictors (albeit with the smallest coefficients.
- Best subset selection with the AIC dropped all five non-informative variables and retained all five true linear predictors.
- Can also choose based on cross-validated prediction accuracy via leaveone-out or *K*-fold.
- What are AIC and BIC?
 - AIC is Akaike's Information Criterion
 - BIC is Bayesian Information Criterion
- The AIC and BIC are measures of relative fit of statistical models.
- Why not simply use mean squared prediction error (MSPE)?

$$MSPE = \frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

AIC and BIC

• As we have seen, the MSPE (mean squared prediction error) measured on the training set is an underestimate of the test set MSPE.



• In particular, the training set MSPE will go down so long as more variables/flexibility are included in the model; however, the test set MSPE will not.

AIC and BIC

The AIC and BIC are both based on -2 times the maximized value of the likelihood (-2LL). k is the number of parameters and LL is the loglikelihood value at the MLE.

$$AIC = 2 * k - 2LL$$
 and $BIC = \ln(n) * k - 2LL$

The likelihood (assuming normally distributed errors) for multiple linear regression is

$$\left(\frac{1}{\sqrt{2\pi}\sigma}\right)^n \exp\left(-\frac{1}{2\sigma^2}|Y-X\beta|^2\right)$$

The log-likelihood is

$$-\frac{n}{2}\log 2\pi - n\log \hat{\sigma} - \frac{1}{2\hat{\sigma}^2}|Y - X\beta|^2$$
This is to residual sum of squares.

squares

Here there is a penalty for more complexity in the model.

Here there is a penalty for magnitude of the LL; smaller is better

AIC and BIC

- As you see from their definitions, AIC and BIC are similarly constructed, with the essential difference being the term multiplied by *k*, the number of parameters estimated.
- As a result (2 vs. log(n)), the BIC tends to penalize model complexity more heavily than the AIC.
- There is debate about which information criteria (there are others aside from AIC and BIC) are "best" and under what circumstances.
- Because the AIC is somewhat more permissive of model complexity than the BIC, it may be preferred for *prediction*.
- When the intent is to create a model for *explanation*, BIC may be preferred because it will tend to produce a simpler (i.e., more easily interpretable) model.

Best Subset and Computational Efficiency

- Best subset selection is computationally demanding.
- For p = 30, for example, best subset will require fitting $2^{30} = 1,073,741,824$ models.
- Whereas, forward selection (described next) will require fitting 1 + 1 + 2 + 3 + ... + 29 + 30 = p(p+1)/2 + 1 = 451 models.
- That said, the ease of fitting Gaussian linear models make even large (i.e., 30+ covariates) problems tractable within a few seconds. This is possible in part due to advances in computational algorithms over the last decades.
- The problem of computational efficiency is quickly made worse, however, by using other models that require iterative numerical methods for solving.
- For example, with Gaussian linear models (which can be solved analytically), a best subset search with 11 predictors takes less than a second.
- By contrast, with a logistic model (solved using Newton's method), 11 predictors takes nearly 2 minutes, and adding each subsequent predictor causes the time to more than double.

Data from the early childhood longitudinal study.

The outcome is 5th grade math score (C6R4MSCL).

The exposure is an indicator that is 1 if student received special education services at or before 3rd grade; 0 otherwise.

Interested in the effect of exposure to special education on math score, controlling for other variables.

