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cv: An R Package for Cross-Validation of Regression Models

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Abstract

The abstract of the article.

Keywords: cross-validation, regression analysis, model selection, R.

1. Cross-validation

Cross-validation (CV) is an essentially simple and intuitively reasonable approach to estimating the predictive accuracy of regression models. CV is developed in many standard sources on regression modeling and "machine learning"—we particularly recommend James, Witten, Hastie, and Tibshirani (2021, Secs. 5.1, 5.3)—and so we will describe the method only briefly here before taking up computational issues and some examples. See Arlot and Celisse (2010) for a wide-ranging, if technical, survey of cross-validation and related methods that emphasizes the statistical properties of CV.

Validating research by replication on independently collected data is a common scientific norm. Emulating this process in a single study by data-division is less common: The data are randomly divided into two, possibly equal-size, parts; the first part is used to develop and fit a statistical model; and then the second part is used to assess the adequacy of the model fit to the first part of the data. Data-division, however, suffers from two problems: (1) Dividing the data decreases the sample size and thus increases sampling error; and (2), even more disconcertingly, particularly in smaller samples, the results can vary substantially based on the random division of the data: See Harrell (2015, Sec. 5.3) for this and other remarks about data-division and cross-validation.

Cross-validation speaks to both of these issues. In CV, the data are randomly divided as equally as possible into several, say k, parts, called "folds." The statistical model is fit k times, leaving each fold out in turn. Each fitted model is then used to predict the response

variable for the cases in the omitted fold. A CV criterion or "cost" measure, such as the mean-squared error ("MSE") of prediction, is then computed using these predicted values. In the extreme k=n, the number of cases in the data, thus omitting individual cases and refitting the model n times—a procedure termed "leave-one-out (LOO) cross-validation."

Because the n models are each fit to n-1 cases, LOO CV produces a nearly unbiased estimate of prediction error. The n regression models are highly statistical dependent, however, based as they are on nearly the same data, and so the resulting estimate of prediction error has relatively large variance. In contrast, estimated prediction error for k-fold CV with k=5 or 10 (commonly employed choices) are somewhat biased but have smaller variance. It is also possible to correct k-fold CV for bias (see below).

2. Examples

2.1. Polynomial regression for the Auto data

The data for this example are drawn from the **ISLR2** package for R, associated with James et al. (2021). The presentation here is close (though not identical) to that in the original source (James et al. 2021, Secs. 5.1, 5.3), and it demonstrates the use of the cv() function in the cv package.¹

The Auto dataset contains information about 392 cars:

```
R> data("Auto", package="ISLR2")
R> head(Auto)
```

	${\tt mpg}$	cylinders	displacement	horsepower	weight	acceleration	year	origin
1	18	8	307	130	3504	12.0	70	1
2	15	8	350	165	3693	11.5	70	1
3	18	8	318	150	3436	11.0	70	1
4	16	8	304	150	3433	12.0	70	1
5	17	8	302	140	3449	10.5	70	1
6	15	8	429	198	4341	10.0	70	1
			name					

```
1 chevrolet chevelle malibu
2 buick skylark 320
3 plymouth satellite
4 amc rebel sst
5 ford torino
6 ford galaxie 500
```

R> dim(Auto)

[1] 392 9

¹James *et al.* (2021) use the cv.glm() function in the **boot** package (Canty and Ripley 2022; Davison and Hinkley 1997). Despite its name, cv.glm() is an independent function and not a method of a cv() generic function.

With the exception of origin (which we don't use here), these variables are largely self-explanatory, except possibly for units of measurement: for details see help("Auto", package="ISLR2"). We'll focus here on the relationship of mpg (miles per gallon) to horsepower, as displayed in

```
R> plot(mpg ~ horsepower, data=Auto)
```

the following scatterplot:

The relationship between the two variables is monotone, decreasing, and nonlinear. Following James *et al.* (2021), we'll consider approximating the relationship by a polynomial regression, with the degree of the polynomial p ranging from 1 (a linear regression) to $10.^2$ Polynomial fits for p = 1 to 5 are shown in the following figure:

The linear fit is clearly inappropriate; the fits for p = 2 (quadratic) through 4 are very similar; and the fit for p = 5 may over-fit the data by chasing one or two relatively high mpg values at the right (but see the CV results reported below).

The following graph shows two measures of estimated (squared) error as a function of polynomial-regression degree: The mean-squared error ("MSE"), defined as $\mathsf{MSE} = \frac{1}{n} \sum_{i=1}^n (y_i - \widehat{y}_i)^2$, and the usual residual variance, defined as $\widehat{\sigma}^2 = \frac{1}{n-p-1} \sum_{i=1}^n (y_i - \widehat{y}_i)^2$. The former necessarily declines with p (or, more strictly, can't increase with p), while the latter gets slightly larger for the largest values of p, with the "best" value, by a small margin, for p = 7.

R> library("cv") # for mse() and other functions

Loading required package: doParallel

Loading required package: foreach

Loading required package: iterators

Loading required package: parallel

²Although it serves to illustrate the use of CV, a polynomial is probably not the best choice here. Consider, for example the scatterplot for log-transformed mpg and horsepower, produced by plot(mpg ~ horsepower, data=Auto, log="xy") (execution of which is left to the reader).

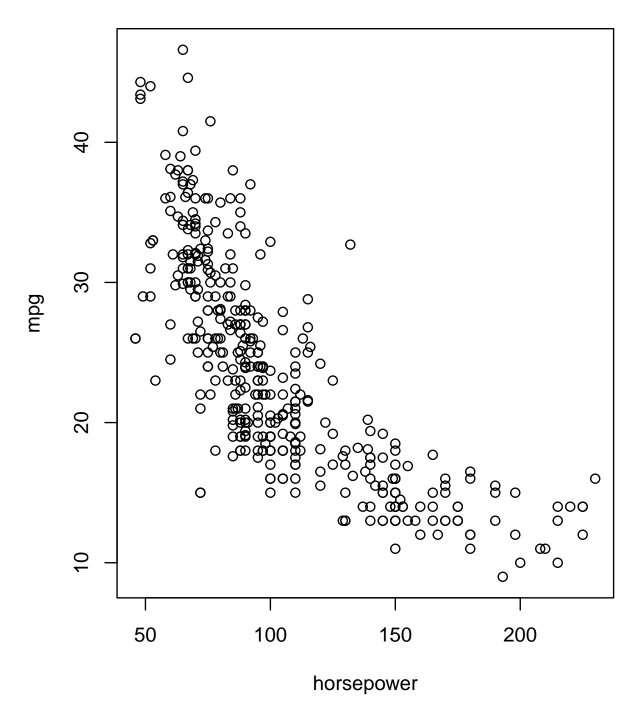


Figure 1: 'mpg' vs 'horsepower' for the 'Auto' data

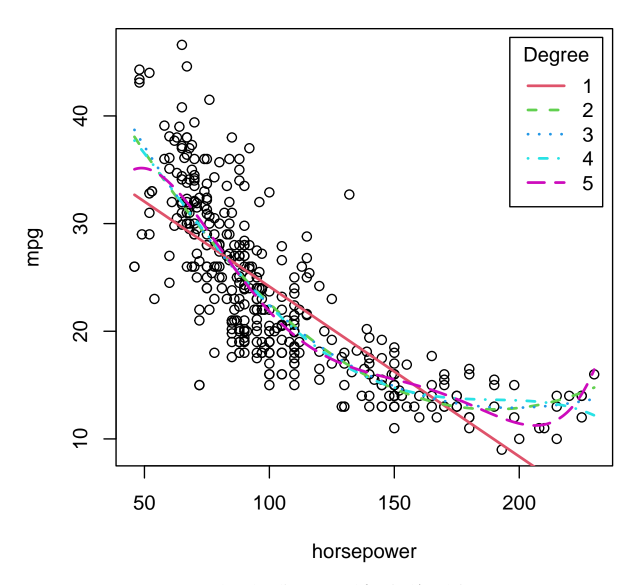


Figure 2: 'mpg' vs 'horsepower' for the 'Auto' data

```
R> var <- mse <- numeric(10)</pre>
R> for (p in 1:10){
    m <- lm(mpg ~ poly(horsepower, p), data=Auto)</pre>
    mse[p] <- mse(Auto$mpg, fitted(m))</pre>
    var[p] <- summary(m)$sigma^2</pre>
+ }
R>
R > plot(c(1, 10), range(mse, var), type="n",
       xlab="Degree of polynomial, p",
       ylab="Estimated Squared Error")
R> lines(1:10, mse, lwd=2, lty=1, col=2, pch=16, type="b")
R> lines(1:10, var, lwd=2, lty=2, col=3, pch=17, type="b")
R> legend("topright", inset=0.02,
         legend=c(expression(hat(sigma)^2), "MSE"),
         lwd=2, lty=2:1, col=3:2, pch=17:16)
The code for this graph uses the mse() function from the cv package to compute the MSE
for each fit.
Using cv()
The generic cv() function has an "lm" method, which by default performs k = 10-fold CV:
R> m.auto <- lm(mpg ~ poly(horsepower, 2), data=Auto)</pre>
R> summary(m.auto)
Call:
lm(formula = mpg ~ poly(horsepower, 2), data = Auto)
Residuals:
               10
                    Median
                                  30
     Min
                                           Max
-14.7135 -2.5943 -0.0859
                              2.2868 15.8961
Coefficients:
                       Estimate Std. Error t value Pr(>|t|)
(Intercept)
                        23.4459
                                    0.2209 106.13
                                                      <2e-16
poly(horsepower, 2)1 -120.1377
                                    4.3739 -27.47
                                                      <2e-16
poly(horsepower, 2)2
                        44.0895
                                    4.3739
                                              10.08
                                                      <2e-16
Residual standard error: 4.374 on 389 degrees of freedom
Multiple R-squared: 0.6876,
                                 Adjusted R-squared: 0.686
              428 on 2 and 389 DF, p-value: < 2.2e-16
F-statistic:
```

R RNG seed set to 534539

R> cv(m.auto)

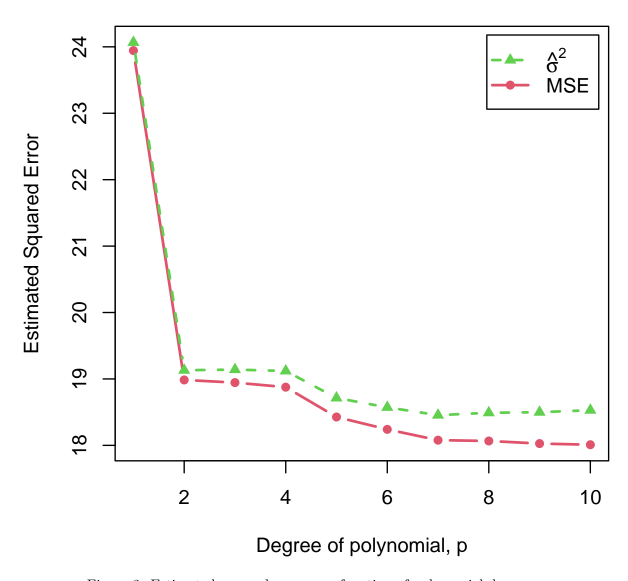


Figure 3: Estimated squared error as a function of polynomial degree, p

10-Fold Cross Validation
method: Woodbury
criterion: mse
cross-validation criterion = 19.22944
bias-adjusted cross-validation criterion = 19.21651
full-sample criterion = 18.98477

The "lm" method by default uses mse() as the CV criterion and the Woodbury matrix identity to update the regression with each fold deleted without having literally to refit the model. Computational details are discussed in the final section of this vignette. The function reports the CV estimate of MSE, a biased-adjusted estimate of the MSE (the bias adjustment is explained in the final section), and the MSE is also computed for the original, full-sample regression. Because the division of the data into 10 folds is random, cv() explicitly (randomly) generates and saves a seed for R's pseudo-random number generator, to make the results replicable. The user can also specify the seed directly via the seed argument to cv().