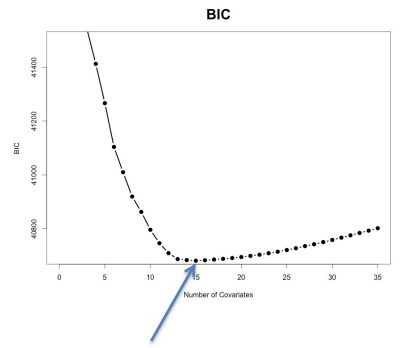
Variable Name	Variable Name Description of Variable		d	r		
DEMOGRAPHI	DEMOGRAPHIC					
GENDER	Male	0, 1	0.38	0.88		
WKWHITE	White	0, 1	0.17	0.79		
WKSESL	Socioeconomic Status	[-4.8, 2.8]	-0.29	0.89		
ACADEMIC						
RIRT	Kindergarten Reading Score	[23.17, 139.36]	-0.65	0.53		
MIRT	Kindergarten Math Score	[11.9, 99.0]	-0.71	0.77		
S2KPUPRI	Public School	0, 1	0.44	0.25		
P1EXPECT	Parental Expectations	Integers 1–6	-0.32	1.22		
P1FIRKDG	First-Time Kindergartener	0, 1	-0.41	3.26		
P1AGEENT	Child's Age at K Entry (Months)	[54, 79]	0.08	1.08		
apprchT1	Approaches to Learning Rating	Integers 1–4	-0.70	1.20		
P1HSEVER	Attended Head Start	0, 1	0.19	1.42		
chg14	Ever Changed Schools	0, 1	0.02	1.09		
SCHOOL COMP	POSITION					
avg_RIRT	Reading IRT	[27.9, 80.0]	-0.23	0.79		
avg_MIRT	Math IRT	[16.1, 66.1]	-0.18	0.82		
avg_SES	SES	[-2.2, 2.5]	-0.16	0.88		
avg_apprchT1	Approaches to Learning	[1.5, 4.0]	-0.14	0.80		
S2KMINOR	Percent Minority Students	Integers 1–5	-0.20	0.77		
FAMILY CONTI	$\mathbf{E}\mathbf{X}\mathbf{T}$					
P1FSTAMP	Received Food Stamps	0, 1	0.12	1.26		
ONEPARENT	One-Parent Family	0, 1	0.13	1.22		
STEPPARENT	Stepparent Family	0, 1	0.05	1.19		
P1NUMSIB	Number of Siblings	[0, 10]	0.16	1.17		
P1HMAFB	Mother's Age at First Birth	Years [12, 45]	-0.26	1.00		
WKCAREPK	Nonparental Pre-K Child Care	0, 1	-0.07	1.14		
HEALTH						
P1EARLY	Number of Days Premature	[0, 112]	0.19	2.05		
wt_ounces	Birth Weight (Ounces)	[17, 214]	-0.11	1.24		
C1FMOTOR	Fine Motor Skills	Integers 0–9	-0.63	1.27		
C1GMOTOR	Gross Motor Skills	Integers 0–8	-0.43	1.54		
PARENT RATIN	NG OF CHILD					
P1HSCALE	Overall Health	Integers 1–5	0.12	1.17		
P1SADLON	Sad/Lonely	Integers 1–4	0.10	1.32		
P1IMPULS	Impulsive	Integers 1–4	0.41	1.55		
P1ATTENI	Attentive	Integers 1–4	0.72	1.45		
P1SOLVE	Problem Solving	Integers 1–4	0.68	1.55		
PSPRONOU	Verbal Communication	Integers 1–4	0.86	1.51		
P1DISABL	Child has Disability	0, 1	0.82	2.38		
OUTCOME VAI		•				
C6R4MSCL	Fifth Grade Math Score	[50.9, 170.7]	-0.77	1.40		

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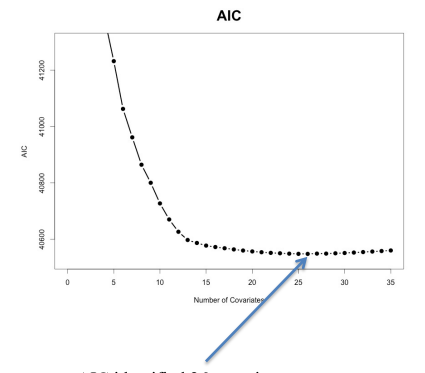
ECLSK Example – Best Subset Selection

```
### Outcome is C6R4MSCL (5th grade math score)
### 36 predictors examined
bs3 <- bestglm(Xy = eclsk1, family = gaussian, IC = "BIC")
bs4 <- bestglm(Xy = eclsk1, family = gaussian, IC = "AIC")</pre>
```

Best subset selection works here because the model is Gaussian. If we were to try a logistic model for binomial data with 36 predictors we would get an error message.



BIC identified 15 covariates as the optimal number.



AIC identified 26 covariates as the optimal number.