To perform LOO CV, we can set the k argument to cv() to the number of cases in the data, here k=392, or, more conveniently, to k="loo" or k="n":

R> cv(m.auto, k="loo")
n-Fold Cross Validation
method: hatvalues
criterion: mse
cross-validation criterion = 19.24821

For LOO CV of a linear model, cv() by default uses the hatvalues from the model fit to the full data for the LOO updates, and reports only the CV estimate of MSE. Alternative methods are to use the Woodbury matrix identity or the "naive" approach of literally refitting the model with each case omitted. All three methods produce exact results for a linear model (within the precision of floating-point computations):

R> cv(m.auto, k="loo", method="naive")
n-Fold Cross Validation
method: naive
criterion: mse
cross-validation criterion = 19.24821
bias-adjusted cross-validation criterion = 19.24787
full-sample criterion = 18.98477

R> cv(m.auto, k="loo", method="Woodbury")
n-Fold Cross Validation
method: Woodbury
criterion: mse
cross-validation criterion = 19.24821
bias-adjusted cross-validation criterion = 19.24787
full-sample criterion = 18.98477

The "naive" and "Woodbury" methods also return the bias-adjusted estimate of MSE and the full-sample MSE, but bias isn't an issue for LOO CV.

This is a small regression problem and all three computational approaches are essentially instantaneous, but it is still of interest to investigate their relative speed. In this comparison, we include the <code>cv.glm()</code> function from the **boot** package, which takes the naive approach, and for which we have to fit the linear model as an equivalent Gaussian GLM. We use the <code>microbenchmark()</code> function from the package of the same name for the timings (Mersmann 2023):

```
R> m.auto.glm <- glm(mpg ~ poly(horsepower, 2), data=Auto)</pre>
R> boot::cv.glm(Auto, m.auto.glm)$delta
[1] 19.24821 19.24787
R> microbenchmark::microbenchmark(
    hatvalues = cv(m.auto, k="loo"),
    Woodbury = cv(m.auto, k="loo", method="Woodbury"),
    naive = cv(m.auto, k="loo", method="naive"),
    cv.glm = boot::cv.glm(Auto, m.auto.glm),
    times=10
+ )
Warning in microbenchmark::microbenchmark(hatvalues = cv(m.auto, k = "loo"), :
less accurate nanosecond times to avoid potential integer overflows
Unit: microseconds
                  min
      expr
                              lq
                                                median
                                       mean
                                                                uq
                                                                          max
 hatvalues
              977.973
                          993.43
                                   1143.433
                                              1173.113
                                                          1202.981
                                                                     1296.256
  Woodbury
             9932.004
                       10178.21 11461.993
                                             10363.201
                                                        10865.943
     naive 215384.152 217809.02 224546.766 219116.505 221670.231 275518.278
    cv.glm 384357.616 388159.59 409612.546 390652.100 441741.913 443226.605
 neval cld
    10 a
    10 a
    10 b
    10
```

On our computer, using the hat values is about an order of magnitude faster than employing Woodbury matrix updates, and more than two orders of magnitude faster than refitting the $\operatorname{model.}^3$

Comparing competing models

³Out of impatience, we asked microbenchmark() to execute each command only 10 times rather than the default 100. With the exception of the last columns, the output is self-explanatory. The last column shows which methods have average timings that are statistically distinguishable. Because of the small number of repetitions (i.e., 10), the "hatvalues" and "Woodbury" methods aren't distinguishable, but the difference between these methods persists when we perform more repetitions—we invite the reader to redo this computation with the default times=100 repetitions.

The cv() function also has a method that can be applied to a list of regression models for the same data, composed using the models() function. For k-fold CV, the same folds are used for the competing models, which reduces random error in their comparison. This result can also be obtained by specifying a common seed for R's random-number generator while applying cv() separately to each model, but employing a list of models is more convenient for both k-fold and LOO CV (where there is no random component to the composition of the n folds).

We illustrate with the polynomial regression models of varying degree for the Auto data (discussed previously), beginning by fitting and saving the 10 models:

```
R> for (p in 1:10){
    assign(paste0("m.", p),
           lm(mpg ~ poly(horsepower, p), data=Auto))
+ }
R> objects(pattern="m\\.[0-9]")
 [1] "m.1" "m.10" "m.2"
                         "m.3"
                                  "m.4"
                                         "m.5"
                                                "m.6"
                                                        "m.7"
                                                               "m.8"
                                                                      "m.9"
R> summary(m.2) # for example, the quadratic fit
Call:
lm(formula = mpg ~ poly(horsepower, p), data = Auto)
Residuals:
     Min
               1Q
                    Median
                                  3Q
                                          Max
-14.7135 -2.5943 -0.0859
                              2.2868
                                      15.8961
Coefficients:
                      Estimate Std. Error t value Pr(>|t|)
(Intercept)
                        23.4459
                                    0.2209
                                            106.13
                                                      <2e-16
poly(horsepower, p)1 -120.1377
                                            -27.47
                                                      <2e-16
                                    4.3739
poly(horsepower, p)2
                        44.0895
                                    4.3739
                                             10.08
                                                      <2e-16
Residual standard error: 4.374 on 389 degrees of freedom
Multiple R-squared: 0.6876,
                                 Adjusted R-squared: 0.686
               428 on 2 and 389 DF, p-value: < 2.2e-16
F-statistic:
We then apply cv() to the list of 10 models (the data argument is required):
R> # 10-fold CV
R> cv.auto.10 <- cv(models(m.1, m.2, m.3, m.4, m.5,
                       m.6, m.7, m.8, m.9, m.10),
```

+ data=Auto, seed=2120)
R> cv.auto.10[1:2] # for the linear and quadratic models

```
Model model.1:
10-Fold Cross Validation
method: Woodbury
cross-validation criterion = 24.24642
bias-adjusted cross-validation criterion = 24.23039
full-sample criterion = 23.94366
Model model.2:
10-Fold Cross Validation
method: Woodbury
cross-validation criterion = 19.34601
bias-adjusted cross-validation criterion = 19.32699
full-sample criterion = 18.98477
R> # LOO CV
R> cv.auto.loo <- cv(models(m.1, m.2, m.3, m.4, m.5,
+
                          m.6, m.7, m.8, m.9, m.10),
                   data=Auto, k="loo")
R> cv.auto.loo[1:2] # linear and quadratic models
Model model.1:
n-Fold Cross Validation
method: hatvalues
cross-validation criterion = 24.23151
Model model.2:
n-Fold Cross Validation
method: hatvalues
cross-validation criterion = 19.24821
```

Because we didn't supply names for the models in the calls to the models() function, the names model.1, model.2, etc., are generated by the function.

Finally, we extract and graph the adjusted MSEs for 10-fold CV and the MSEs for LOO CV:

Alternatively, we can use the plot() method for "cvModList" objects to compare the models, though with separate graphs for 10-fold and LOO CV:

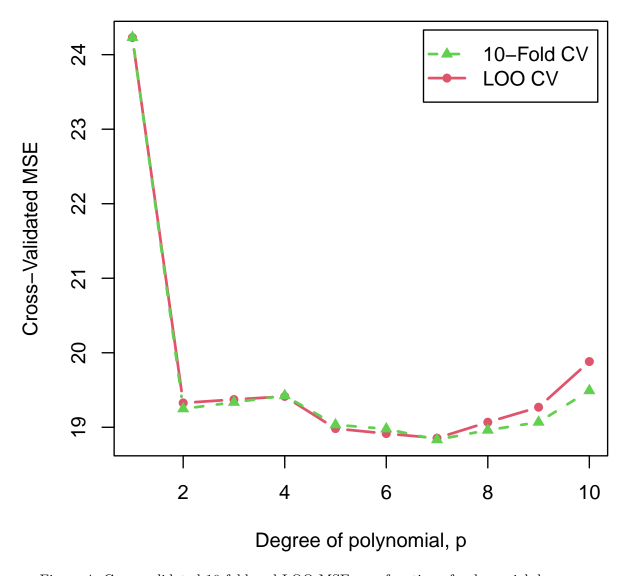


Figure 4: Cross-validated 10-fold and LOO MSE as a function of polynomial degree, p

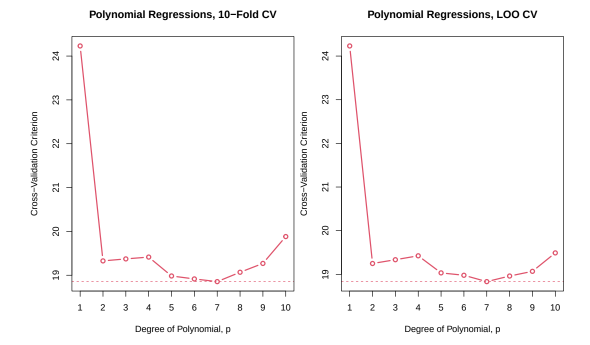


Figure 5: Cross-validated 10-fold and LOO MSE as a function of polynomial degree, p

```
R> plot(cv.auto.10, main="Polynomial Regressions, 10-Fold CV",
+ axis.args=list(labels=1:10), xlab="Degree of Polynomial, p")
R> plot(cv.auto.loo, main="Polynomial Regressions, LOO CV",
+ axis.args=list(labels=1:10), xlab="Degree of Polynomial, p")
```

In this example, 10-fold and LOO CV produce generally similar results, and also results that are similar to those produced by the estimated error variance $\hat{\sigma}^2$ for each model, reported above (except for the highest-degree polynomials, where the CV results more clearly suggest over-fitting).

2.2. Logistic regression for the Mroz data

The Mroz data set from the **carData** package (associated with Fox and Weisberg 2019) has been used by several authors to illustrate binary logistic regression; see, in particular Fox and Weisberg (2019). The data were originally drawn from the U.S. Panel Study of Income Dynamics and pertain to married women. Here are a few cases in the data set:

```
R> data("Mroz", package="carData")
R> head(Mroz, 3)

Ifp k5 k618 age wc hc lwg inc
1 yes 1     0 32 no no 1.2101647 10.91
2 yes 0     2 30 no no 0.3285041 19.50
3 yes 1     3 35 no no 1.5141279 12.04
R> tail(Mroz, 3)
```

```
lfp k5 k618 age wc hc
                                lwg
                                       inc
                                   9.952
751 no
              0 43 no no 0.8881401
                60 no no 1.2249736 24.984
752 no
753
              3 39 no no 0.8532125 28.363
```

The response variable in the logistic regression is 1fp, labor-force participation, a factor coded "yes" or "no". The remaining variables are predictors:

- k5, number of children 5 years old of younger in the woman's household;
- k618, number of children between 6 and 18 years old;
- age, in years;
- wc, wife's college attendance, "yes" or "no";
- hc, husband's college attendance;
- lwg, the woman's log wage rate if she is employed, or her imputed wage rate, if she is not (a variable that Fox and Weisberg 2019, show is problematically defined); and
- inc, family income, in \$1000s, exclusive of wife's income.

We use the glm() function to fit a binary logistic regression to the Mroz data:

```
R> m.mroz <- glm(lfp ~ ., data=Mroz, family=binomial)</pre>
R> summary(m.mroz)
Call:
glm(formula = lfp ~ ., family = binomial, data = Mroz)
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
(Intercept) 3.182140
                        0.644375 4.938 7.88e-07
k5
            -1.462913
                        0.197001 -7.426 1.12e-13
k618
            -0.064571 0.068001 -0.950 0.342337
            -0.062871 0.012783 -4.918 8.73e-07
age
wcyes
             0.807274
                        0.229980
                                  3.510 0.000448
hcyes
             0.111734
                       0.206040
                                  0.542 0.587618
             0.604693
                        0.150818
                                   4.009 6.09e-05
lwg
                        0.008208 -4.196 2.71e-05
            -0.034446
inc
(Dispersion parameter for binomial family taken to be 1)
```

```
degrees of freedom
    Null deviance: 1029.75
                            on 752
Residual deviance: 905.27
                            on 745
                                    degrees of freedom
AIC: 921.27
```

Number of Fisher Scoring iterations: 4

```
R> BayesRule(ifelse(Mroz$lfp == "yes", 1, 0),
            fitted(m.mroz, type="response"))
```

```
[1] 0.3067729
attr(,"casewise loss")
[1] "y != round(yhat)"
```

In addition to the usually summary output for a GLM, we show the result of applying the BayesRule() function from the \mathbf{cv} package to predictions derived from the fitted model. Bayes rule, which predicts a "success" in a binary regression model when the fitted probability of success [i.e., $\phi = \Pr(y=1)$] is $\hat{\phi} \geq .5$ and a "failure" if $\hat{\phi} < .5$.⁴ The first argument to BayesRule() is the binary $\{0, 1\}$ response, and the second argument is the predicted probability of success. BayesRule() returns the proportion of predictions that are *in error*, as appropriate for a "cost" function.

The value returned by BayesRule() is associated with an "attribute" named "casewise loss" and set to "y != round(yhat)", signifying that the Bayes rule CV criterion is computed as the mean of casewise values, here 0 if the prediction for a case matches the observed value and 1 if it does not (signifying a prediction error). The mse() function for numeric responses is also calculated as a casewise average. Some other criteria, such as the median absolute error, computed by the medAbsErr() function in the cv package, aren't averages of casewise components. The distinction is important because, to our knowledge, the statistical theory of cross-validation, for example, in Davison and Hinkley (1997), Bates, Hastie, and Tibshirani (2023), and Arlot and Celisse (2010), is developed for CV criteria like MSE that are means of casewise components. As a consequence, we limit computation of bias adjustment and confidence intervals (see below) to criteria that are casewise averages.

In this example, the fitted logistic regression incorrectly predicts 31% of the responses; we expect this estimate to be optimistic given that the model is used to "predict" the data to which it is fit.

The "glm" method for cv() is largely similar to the "lm" method, although the default algorithm, selected explicitly by method="exact", refits the model with each fold removed (and is thus equivalent to method="naive" for "lm" models). For generalized linear models, method="Woodbury" or (for LOO CV) method="hatvalues" provide approximate results (see the last section of the vignette for details):

```
R> cv(m.mroz, criterion=BayesRule, seed=248)

R RNG seed set to 248

10-Fold Cross Validation
method: exact
criterion: BayesRule
cross-validation criterion = 0.3240372
bias-adjusted cross-validation criterion = 0.3195223
95% CI for bias-adjusted CV criterion = (0.2860721, 0.3529724)
full-sample criterion = 0.3067729
```

R> cv(m.mroz, criterion=BayesRule, seed=248, method="Woodbury")

⁴BayesRule() does some error checking; BayesRule2() is similar, but omits the error checking, and so can be faster for large problems.

R RNG seed set to 248

10-Fold Cross Validation
method: Woodbury
criterion: BayesRule
cross-validation criterion = 0.3240372
bias-adjusted cross-validation criterion = 0.3192577
95% CI for bias-adjusted CV criterion = (0.2858076, 0.3527079)
full-sample criterion = 0.3067729

To ensure that the two methods use the same 10 folds, we specify the seed for R's random-number generator explicitly; here, and as is common in our experience, the "exact" and "Woodbury" algorithms produce nearly identical results. The CV estimates of prediction error are slightly higher than the estimate based on all of the cases.

The printed output includes a 95% confidence interval for the bias-adjusted Bayes rule CV criterion. Bates *et al.* (2023) show that these confidence intervals are unreliable for models fit to small samples, and by default cv() computes them only when the sample size is 400 or larger and when the CV criterion employed is an average of casewise components, as is the case for Bayes rule. See the final section of the vignette for details of the computation of confidence intervals for bias-adjusted CV criteria.

Here are results of applying LOO CV to the Mroz model, using both the exact and the approximate methods:

R> cv(m.mroz, k="loo", criterion=BayesRule) n-Fold Cross Validation method: exact criterion: BayesRule cross-validation criterion = 0.3200531 bias-adjusted cross-validation criterion = 0.3183001 95% CI for bias-adjusted CV criterion = (0.2849584, 0.3516418) full-sample criterion = 0.3067729R> cv(m.mroz, k="loo", criterion=BayesRule, method="Woodbury") n-Fold Cross Validation method: Woodbury criterion: BayesRule cross-validation criterion = 0.3200531 bias-adjusted cross-validation criterion = 0.3183001 95% CI for bias-adjusted CV criterion = (0.2849584, 0.3516418) full-sample criterion = 0.3067729

R> cv(m.mroz, k="loo", criterion=BayesRule, method="hatvalues")

```
n-Fold Cross Validation
method: hatvalues
criterion: BayesRule
cross-validation criterion = 0.3200531
```

To the number of decimal digits shown, the three methods produce identical results for this example.

As for linear models, we report some timings for the various cv() methods of computation in LOO CV as well as for the cv.glm() function from the **boot** package (which, recall, refits the model with each case removed, and thus is comparable to cv() with method="exact"):

```
R> microbenchmark::microbenchmark(
    hatvalues=cv(m.mroz, k="loo", criterion=BayesRule, method="hatvalues"),
    Woodbury=cv(m.mroz, k="loo", criterion=BayesRule, method="Woodbury"),
    exact=cv(m.mroz, k="loo", criterion=BayesRule),
    cv.glm=boot::cv.glm(Mroz, m.mroz,
                 cost=BayesRule),
    times=10)
Unit: milliseconds
      expr
                   min
                                 lq
                                           mean
                                                      median
                                                                      uq
 hatvalues
              1.316141
                           1.326842
                                       1.654063
                                                    1.407571
                                                                1.442462
  Woodbury
             37.127427
                          38.641639
                                      39.368192
                                                   39.356904
                                                               40.562202
     exact 1757.625925 1827.861385 1905.072191 1911.421128 1951.049944
    cv.glm 2062.693354 2122.649130 2149.761134 2140.079440 2184.788525
         max neval cld
    4.022428
                10 a
   41.060188
                10 a
 2128.307540
                10
                    b
 2299.544573
                10
```

There is a substantial time penalty associated with exact computations.

3. Cross-validating mixed-effects models

The fundamental analogy for cross-validation is to the collection of new data. That is, predicting the response in each fold from the model fit to data in the other folds is like using the model fit to all of the data to predict the response for new cases from the values of the predictors for those new cases. As we explained, the application of this idea to independently sampled cases is straightforward—simply partition the data into random folds of equal size and leave each fold out in turn, or, in the case of LOO CV, simply omit each case in turn.

In contrast, mixed-effects models are fit to *dependent* data, in which cases as clustered, such as hierarchical data, where the clusters comprise higher-level units (e.g., students clustered in schools), or longitudinal data, where the clusters are individuals and the cases repeated

observations on the individuals over time.⁵

We can think of two approaches to applying cross-validation to clustered data:⁶

- 1. Treat CV as analogous to predicting the response for one or more cases in a *newly observed cluster*. In this instance, the folds comprise one or more whole clusters; we refit the model with all of the cases in clusters in the current fold removed; and then we predict the response for the cases in clusters in the current fold. These predictions are based only on fixed effects because the random effects for the omitted clusters are presumably unknown, as they would be for data on cases in newly observed clusters.
- 2. Treat CV as analogous to predicting the response for a newly observed case in an existing cluster. In this instance, the folds comprise one or more individual cases, and the predictions can use both the fixed and random effects.

3.1. Example: The High-School and Beyond data

Following their use by Raudenbush and Bryk (2002), data from the 1982 *High School and Beyond* (HSB) survey have become a staple of the literature on mixed-effects models. The HSB data are used by Fox and Weisberg (2019, Sec. 7.2.2) to illustrate the application of linear mixed models to hierarchical data, and we'll closely follow their example here.

The HSB data are included in the MathAchieve and MathAchSchool data sets in the nlme package (Pinheiro and Bates 2000). MathAchieve includes individual-level data on 7185 students in 160 high schools, and MathAchSchool includes school-level data:

```
R> data("MathAchieve", package="nlme")
R> dim(MathAchieve)
[1] 7185
            6
R> head(MathAchieve, 3)
Grouped Data: MathAch ~ SES | School
  School Minority
                      Sex
                             SES MathAch MEANSES
    1224
               No Female -1.528
                                    5.876
                                           -0.428
1
2
    1224
               No Female -0.588
                                  19.708
                                           -0.428
                     Male -0.528
3
    1224
               No
                                  20.349
                                           -0.428
```

R> tail(MathAchieve, 3)

⁵There are, however, more complex situations that give rise to so-called *crossed* (rather than *nested*) random effects. For example, consider students within classes within schools. In primary schools, students typically are in a single class, and so classes are nested within schools. In secondary schools, however, students typically take several classes and students who are together in a particular class may not be together in other classes; consequently, random effects based on classes within schools are crossed. The lmer() function in the lme4 package is capable of modeling both nested and crossed random effects, and the cv() methods for mixed models in the cv package pertain to both nested and crossed random effects. We present an example of the latter later in the vignette.

⁶We subsequently discovered that Vehtari (2023, Section 8) makes similar points.

```
Grouped Data: MathAch ~ SES | School
     School Minority
                        Sex
                                SES MathAch MEANSES
7183
       9586
                  No Female 1.332 19.641
                                              0.627
7184
       9586
                  No Female -0.008
                                    16.241
                                              0.627
7185
       9586
                  No Female 0.792 22.733
                                              0.627
R> data("MathAchSchool", package="nlme")
R> dim(MathAchSchool)
Γ17 160
R> head(MathAchSchool, 2)
     School Size Sector PRACAD DISCLIM HIMINTY MEANSES
1224
       1224 842 Public
                          0.35
                                  1.597
1288
       1288 1855 Public
                           0.27
                                  0.174
                                              0
                                                  0.128
R> tail(MathAchSchool, 2)
                   Sector PRACAD DISCLIM HIMINTY MEANSES
     School Size
9550
       9550 1532
                   Public
                            0.45
                                    0.791
                                                0
                                                    0.059
9586
       9586
            262 Catholic
                             1.00 - 2.416
                                                    0.627
```

The first few students are in school number 1224 and the last few in school 9586.