ECLSK Example - Results

BIC Results

Estimate S	Std. Error	t value Pr	(> t)		
(Intercept)	92.52027	3.30622	27.984	< 2e-16	***
GENDER	6.20534	0.38073	16.299	< 2e-16	***
WKWHITE	3.19381	0.46598	6.854	7.76e-12	***
WKSESL	2.14455	0.34933	6.139	8.73e-10	***
MIRT	1.17759	0.02432	48.427	< 2e-16	***
S2KPUPRI	5.52234	0.49119	11.243	< 2e-16	***
P1FIRKDG	12.04908	1.07558	11.202	< 2e-16	***
P1AGEENT	-0.72637	0.04860	-14.946	< 2e-16	***
apprchT1	2.68547	0.32903	8.162	3.85e-16	***
P1HSEVER	-3.60635	0.61973	-5.819	6.16e-09	***
ONEPARENT	-1.90534	0.52685	-3.616	0.000301	***
P1HMAFB	0.21746	0.04114	5.286	1.29e-07	***
C1FMOTOR	1.67406	0.10746	15.579	< 2e-16	***
P1SOLVE	-1.16148	0.34532	-3.364	0.000773	***
avg_SES	3.14823	0.53396	5.896	3.89e-09	***
F5SPECS	-7.17163	0.81787	-8.769	< 2e-16	***

Special Ed indicator is significant in both models and the treatment effect estimate is about the same (-7.17 vs. -7.14).

AIC Results

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 96.88775
                         4.24281 22.836 < 2e-16 ***
GENDER
                         0.38889 15.824 < 2e-16 ***
              6.15373
WKWHITE
              2.27241
                         0.55271
                                    4.111 3.98e-05 ***
WKSESL
              1.92348
                         0.35772
                                    5.377 7.81e-08 ***
MIRT
              1.16513
                         0.02451 47.540
                                          < 2e-16
S2KPUPRI
              5.57930
                         0.49124 11.358
                                         < 2e-16 ***
P1EXPECT
              0.41871
                         0.19160
                                    2.185 0.028893 *
             12.05349
P1FIRKDG
                         1.07801 11.181
                                          < 2e-16
             -0.72955
                         0.04909 - 14.861
P1AGEENT
                                           < 2e-16
              3.03527
apprchT1
                         0.38177
                                   7.951 2.14e-15 ***
P1HSEVER
             -2.98581
                         0.64604 -4.622 3.87e-06 ***
             -1.58801
P1FSTAMP
                         0.67575 -2.350 0.018799 *
             -0.49038
                         0.16931
                                  -2.896 0.003787 **
S2KMINOR
                         0.55536 -2.203 0.027595 *
             -1.22370
ONEPARENT
              0.21800
P1HMAFB
                         0.04131
                                    5.278 1.35e-07 ***
WKCAREPK
             -1.09543
                         0.52133 -2.101 0.035657 *
P1EARLY
              0.04088
                                    2.264 0.023633 *
                         0.01806
wt ounces
              0.03510
                         0.01065
                                    3.295 0.000987 ***
C1FMOTOR
              1.68800
                         0.10862 15.541 < 2e-16 ***
             -0.20252
C1GMOTOR
                         0.10899
                                  -1.858 0.063187 .
P1ATTENI
             -0.56043
                         0.33260
                                  -1.685 0.092030 .
             -0.82948
P1SOLVE
                         0.36751
                                  -2.257 0.024037 *
             -0.64512
                         0.32960 -1.957 0.050353 .
P1PRONOU
P1DISABL
              0.85005
                         0.58371
                                   1.456 0.145356
avg SES
              2.92792
                         0.55637
                                    5.263 1.46e-07 ***
                         0.70086 -2.501 0.012410 *
avg apprchT1 -1.75277
             -7.14141
                                  -8.605 < 2e-16 ***
F5SPECS
                         0.82992
```

Forward Stepwise Selection

- Forward and backward stepwise approaches to model selection approximate the best subset solution by working with a (much!) more restricted set of models.
- Forward stepwise selection algorithm:
 - 1. Let M_0 denote the null model (i.e., no predictors).
 - 2. For k = 0, ..., p 1:
 - a) Consider all p k models that increase the predictors in M_k with one additional parameter.
 - b) Choose the best among the p k models and call it M_{k+1} . Note that "best" is defined in terms of smallest SSE.
 - 3. Select a single best model from $\{M0, ..., M_k\}$ via CV, AIC, BIC, etc.
- In total, this will involve fitting 1 null model and p k models in the kth iteration, for k = 0, 1, ..., p 1.
- This is in contrast to 2^p for best subset selection.

ECLSK Example – Forward Stepwise Selection

• Several packages in R can run forward selection. The "stepAIC" function in package MASS is what we will use.

```
library (MASS)
min.model <- lm(C6R4MSCL ~ 1, data = eclsk1)</pre>
max.model <- lm(C6R4MSCL ~ ., data = eclsk1)</pre>
scp <- list(lower = min.model, upper = max.model)</pre>
fwd <- step(min.model,</pre>
             direction = 'forward',
             scope = scp)
fwd$coefficients
 (Intercept)
                                  WKSESL
                                              C1FMOTOR
                      MIRT
                                                              GENDER
                                                                          P1AGEENT
 96.88775211
                1.16513347
                              1.92348365
                                            1.68800475
                                                          6.15372996
                                                                       -0.72954903
    P1FIRKDG
                  apprchT1
                                 WKWHITE
                                              S2KPUPRI
                                                             F5SPECS
                                                                           avg SES
 12.05348816
                3.03526705
                              2.27240622
                                            5.57929998
                                                         -7.14141261
                                                                        2.92792434
    P1HSEVER
                   P1HMAFB
                               ONEPARENT
                                               P1SOLVE
                                                            S2KMINOR
                                                                         wt ounces
 -2.98581062
                0.21799665
                             -1.22369547
                                           -0.82947802
                                                         -0.49037825
                                                                        0.03510179
avg apprchT1
                                                            WKCAREPK
                  P1EXPECT
                                 P1EARLY
                                              P1FSTAMP
                                                                          P1PRONOU
 -1.75276672
                0.41870822
                              0.04087811
                                           -1.58800536
                                                         -1.09542548
                                                                       -0.64512367
    C1GMOTOR
                  P1ATTENT
                                P1DTSABL
 -0.20252002
               -0.56042548
                              0.85004828
```

Note that AIC via best subset also identified the same 26 covariates.

ECLSK Example – Forward Stepwise Selection

• To use the BIC we modify the argument k (default is 2 for AIC).

```
fwd2 <- stepAIC(min.model,</pre>
                direction = 'forward',
                scope = scp,
                k = log(nrow(eclsk1)))
fwd2$coefficients
(Intercept)
                   MIRT
                             WKSESL
                                        C1FMOTOR
                                                      GENDER
                                                                P1AGEENT
                                                                             P1FIRKDG
92.5202705
              1.1775938
                          2.1445523
                                      1.6740648
                                                   6.2053449 -0.7263671 12.0490844
  apprchT1
                WKWHITE
                           S2KPUPRI
                                         F5SPECS
                                                     avg SES
                                                                P1HSEVER
                                                                              P1HMAFB
  2.6854711
                          5.5223400 -7.1716267
                                                   3.1482347 -3.6063459
                                                                            0.2174640
              3.1938133
  ONEPARENT
                P1SOLVE
-1.9053391
             -1.1614815
```

- Note that BIC via best subset also identified the same 15 covariates.
- The two methods will not always agree.

Backward Stepwise Selection

- Backward stepwise selection is also a much more efficient alternative to best subset selection.
- Whereas forward stepwise selection begins with a null model and moves forward one predictor at a time, backward stepwise selection begins with the full model (i.e., containing all *p* predictors) and then removes the least useful predictors one at a time.
- Backward stepwise selection algorithm (ISLR, p. 209):
 - 1. Let M_p denote the *full* model, containing all p predictors.
 - 2. For k = p, p 1, p 2, ..., 1:
 - a) Consider all k models that contain all but one of the predictors in M_k , for a total of k-1 predictors.
 - b) Select the best of the k models and label it M_{k-1} . At this stage, *best* is defined by smallest residual sum of squares.
 - 3. Select the single best model from M_0, \ldots, M_p via cross-validation prediction error, AIC, BIC, etc.

Backward Subset Selection