We'll use only the School, SES (students' socioeconomic status), and MathAch (their score on a standardized math-achievement test) variables in the MathAchieve data set, and Sector ("Catholic" or "Public") in the MathAchSchool data set.

Some data-management is required before fitting a mixed-effects model to the HSB data, for which we use the **dplyr** package (Wickham, François, Henry, Müller, and Vaughan 2023):

```
R> library("dplyr")
Attaching package: 'dplyr'
The following objects are masked from 'package:stats':
    filter, lag
The following objects are masked from 'package:base':
    intersect, setdiff, setequal, union
R> MathAchieve %>% group_by(School) %>%
    summarize(mean.ses = mean(SES)) -> Temp
R> Temp <- merge(MathAchSchool, Temp, by="School")</pre>
```

In the process, we created two new school-level variables: meanses, which is the average SES for students in each school; and cses, which is school-average SES centered at its mean. For details, see Fox and Weisberg (2019, Sec. 7.2.2).

Still following Fox and Weisberg, we proceed to use the lmer() function in the lme4 package (Bates, Mächler, Bolker, and Walker 2015) to fit a mixed model for math achievement to the HSB data:

```
R> library("lme4")
```

Loading required package: Matrix

```
Linear mixed model fit by REML ['lmerMod']
Formula: mathach ~ mean.ses * cses + sector * cses + (cses | school)
   Data: HSB
```

REML criterion at convergence: 46503.7

Scaled residuals:

```
Min 1Q Median 3Q Max -3.15926 -0.72319 0.01704 0.75444 2.95822
```

Random effects:

Groups	Name	Variance	Std.Dev	. Corr
school	(Intercept)	2.380	1.5426	
	cses	0.101	0.3179	0.39
Residual		36.721	6.0598	
Number of	obs: 7185,	groups:	school,	160

Fixed effects:

	Fetimata	Std. Error	+ 172]110
	претшесе	bta. Liioi	t varue
(Intercept)	12.1279	0.1993	60.856
mean.ses	5.3329	0.3692	14.446
cses	2.9450	0.1556	18.928
sectorCatholic	1.2266	0.3063	4.005
mean.ses:cses	1.0393	0.2989	3.477
cses:sectorCatholic	-1.6427	0.2398	-6.851

```
We can then cross-validate at the cluster (i.e., school) level,
R> cv(hsb.lmer, k=10, clusterVariables="school", seed=5240)
R RNG seed set to 5240
10-Fold Cross Validation based on 160 {school} clusters
cross-validation criterion = 39.15662
bias-adjusted cross-validation criterion = 39.14844
95% CI for bias-adjusted CV criterion = (38.06554, 40.23135)
full-sample criterion = 39.00599
or at the case (i.e., student) level,
R> cv(hsb.lmer, seed=1575)
R RNG seed set to 1575
Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
Model failed to converge with max|grad| = 0.00587228 (tol = 0.002, component 1)
boundary (singular) fit: see help('isSingular')
10-Fold Cross Validation
cross-validation criterion = 37.44473
bias-adjusted cross-validation criterion = 37.33801
95% CI for bias-adjusted CV criterion = (36.28761, 38.38841)
```

For cluster-level CV, the clusterVariables argument tells cv() how the clusters are defined. Were there more than one clustering variable, say classes within schools, these would be provided as a character vector of variable names: clusterVariables = c("school", "class"). For cluster-level CV, the default is k = "loo", that is, leave one cluster out at a time; we instead specify k = 10 folds of clusters, each fold therefore comprising 160/10 = 16 schools.

full-sample criterion = 36.06767

If the clusterVariables argument is omitted, then case-level CV is employed, with $\tt k=10$ folds as the default, here each with $7185/10\approx719$ students. Notice that one of the 10 models refit with a fold removed failed to converge. Convergence problems are common in mixed-effects modeling. The apparent issue here is that an estimated variance component is close to or equal to 0, which is at a boundary of the parameter space. That shouldn't disqualify the fitted model for the kind of prediction required for cross-validation.

There is also a cv() method for linear mixed models fit by the lme() function in the nlme package, and the arguments for cv() in this case are the same as for a model fit by lmer() or glmer(). We illustrate with the mixed model fit to the HSB data:

```
R> library(nlme)
R> hsb.lme <- lme(mathach ~ mean.ses*cses + sector*cses,
                 random = ~ cses | school, data=HSB,
                control=list(opt="optim"))
R> summary(hsb.lme)
Linear mixed-effects model fit by REML
  Data: HSB
      AIC
               BIC
                      logLik
  46524.78 46593.57 -23252.39
Random effects:
 Formula: ~cses | school
 Structure: General positive-definite, Log-Cholesky parametrization
           StdDev
                     Corr
(Intercept) 1.54117685 (Intr)
           0.01817364 0.006
cses
Residual
           6.06349216
Fixed effects: mathach ~ mean.ses * cses + sector * cses
                      Value Std.Error DF t-value p-value
                 12.128207 0.1991964 7022 60.88567
(Intercept)
                                                      0e+00
                   5.336665 0.3689784 157 14.46335
mean.ses
                                                      0e+00
                   2.942145 0.1512240 7022 19.45554 0e+00
cses
sectorCatholic
                   1.224531 0.3061139 157 4.00025
                                                     1e-04
mean.ses:cses
                   1.044406 0.2910747 7022 3.58810 3e-04
cses:sectorCatholic -1.642148 0.2331162 7022 -7.04433 0e+00
 Correlation:
                   (Intr) men.ss cses sctrCt mn.ss:
                    0.256
mean.ses
                    0.000 0.000
cses
sectorCatholic
                   -0.699 -0.356 0.000
                    0.000 0.000 0.295 0.000
mean.ses:cses
cses:sectorCatholic 0.000 0.000 -0.696 0.000 -0.351
Standardized Within-Group Residuals:
                    Q1
                               Med
                                            QЗ
-3.17010624 -0.72487654 0.01489162 0.75426269 2.96549829
Number of Observations: 7185
Number of Groups: 160
R> cv(hsb.lme, k=10, clusterVariables="school", seed=5240)
R RNG seed set to 5240
10-Fold Cross Validation based on 160 {school} clusters
cross-validation criterion = 39.1569
```

```
bias-adjusted cross-validation criterion = 39.14881
95% CI for bias-adjusted CV criterion = (38.06591, 40.23171)
full-sample criterion = 39.0062

R> cv(hsb.lme, seed=1575)

R RNG seed set to 1575

10-Fold Cross Validation
cross-validation criterion = 37.44163
bias-adjusted cross-validation criterion = 37.40222
95% CI for bias-adjusted CV criterion = (36.35136, 38.45309)
full-sample criterion = 36.14707
```

We used the same random-number generator seeds as in the previous example cross-validating the model fit by lmer(), and so the same folds are employed in both cases. The estimated covariance components and fixed effects in the summary output differ slightly between the lmer() and lme() solutions, although both functions seek to maximize the REML criterion. This is, of course, to be expected when different algorithms are used for numerical optimization. To the precision reported, the cluster-level CV results for the lmer() and lme() models are identical, while the case-level CV results are very similar but not identical.

3.2. Example: Contrived hierarchical data

We introduce an artificial data set that exemplifies aspects of cross-validation particular to hierarchical models. Using this data set, we show that model comparisons employing cluster-based and those employing case-based cross-validation may not agree on a "best" model. Furthermore, commonly used measures of fit, such as mean-squared error, do not necessarily become smaller as models become larger, even when the models are nested, and even when the measure of fit is computed for the whole data set.

Consider a researcher studying improvement in a skill, yodeling, for example, among students enrolled in a four-year yodeling program. The plan is to measure each student's skill level at the beginning of the program and every year thereafter until the end of the program, resulting in five annual measurements for each student. It turns out that yodeling appeals to students of all ages, and students enrolling in the program range in age from 20 to 70. Moreover, participants' untrained yodeling skill is similar at all ages, as is their rate of progress with training. All students complete the four-year program.

The researcher, who has more expertise in yodeling than in modeling, decides to model the response, y, yodeling skill, as a function of age, x, reasoning that students get older during their stay in the program, and (incorrectly) that age can serve as a proxy for elapsed time. The researcher knows that a mixed model should be used to account for clustering due to the expected similarity of measurements taken from each student.

⁷The observant reader will notice that we set the argument control=list(opt="optim") in the call to lme(), changing the optimizer employed from the default "nlminb". We did this because with the default optimizer, lme() encountered the same convergence issue as lmer(), but rather than issuing a warning, lme() failed, reporting an error. As it turns out, setting the optimizer to "optim" avoids this problem.

We start by generating the data, using parameters consistent with the description above and meant to highlight the issues that arise in cross-validating mixed-effects models:⁸

```
R> # Parameters:
R> set.seed(9693)
R> Nb <- 100
                # number of groups
R> Nw <- 5
                 # number of individuals within groups
R> Bb <- 0
                 # between-group regression coefficient on group mean
R> SDre <- 2.0
               # between-group SD of random level relative to group mean of x
R> SDwithin <- 0.5 # within group SD
R> Bw <- 1
                    # within group effect of x
R> Ay <- 10
                    # intercept for response
R > Ax < -20
                    # starting level of x
R > Nx < - Nw*10
                    # number of distinct x values
R>
R> Data <- data.frame(</pre>
    group = factor(rep(1:Nb, each=Nw)),
    x = Ax + rep(1:Nx, length.out = Nw*Nb)
+ ) />
    within(
      {
        xm <- ave(x, group, FUN = mean) # within-group mean
        y \leftarrow Ay +
          Bb * xm +
                                        # contextual effect
          Bw * (x - xm) +
                                        # within-group effect
          rnorm(Nb, sd=SDre)[group] + # random level by group
          rnorm(Nb*Nw, sd=SDwithin)
                                        # random error within groups
    )
```

Here is a scatterplot of the data for a representative group of 10 (without loss of generality, the first 10) of 100 students, showing the 95% concentration ellipse for each cluster:⁹

⁸The regsubsets() function computes several measures of model predictive performance, including the R^2 and R^2 adjusted for degrees of freedom, the residual sums of squares, Mallows's C_p , and the BIC. Several of these are suitable for comparing models with differing numbers of coefficients—we use the BIC below—but all necessarily agree when comparing models with the *same* number of coefficients.

⁹We find it convenient to use the **lattice** (Sarkar 2008) and **latticeExtra** (Sarkar and Andrews 2022) packages for this and other graphs in this section.

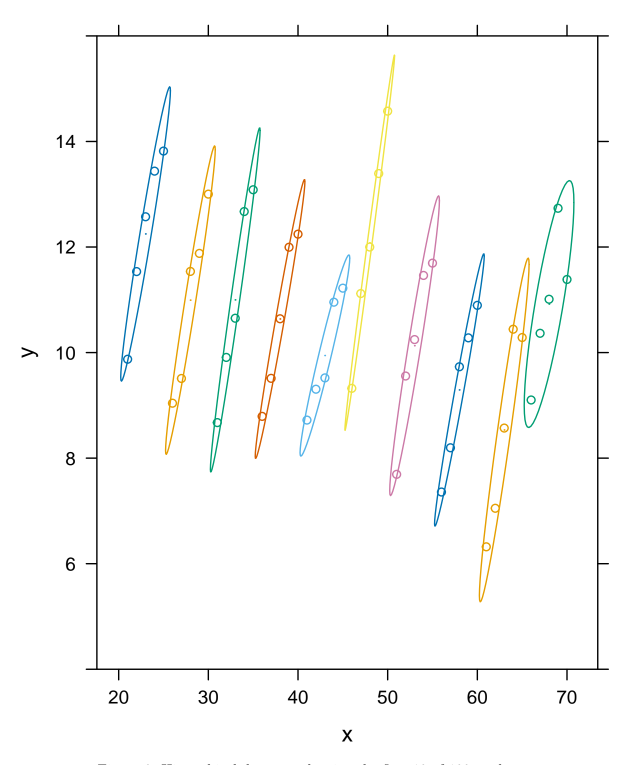


Figure 6: Hierarchical data set, showing the first 10 of 100 students.

The between-student effect of age is 0 but the within-student effect is 1. Due to the large variation in ages between students, the least-squares regression of yodeling skill on age (for the 500 observations among all 100 students) produces an estimated slope close to 0 (though with a small p-value), because the slope is heavily weighted toward the between-student effect:

```
R> summary(lm(y ~ x, data=Data))
```

```
Call:
```

 $lm(formula = y \sim x, data = Data)$

Residuals:

Min 1Q Median 3Q Max -5.7713 -1.6583 -0.0894 1.5520 7.6240

Coefficients:

Residual standard error: 2.347 on 498 degrees of freedom Multiple R-squared: 0.01632, Adjusted R-squared: 0.01435 F-statistic: 8.263 on 1 and 498 DF, p-value: 0.004219

The initial mixed-effects model that we fit to the data is a simple random-intercepts model:

```
R> # random intercept only:
R> mod.0 <- lmer(y ~ 1 + (1 | group), Data)
R> summary(mod.0)
```

Linear mixed model fit by REML ['lmerMod']
Formula: y ~ 1 + (1 | group)
 Data: Data

REML criterion at convergence: 2103.1

Scaled residuals:

Min 1Q Median 3Q Max -2.03514 -0.72645 -0.01169 0.78477 2.04377

Random effects:

Groups Name Variance Std.Dev. group (Intercept) 2.900 1.703
Residual 2.712 1.647
Number of obs: 500, groups: group, 100

Fixed effects:

```
Estimate Std. Error t value (Intercept) 10.0018 0.1855 53.91
```

We will shortly consider three other, more complex, mixed models; because of data-management considerations, it is convenient to fit them now, but we defer discussion of these models:

```
R> # effect of x and random intercept:
R> mod.1 <- lmer(y ~ x + (1 | group), Data)
R>
R> # effect of x, contextual (student) mean of x, and random intercept:
R> mod.2 <- lmer(y ~ x + xm + (1 | group), Data)
R> # equivalent to y ~ I(x - xm) + xm + (1 | group)
R>
R> # model generating the data (where Bb = 0)
R> mod.3 <- lmer(y ~ I(x - xm) + (1 | group), Data)
```

We proceed to obtain predictions from the random-intercept model (mod.0) and the other models (mod.1, mod.2, and mod.3) based on fixed effects alone, as would be used for cross-validation based on clusters (i.e., students), and for fixed and random effects—so-called best linear unbiased predictions or BLUPs—as would be used for cross-validation based on cases (i.e., occasions within students):

We then prepare the data for plotting:

Predictions based on the random-intercept model mod.0 for the first 10 students are shown in the following graph:

The fixed-effect predictions for the various individuals are identical—the estimated fixed-effects intercept or estimated general mean of y—while the BLUPs are the sums of the fixed-effects intercept and the random intercepts, and are only slightly shrunken towards the general mean. Because in our artificial data there is no population relationship between age and skill, the fixed-effect-only predictions and the BLUPs are not very different.

Our next model, mod.1, includes a fixed intercept and fixed effect of x along with a random intercept:

```
R> summary(mod.1)
Linear mixed model fit by REML ['lmerMod']
Formula: y \sim x + (1 \mid group)
   Data: Data
REML criterion at convergence: 1564.5
Scaled residuals:
                                 ЗQ
     Min
               10
                  Median
                                         Max
-2.90160 -0.63501 0.01879 0.55407 2.82932
Random effects:
 Groups
                      Variance Std.Dev.
          (Intercept) 192.9406 13.8903
 group
                        0.2569 0.5068
 Residual
Number of obs: 500, groups: group, 100
Fixed effects:
             Estimate Std. Error t value
(Intercept) -33.91892
                         1.56446 -21.68
              0.96529
                         0.01581
                                   61.05
Correlation of Fixed Effects:
  (Intr)
x - 0.460
```

Model: y ~ 1 + (1 | group)

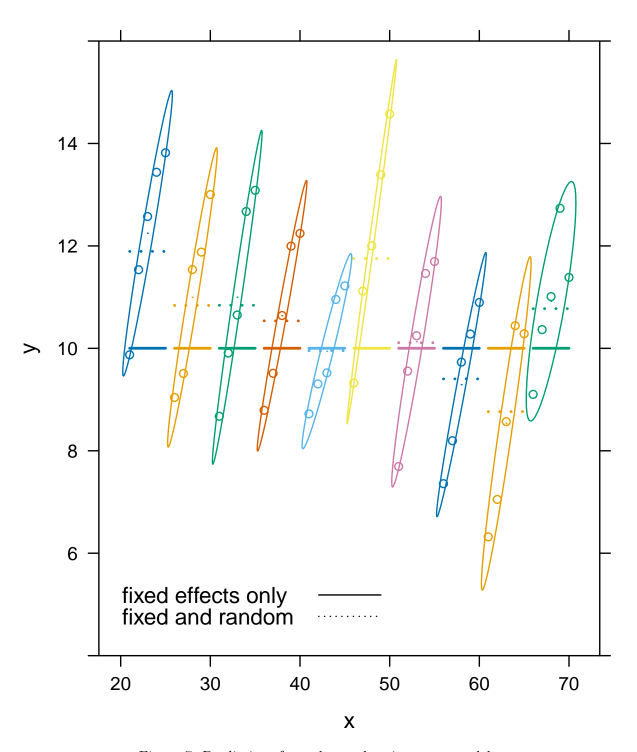


Figure 7: Predictions from the random intercept model.

Predictions from this model appear in the following graph:

The BLUPs fit the observed data very closely, but predictions based on the fixed effects alone, with a common intercept and slope for all clusters, are very poor—indeed, much worse than the fixed-effects-only predictions based on the simpler random-intercept model, mod.0. We therefore anticipate (and show later in this section) that case-based cross-validation will prefer mod1 to mod0, but that cluster-based cross-validation will prefer mod0 to mod1.

Our third model, mod.2, includes the contextual effect of x—that is, the cluster mean xm—along with x and the intercept in the fixed-effect part of the model, and a random intercept:

```
R> summary(mod.2)
Linear mixed model fit by REML ['lmerMod']
Formula: y \sim x + xm + (1 \mid group)
   Data: Data
REML criterion at convergence: 1169.2
Scaled residuals:
     Min
               1Q
                    Median
                                  3Q
                                          Max
-2.98466 -0.63750 0.00191 0.55682 2.73246
Random effects:
 Groups
          Name
                       Variance Std.Dev.
          (Intercept) 3.3986
                                1.8435
 group
 Residual
                       0.2552
                                0.5052
Number of obs: 500, groups: group, 100
Fixed effects:
            Estimate Std. Error t value
(Intercept) 9.47866
                         0.61705
                                   15.36
х
             0.99147
                         0.01597
                                   62.07
            -0.97998
                         0.02055 -47.68
xm
```

Model: $y \sim 1 + x + (1 | group)$

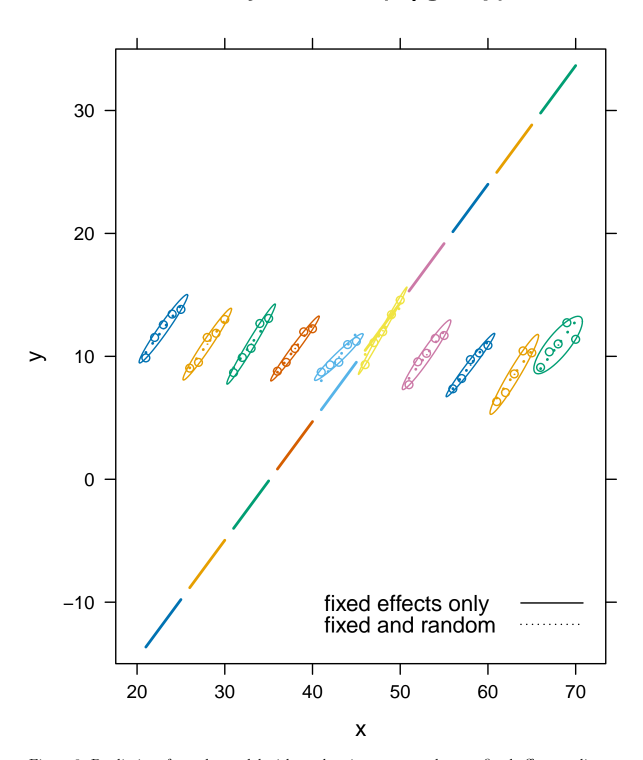


Figure 8: Predictions from the model with random intercepts and x as a fixed-effect predictor.

```
Correlation of Fixed Effects:
(Intr) x
x 0.000
xm -0.600 -0.777
```

This model is equivalent to fitting $y \sim I(x - xm) + xm + (1 | group)$, which is the model that generated the data once the coefficient of the contextual predictor xm is set to 0 (as it is in mod.3, discussed below).

Predictions from model mod.2 appear in the following graph:

Depending on the estimated variance parameters of the model, a mixed model like mod.2 will apply varying degrees of shrinkage to the random-intercept BLUPs that correspond to variation in the heights of the parallel fitted lines for the individual students. In our contrived data, the mod.2 applies little shrinkage, allowing substantial variability in the heights of the fitted lines, which closely approach the observed values for each student. The fit of the mixed model mod.2 is consequently similar to that of a fixed-effects model with age and a categorical predictor for individual students (i.e., treating students as a factor, and not shown here).

The mixed model mod.2 therefore fits individual observations well, and we anticipate a favorable assessment using individual-based cross-validation. In contrast, the large variability in the BLUPs results in larger residuals for predictions based on fixed effects alone, and so we expect that cluster-based cross-validation won't show an advantage for model mod.2 compared to the smaller model mod.0, which includes only fixed and random intercepts.

Had the mixed model applied considerable shrinkage, then neither cluster-based nor case-based cross-validation would show much improvement over the random-intercept-only model. In our experience, the degree of shrinkage does not vary smoothly as parameters are changed but tends to be "all or nothing," and near the tipping point, the behavior of estimates can be affected considerably by the choice of algorithm used to fit the model.