```
bwd <- stepAIC(max.model,</pre>
                                                  Note we now use
               direction = 'backward',
                                                  max.model here instead of
               scope = scp)
                                                  min.model.
bwd$coefficients
(Intercept)
                                             WKSESL
                                                                     S2KPUPRI
                  GENDER
                              WKWHITE
                                                            MIRT
 96.16423579
               6.15303750
                            2.34469806
                                         1.92016994
                                                                    5.48863902
                                                       1.17493417
                                           apprchT1
    P1EXPECT
                 P1FIRKDG
                              P1AGEENT
                                                         P1HSEVER
                                                                      P1FSTAMP
                                         2.94808419 -2.96083960 -1.56001714
  0.42597045
              11.96839592
                           -0.71978192
    S2KMINOR
                ONEPARENT
                               P1HMAFB
                                            WKCAREPK
                                                          P1EARLY
                                                                     wt ounces
 -0.56437253
              -1.23950171 0.21721213 -1.06344788
                                                       0.04315746
                                                                    0.03569199
    C1FMOTOR
                 C1GMOTOR
                              P1ATTENT
                                             P1SOLVE
                                                         P1PRONOU
                                                                      avg RIRT
  1.68755103 - 0.20388550 - 0.54930933 - 0.80559386
                                                     -0.53292879
                                                                    0.08605165
    avg MIRT
                  avg SES avg apprchT1
                                             F5SPECS
 -0.12525895
               3.04132473 -1.57149298 -6.96682986
```

- avg_MIRT and avg_RIRT were selected by backward subset selection, but not by best or forward.
- P1DISABL was selected by best and forward but not by backward.

Limitations of Stepwise Selection

- Limitation 1: Inflation of Type I error rate when testing the significance of predictors.
- Stepwise routines fit many models along the way to the final formulation, testing many (or all in the case of best subset selection) possible combinations of covariates.
- Because so many combinations are tested, some are bound to be significant by chance.
- Thus, hypothesis testing of regression coefficients after running a stepwise selection routine will typically show that nearly every variable retained is a "significant" predictor of the outcome.
- The problem here lies in the fact that so many models were fit. It is, therefore, not appropriate to interpret the statistical significance of regression coefficients selected by stepwise selection routines at face value.
- For the purpose of making good *predictions*, on the other hand, stepwise selection methods are very useful because they can eliminate non-informative variables.

Limitations of Stepwise Selection

• We can demonstrate the potential to make Type I errors through simulation. Generate Y *completely unrelated* to X1 through X20.

```
### Generate random noise and use stepwise approach
noise \leftarrow function (N = 400, p = 20) {
  X <- rmvnorm(N, sigma = diag(p))</pre>
  Y \leftarrow rnorm(N, sd = 4)
  df <- data.frame(cbind(X,Y))</pre>
  names(df) <- c(paste0("X", 1:p), "Y")</pre>
  df
set.seed(1355)
df6 <- noise()
summary(lm(Y \sim X, df6))
bestglm(Xy = df6, family = gaussian, IC = "AIC")
               Estimate Std. Error t value
                                                  Pr(>|t|)
(Intercept) -0.3564621 0.2014851 -1.769174 0.077638465
Х7
             -0.3251008 0.2075722 -1.566206 0.118103234
X11
             -0.6132137 0.2182780 -2.809324 0.005211912
            -0.3750686 0.2065032 -1.816285 0.070086522
X14
            -0.3229027 0.2140893 -1.508262 0.132288968
X16
X20
             -0.3001960 0.1912377 -1.569753 0.117275226
```

Best subset selection using the AIC identifies X11 as a significant predictor of Y even though Y and Xs are independent.

Using the BIC also identifies X11 as significant.

Furthermore, the linear regression of Y on all Xs yielded no significant relationships at alpha = 0.05.

Limitations of Stepwise Selection

- Limitation 2: functional form assumptions are strong.
- Linear and generalized linear stepwise procedures assume the functional form of the model is a subset of the most complex model you specify.
- For example, if you do not specify any squared terms or interactions in the "upper" model, the assumption is that all predictors are linearly related to the outcome.
- As a result the "best" model will be selected within the linear models.