Finally, mod.3 directly estimates the model used to generate the data. As mentioned, it is a constrained version of mod.2, with the coefficient of xm set to 0, and with x expressed as a deviation from the cluster mean xm:

```
R> summary(mod.3)
```

Model: $y \sim 1 + x + xm + (1 | group)$

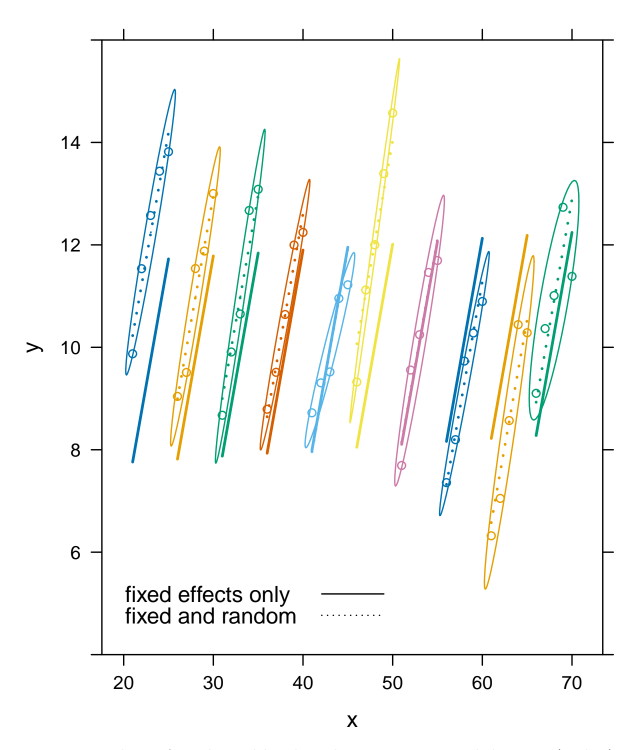


Figure 9: Predictors from the model with random intercepts, x, and the group (student) mean of x as predictors.

```
Linear mixed model fit by REML ['lmerMod']
Formula: y \sim I(x - xm) + (1 | group)
   Data: Data
REML criterion at convergence: 1163.2
Scaled residuals:
     Min
               1Q
                   Median
                                  3Q
                                          Max
-2.97703 -0.63204 0.00627 0.56032 2.72489
Random effects:
 Groups
          Name
                      Variance Std.Dev.
          (Intercept) 3.3913
 group
                               1.8415
 Residual
                      0.2552
                               0.5052
Number of obs: 500, groups: group, 100
Fixed effects:
            Estimate Std. Error t value
(Intercept) 10.00176
                        0.18553
                                  53.91
I(x - xm)
            0.99147
                        0.01597
                                  62.07
Correlation of Fixed Effects:
          (Intr)
I(x - xm) 0.000
The predictions from mod.3 are therefore similar to those from mod.2:
R> (plot +
+
    xyplot(fit ~ x, subset(Data_long, modelcode == "mod3" & effect == "fe"),
           groups=group, type="1", 1wd=2) +
    xyplot(fit ~ x, subset(Data_long, modelcode == "mod3" & effect == "re"),
           groups=group, type="1", 1wd=2, 1ty=3)
+ ) |> update(
    main="Model: y \sim 1 + I(x - xm) + (1 | group)",
    ylim=c(4, 16),
    key=list(
      corner=c(0.05, 0.05),
      text=list(c("fixed effects only", "fixed and random")),
      lines=list(lty=c(1, 3)))
```

We next carry out case-based cross-validation, which, as we have explained, is based on both fixed and predicted random effects (i.e., BLUPs), and cluster-based cross-validation, which is based on fixed effects only. In order to reduce between-model random variability in comparisons of models, we apply cv() to the list of models created by the models() function (introduced previously), performing cross-validation with the same folds for each model:

Model: $y \sim 1 + I(x - xm) + (1 | group)$

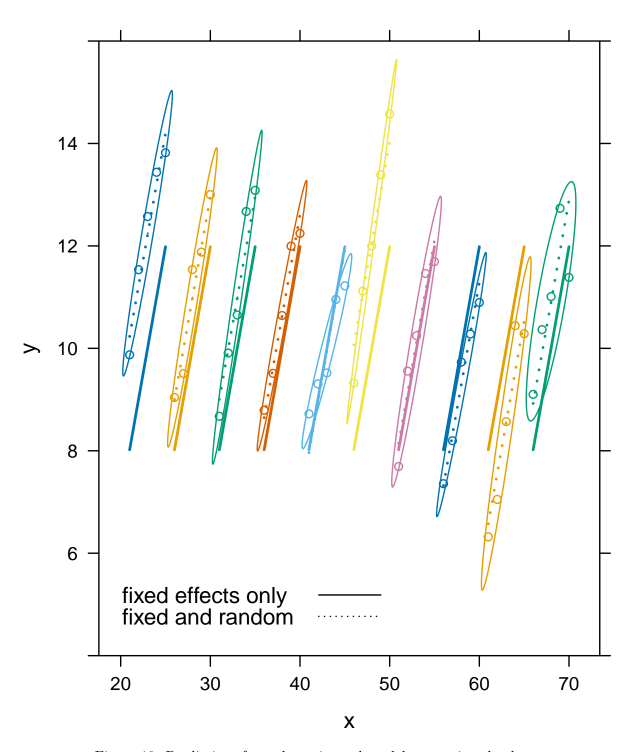


Figure 10: Predictions from the estimated model generating the data.

```
R> cvs_clusters <- cv(modlist, data=Data, cluster="group", k=10, seed=6449)
R> plot(cvs_clusters, main="Model Comparison, Cluster-Based CV")
R> cvs_cases <- cv(modlist, data=Data, seed=9693)</pre>
```

R> plot(cvs cases, main="Model Comparison, Case-Based CV")

In summary, model mod.1, with x alone and without the contextual mean of x, is assessed as fitting very poorly by cluster-based CV, but relatively much better by case-based CV. Model mod.2, which includes both x and its contextual mean, produces better results using both cluster-based and case-based CV. The data-generating model, mod.3, which includes the fixed effect of x - xm in place of separate terms in x and xm, isn't distinguishable from model mod.2, which includes x and xm separately, even though mod.2 has an unnecessary parameter (recall that the population coefficient of xm is 0 when x is expressed as deviations from the contextual mean). These conclusions are consistent with our observations based on graphing predictions from the various models, and they illustrate the desirability of assessing mixed-effect models at different hierarchical levels.

3.3. Example: Crossed random effects

Crossed random effects arise when the structure of the data aren't strictly hierarchical. Nevertheless, crossed and nested random effects can be handled in much the same manner, by refitting the mixed-effects model to the data with a fold of clusters or cases removed and using the refitted model to predict the response in the removed fold.

We'll illustrate with data on pig growth, introduced by Diggle, Liang, and Zeger (1994, Table 3.1). The data are in the Pigs data frame in the cv package:

```
R> head(Pigs, 9)
  id week weight
        1
            24.0
2
   1
        2
            32.0
3
            39.0
   1
        3
4
   1
        4
            42.5
5
   1
        5
            48.0
6
   1
        6
            54.5
7
        7
            61.0
8
   1
        8
            65.0
9
   1
        9
            72.0
R> head(xtabs(~ id + week, data=Pigs), 3)
   week
id 123456789
  1 1 1 1 1 1 1 1 1 1
  2 1 1 1 1 1 1 1 1 1
  3 1 1 1 1 1 1 1 1 1
```

Model Comparison, Cluster-Based CV

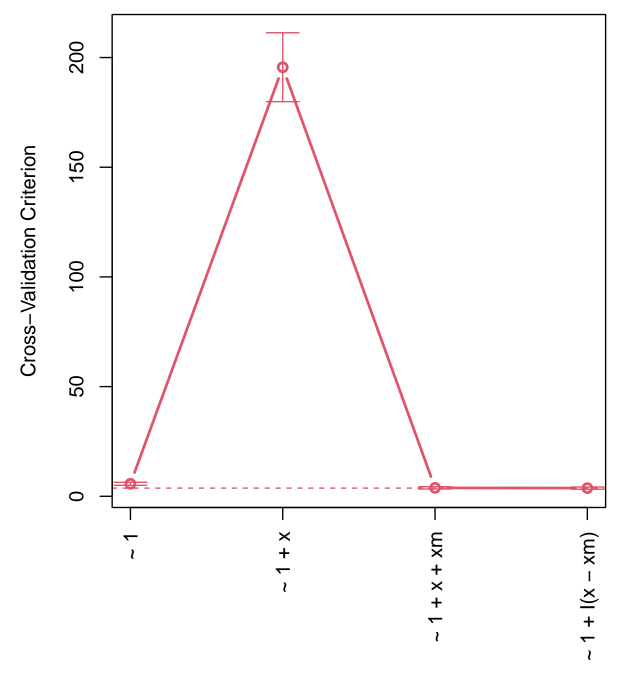


Figure 11: 10-fold cluster-based cross-validation comparing random intercept models with varying fixed effects. The error bars show the 95% confidence interval around the CV estimate of the MSE for each model.

Model Comparison, Case-Based CV

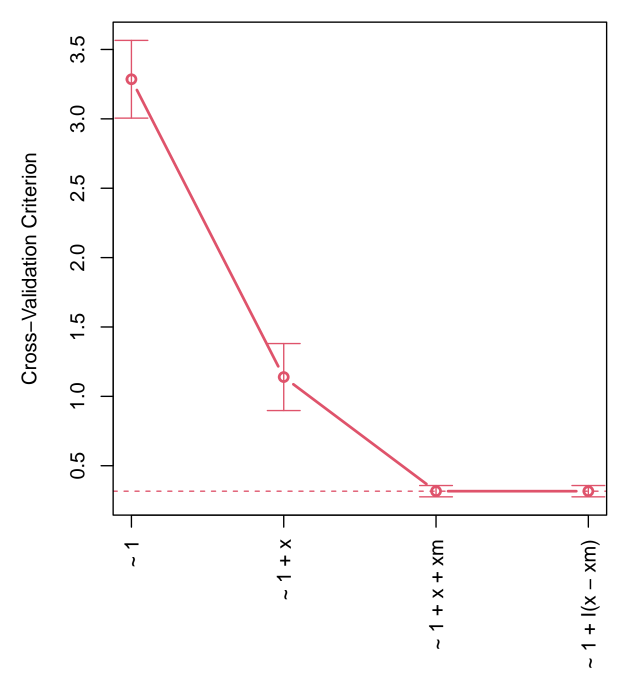


Figure 12: 10-fold case-based cross-validation comparing random intercept models with varying fixed effects.

Each of 48 pigs is observed weekly over a period of 9 weeks, with the weight of the pig recorded in kg. The data are in "long" format, as is appropriate for use with the lmer() function in the lme4 package. The data are very regular, with no missing cases.

The following graph, showing the growth trajectories of the pigs, is similar to Figure 3.1 in Diggle *et al.* (1994); we add an overall least-squares line and a loess smooth, which are nearly indistinguishable:

The individual "growth curves" and the overall trend are generally linear, with some tendency for variability of pig weight to increase over weeks (a feature of the data that we ignore in the mixed model that we fit to the data below).

The **Stata** mixed-effects models manual proposes a model with crossed random effects for the **Pigs** data (StataCorp LLC 2023, page 37):

[S]uppose that we wish to fit

weight_{ij} =
$$\beta_0 + \beta_1$$
week_{ij} + $u_i + v_j + \varepsilon_{ij}$
for the $i = 1, ..., 9$ weeks and $j = 1, ..., 48$ pigs and
 $u_i \sim N(0, \sigma_u^2); v_j \sim N(0, \sigma_v^2); \varepsilon_{ij} \sim N(0, \sigma_\varepsilon^2)$

all independently. That is, we assume an overall population-average growth curve $\beta_0 + \beta_1$ week and a random pig-specific shift. In other words, the effect due to week, u_i , is systematic to that week and common to all pigs. The rationale behind [this model] could be that, assuming that the pigs were measured contemporaneously, we might be concerned that week-specific random factors such as weather and feeding patterns had significant systematic effects on all pigs.

Although we might prefer an alternative model, ¹⁰ we think that this is a reasonable specification.

¹⁰These are repeated-measures data, which would be more conventionally modeled with autocorrelated errors within pigs. The lme() function in the nlme package, for example, is capable of fitting a mixed-model of this form.

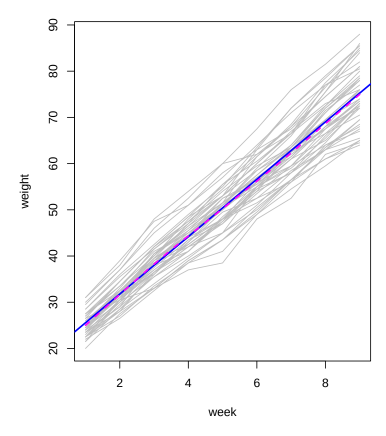


Figure 13: Growth trajectories for 48 pigs, with overall least-squares line (sold blue) and loess line (broken magenta).

The **Stata** manual fits the mixed model by maximum likelihood (rather than REML), and we duplicate the results reported there using <code>lmer()</code>:

```
R > m.p < -lmer(weight ~ week + (1 | id) + (1 | week),
              data=Pigs, REML=FALSE, # i.e., ML
              control=lmerControl(optimizer="bobyqa"))
+
R> summary(m.p)
Linear mixed model fit by maximum likelihood ['lmerMod']
Formula: weight ~ week + (1 | id) + (1 | week)
   Data: Pigs
Control: lmerControl(optimizer = "bobyga")
     AIC
              BIC
                    logLik deviance df.resid
  2037.6
           2058.0 -1013.8
                             2027.6
                                         427
Scaled residuals:
             1Q Median
                             3Q
-3.7750 -0.5418 0.0054 0.4762 3.9816
Random effects:
 Groups
          Name
                      Variance Std.Dev.
 id
          (Intercept) 14.83622 3.8518
 week
          (Intercept) 0.08499 0.2915
 Residual
                       4.29733 2.0730
Number of obs: 432, groups: id, 48; week, 9
Fixed effects:
            Estimate Std. Error t value
(Intercept) 19.35561
                        0.63340
                                  30.56
             6.20990
                        0.05393 115.14
Correlation of Fixed Effects:
     (Intr)
week -0.426
```

We opt for the non-default "bobyqa" optimizer because it provides more numerically stable results for subsequent cross-validation in this example.

We can then cross-validate the model by omitting folds composed of pigs, folds composed of weeks, or folds composed of pig-weeks (which in the Pigs data set correspond to individual cases, using only the fixed effects):

```
R> cv(m.p, clusterVariables="id")
n-Fold Cross Validation based on 48 {id} clusters
cross-validation criterion = 19.97309
```

```
bias-adjusted cross-validation criterion = 19.96496
95% CI for bias-adjusted CV criterion = (17.1249, 22.80502)
full-sample criterion = 19.20076

R> cv(m.p, clusterVariables="week")
```

boundary (singular) fit: see help('isSingular')

n-Fold Cross Validation based on 9 {week} clusters cross-validation criterion = 19.31219 bias-adjusted cross-validation criterion = 19.30499 95% CI for bias-adjusted CV criterion = (16.56624, 22.04373) full-sample criterion = 19.20076

R> cv(m.p, clusterVariables=c("id", "week"), k=10, seed=8469)

R RNG seed set to 8469

10-Fold Cross Validation based on 432 {id, week} clusters cross-validation criterion = 19.2352 bias-adjusted cross-validation criterion = 19.23294 95% CI for bias-adjusted CV criterion = (16.49263, 21.97325) full-sample criterion = 19.20076

We can also cross-validate the individual cases taking account of the random effects (employing the same 10 folds):

R > cv(m.p, k=10, seed=8469)

R RNG seed set to 8469

10-Fold Cross Validation cross-validation criterion = 5.15835 bias-adjusted cross-validation criterion = 5.072933 95% CI for bias-adjusted CV criterion = (4.122952, 6.022913) full-sample criterion = 3.795972

Because these predictions are based on BLUPs, they are more accurate than the predictions based only on fixed effects.¹¹ As well, the difference between the MSE computed for the model

¹¹Even though there is only one observation per combination of pigs and weeks, we can use the BLUP for the omitted case because of the crossed structure of the random effects; that is each pig-week has a pig random effect and a week random effect. Although it probably isn't sensible, we can imagine a mixed model for the pig data that employs nested random effects, which would be specified by lmer(weight ~ week + (1 | id/week), data=Pigs)—that is, a random intercept that varies by combinations of id (pig) and week. This model can't be fit, however: With only one case per combination of id and week, the nested random-effect variance is indistinguishable from the case-level variance.

fit to the full data and the CV estimates of the MSE is greater here than for cluster-based predictions.

4. Replicating cross-validation

Assuming that the number of cases n is a multiple of the number of folds k—a slightly simplifying assumption—the number of possible partitions of cases into folds is $\frac{n!}{[(n/k)!]^k}$, a number that grows very large very quickly. For example, for n=10 and k=5, so that the folds are each of size n/k=2, there are 113,400 possible partitions; for n=100 and k=5, where n/k=20, still a small problem, the number of possible partitions is truly astronomical, 1.09×10^{66} .

Because the partition into folds that's employed is selected randomly, the resulting CV criterion estimates are subject to sampling error. (An exception is LOO cross-validation, which is not at all random.) To get a sense of the magnitude of the sampling error, we can repeat the CV procedure with different randomly selected partitions into folds. All of the CV functions in the **cv** package are capable of repeated cross-validation, with the number of repetitions controlled by the **reps** argument, which defaults to 1.

Here, for example, is 10-fold CV for the Mroz logistic regression, repeated 5 times:

```
R> cv(m.mroz, criterion=BayesRule, seed=248, reps=5,
     method="Woodbury")
R RNG seed set to 248
R RNG seed set to 68134
R RNG seed set to 767359
R RNG seed set to 556270
R RNG seed set to 882966
Replicate 1:
10-Fold Cross Validation
method: Woodbury
criterion: BayesRule
cross-validation criterion = 0.3200531
bias-adjusted cross-validation criterion = 0.3130109
95% CI for bias-adjusted CV criterion = (0.2796692, 0.3463526)
full-sample criterion = 0.3067729
Replicate 2:
10-Fold Cross Validation
method: Woodbury
```

criterion: BayesRule cross-validation criterion = 0.3160691 bias-adjusted cross-validation criterion = 0.311697 95% CI for bias-adjusted CV criterion = (0.2784665, 0.3449274) full-sample criterion = 0.3067729 Replicate 3: 10-Fold Cross Validation method: Woodbury criterion: BayesRule cross-validation criterion = 0.314741 bias-adjusted cross-validation criterion = 0.3086212 95% CI for bias-adjusted CV criterion = (0.2754285, 0.341814) full-sample criterion = 0.3067729 Replicate 4: 10-Fold Cross Validation method: Woodbury criterion: BayesRule cross-validation criterion = 0.3240372 bias-adjusted cross-validation criterion = 0.318069 95% CI for bias-adjusted CV criterion = (0.2846189, 0.3515192) full-sample criterion = 0.3067729 Replicate 5: 10-Fold Cross Validation method: Woodbury criterion: BayesRule cross-validation criterion = 0.3240372 bias-adjusted cross-validation criterion = 0.3192577 95% CI for bias-adjusted CV criterion = (0.2858076, 0.3527079) full-sample criterion = 0.3067729 Average: 10-Fold Cross Validation method: Woodbury criterion: BayesRule cross-validation criterion = 0.3198318 (0.003886968) bias-adjusted cross-validation criterion = 0.3139445 (0.004009339) full-sample criterion = 0.3067729

When reps > 1, the result returned by cv() is an object of class "cvList"—literally a list of "cv" objects. The results are reported for each repetition and then averaged across repetitions, with the standard deviations of the CV criterion and the biased-adjusted CV criterion given in parentheses. In this example, there is therefore little variation across repetitions, increasing our confidence in the reliability of the results.

Notice that the seed that's set in the cv() command pertains to the first repetition and the

seeds for the remaining repetitions are then selected pseudo-randomly.¹² Setting the first seed, however, makes the entire process easily replicable, and the seed for each repetition is stored in the corresponding element of the "cvList" object (which isn't, however, saved in the example).

It's also possible to replicate CV when comparing competing models via the cv() method for "modList" objects. Recall our comparison of polynomial regressions of varying degree fit to the Auto data; we performed 10-fold CV for each of 10 models. Here, we replicate that process 5 times for each model and graph the results:

The graph shows both the average CV criterion and its range for each of the competing models.

5. Cross-validating model selection

5.1. A preliminary example

As Hastie, Tibshirani, and Friedman (2009, Sec. 7.10.2: "The Wrong and Right Way to Do Cross-validation") explain, if the whole data are used to select or fine-tune a statistical model, subsequent cross-validation of the model is intrinsically misleading, because the model is selected to fit the whole data, including the part of the data that remains when each fold is removed.

The following example is similar in spirit to one employed by Hastie *et al.* (2009). Suppose that we randomly generate n=1000 independent observations for a response variable variable $y \sim N(\mu=10,\sigma^2=0)$, and independently sample 1000 observations for p=100 "predictors," x_1,\ldots,x_{100} , each from $x_j \sim N(0,1)$. The response has nothing to do with the predictors and so the population linear-regression model $y_i = \alpha + \beta_1 x_{i1} + \cdots + \beta_{100} x_{i,100} + \varepsilon_i$ has $\alpha=10$ and all $\beta_i=0$.

¹²Because of the manner in which the computation is performed, the order of the replicates in the "cvList" object returned by cv() isn't the same as the order in which the replicates are computed. Each element of the result, however, is a "cv" object with the correct random-number seed saved, and so this technical detail can be safely ignored. The individual "cv" objects are printed in the order in which they are stored rather than the order in which they are computed.

Model Comparison Averaged Across 5 Replications

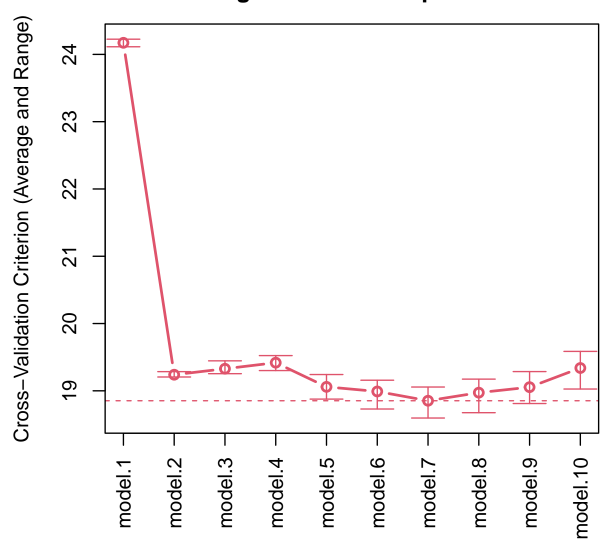


Figure 14: Replicated cross-validated 10-fold CV as a function of polynomial degree, p

```
      y
      X.1
      X.2
      X.3
      X.4
      X.5

      1 10.031647 -1.2388628 -0.2648705 -0.03539048 -2.57697337 0.81104761
      0.81104761

      2 9.664989 0.1228689 -0.1774440 0.37290421 -0.93513788 0.62867324

      3 10.023249 -0.9505172 -0.7348667 -1.05978180 0.88294443 0.02391808

      4 8.990969 1.1357103 0.3241085 0.11036901 1.37630285 -0.42211426

      5 9.071249 1.4947403 1.8753802 0.10574793 0.29213991 -0.18456833

      6 11.349283 -0.1845331 -0.7803709 -1.23803778 -0.01094861 0.69103395
```

Least-squares provides accurate estimates of the regression constant $\alpha=10$ and the error variance $\sigma^2=1$ for the "null model" including only the regression constant; moreover, the omnibus F-test of the correct null hypothesis that all of the β s are 0 for the "full model" with all 100 xs is associated with a large p-value:

```
R> m.full <- lm(y ~ ., data=D)
R> m.null <- lm(y ~ 1, data=D)
R> anova(m.null, m.full)
```

Analysis of Variance Table

```
Model 1: y ~ 1
Model 2: y \sim X.1 + X.2 + X.3 + X.4 + X.5 + X.6 + X.7 + X.8 + X.9 + X.10 +
    X.11 + X.12 + X.13 + X.14 + X.15 + X.16 + X.17 + X.18 + X.19 +
    X.20 + X.21 + X.22 + X.23 + X.24 + X.25 + X.26 + X.27 + X.28 +
    X.29 + X.30 + X.31 + X.32 + X.33 + X.34 + X.35 + X.36 + X.37 +
    X.38 + X.39 + X.40 + X.41 + X.42 + X.43 + X.44 + X.45 + X.46 +
    X.47 + X.48 + X.49 + X.50 + X.51 + X.52 + X.53 + X.54 + X.55 +
    X.56 + X.57 + X.58 + X.59 + X.60 + X.61 + X.62 + X.63 + X.64 +
    X.65 + X.66 + X.67 + X.68 + X.69 + X.70 + X.71 + X.72 + X.73 +
    X.74 + X.75 + X.76 + X.77 + X.78 + X.79 + X.80 + X.81 + X.82 +
    X.83 + X.84 + X.85 + X.86 + X.87 + X.88 + X.89 + X.90 + X.91 +
    X.92 + X.93 + X.94 + X.95 + X.96 + X.97 + X.98 + X.99 + X.100
  Res.Df
            RSS Df Sum of Sq
                                   F Pr(>F)
1
     999 973.65
     899 888.44 100
2
                       85.208 0.8622 0.825
```

R> summary(m.null)

```
Call:
```

```
lm(formula = y ~ 1, data = D)
```

Residuals:

```
Min 1Q Median 3Q Max -3.4585 -0.6809 0.0190 0.6365 2.9346
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|) (Intercept) 9.93704 0.03122 318.3 <2e-16
```

Residual standard error: 0.9872 on 999 degrees of freedom

Next, using the stepAIC() function in the MASS package (Venables and Ripley 2002), let us perform a forward stepwise regression to select a "best" model, starting with the null model, and using AIC as the model-selection criterion (see the help page for stepAIC() for details):¹³

Coefficients:

	${\tt Estimate}$	Std. Error	t value	Pr(> t)
(Intercept)	9.93716	0.03098	320.803	< 2e-16
X.99	-0.09103	0.03082	-2.953	0.00322
X.90	-0.08205	0.03135	-2.617	0.00901
X.87	-0.06942	0.03105	-2.235	0.02561
X.40	-0.04759	0.03076	-1.547	0.12211
X.65	-0.05523	0.03147	-1.755	0.07952
X.91	0.05245	0.03084	1.700	0.08937
X.53	-0.04921	0.03048	-1.615	0.10672
X.45	0.05543	0.03182	1.742	0.08183
X.31	0.04525	0.03108	1.456	0.14570
X.56	0.05433	0.03273	1.660	0.09723
X.61	-0.05085	0.03170	-1.604	0.10908
X.60	-0.05133	0.03194	-1.607	0.10832
X.46	0.05158	0.03272	1.576	0.11529
X.35	0.04696	0.03146	1.493	0.13584
X.92	0.04430	0.03100	1.429	0.15329

Residual standard error: 0.9725 on 984 degrees of freedom

 $^{^{13}}$ It's generally advantageous to start with the largest model, here the one with 100 predictors, and proceed by backward elimination. In this demonstration, however, where all of the β s are really 0, the selected model will be small, and so we proceed by forward selection from the null model to save computing time.

```
Multiple R-squared: 0.04419, Adjusted R-squared: 0.02962
F-statistic: 3.033 on 15 and 984 DF, p-value: 8.338e-05
R> mse(D$y, fitted(m.select))
[1] 0.9306254
attr(,"casewise loss")
[1] "(y - yhat)^2"
```

The resulting model has 15 predictors, a very modest $R^2 = .044$, but a small p-value for its omnibus F-test (which, of course, is entirely spurious because the same data were used to select and test the model). The MSE for the selected model is smaller than the true error variance $\sigma^2 = 1$, as is the estimated error variance for the selected model, $\hat{\sigma}^2 = 0.973^2 = 0.947$. If we cross-validate the selected model, we also obtain an optimistic estimate of its predictive

If we cross-validate the selected model, we also obtain an optimistic estimate of its predictive power (although the confidence interval for the bias-adjusted MSE includes 1):

```
R> cv(m.select, seed=2529)

R RNG seed set to 2529

10-Fold Cross Validation
method: Woodbury
criterion: mse
cross-validation criterion = 0.9593695
bias-adjusted cross-validation criterion = 0.9578478
95% CI for bias-adjusted CV criterion = (0.8766138, 1.039082)
full-sample criterion = 0.9306254
```

The cvSelect() function in the cv package allows us to cross-validate the whole model-selection procedure. The first argument to cvSelect() is a model-selection function capable of refitting the model with a fold omitted and returning a CV criterion. The selectStepAIC() function, also in cv and based on stepAIC(), is suitable for use with cvSelect():

The other arguments to cvSelect() are:

- data, the data set to which the model is fit;
- seed, an optional seed for R's pseudo-random-number generator; as for cv(), if the seed isn't supplied by the user, a seed is randomly selected and saved;
- additional arguments required by the model-selection function, here the starting model argument, the direction of model selection, and the scope of models considered (from the model with only a regression constant to the model with all 100 predictors).

By default, cvSelect() performs 10-fold CV, and produces an estimate of MSE for the model-selection procedure even *larger* than the true error variance, $\sigma^2 = 1$.

Also by default, when the number of folds is 10 or fewer, cvSelect() saves the coefficients of the selected models. In this example, the compareFolds() function reveals that the variables retained by the model-selection process in the several folds are quite different:

R> compareFolds(cv.select)

	(Intercep	ot) X.	87 X.	.90 X	.99	X.91	X.54	X.53	X.56
Fold 1	9.91	87 -0.06	315 -0.09	994 -0.0	942 0.	0512 0	.0516		
Fold 2	9.94	51 -0.07	45 -0.08	399 -0.0	614	0	.0587	0	.0673
Fold 3	9.94	23 -0.07	783 -0.07	718 -0.0	987 0.	0601		0	.0512
Fold 4	9.94 9.94	10 -0.08	360 -0.08	331 -0.0	867 0.	0570	-0.	0508	
Fold 5	9.94	21 -0.06	359 -0.08	349 -0.1	004 0.	0701 0	.0511 -0.	0487 0	. 0537
Fold 6	9.96	33 -0.07	733 -0.08	374 -0.0	960 0.	0555 0	.0629 -0.	0478	
Fold 7	9.92	79 -0.06	318 -0.09	960 -0.0	838 0.	0533	-0.	0464	
Fold 8	9.94	53 -0.06	310 -0.08	311 -0.0	818	0	.0497 -0.	0612 0	.0560
Fold 9	9.91	73 -0.06	63 -0.08	394 -0.1	100 0.	0504 0	.0524	0	.0747
Fold 10	9.94	49 -0.07	45 -0.09	906 -0.0	891 0.	0535 0	.0482 -0.	0583 0	.0642
									3 X.60
Fold 1		_	-0.0590			-0.045	6 0.0658	0.060	3
Fold 2					0.0607		0.0487	•	
	-0.0496								
Fold 4	-0.0597	0.0579 -	-0.0531		0.0519	-0.056	6		-0.0519
Fold 5				0.0587				0.052	7 -0.0603
Fold 6	-0.0596	0.0552		0.0474					
Fold 7		0.0572		0.0595					
Fold 8		0.0547 -	-0.0617	0.0453	0.0493	-0.061	3 0.0591	0.070	3 -0.0588
Fold 9	-0.0552	0.0573 -	-0.0635	0.0492		-0.051	3 0.0484		-0.0507
Fold 10	-0.0558								0
	X.61	X.8	X.28	X.29	X.31	Х.З	5 X.70	X.8	9 X.17
Fold 1	-0.0490		0.0616 -	-0.0537			0.0638	3	
					0.0568			0.052	3
									0.0527
Fold 5		0.0425			0.0672	0.061	3	0.049	3
Fold 6		0.0559	-	-0.0629	0.0498		0.0487	•	

Fold 7								0.0611	0.0472
Fold 8	-0.0719						0.0586		
Fold 9			0.0525						
Fold 10	-0.0580					0.0603			
	X.25	X.4	X.64	X.81	X.97	X.11	X.2	X.33	X.47
Fold 1					0.0604		0.0575		
Fold 2	0.0478		0.0532	0.0518					
Fold 3				0.0574				0.0473	
Fold 4			0.0628						
Fold 5	0.0518								
Fold 6						0.0521			
Fold 7		0.0550							
Fold 8									
Fold 9					0.0556				0.0447
Fold 10		0.0516							
	X.6	X.72	X.73	X.77	X.79	X.88			
Fold 1	0.0476								
Fold 2			0.0514						
Fold 3									
Fold 4					-0.0473				
Fold 5		0.0586				0.07			
Fold 6				-0.0489					
Fold 7									
Fold 8									
Fold 9									
Fold 10									

5.2. Mroz's logistic regression revisited

For a contrasting example we apply model selection to Mroz's logistic regression for married women's labor-force participation. First, recall the logistic regression model that we fit to the Mroz data:

```
hcyes
            0.111734 0.206040
                                  0.542 0.587618
            lwg
           -0.034446 0.008208 -4.196 2.71e-05
inc
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 1029.75 on 752
                                   degrees of freedom
Residual deviance: 905.27
                           on 745
                                   degrees of freedom
AIC: 921.27
Number of Fisher Scoring iterations: 4
Applying stepwise model selection Mroz's logistic regression, using BIC as the model-selection
criterion (via the argument k=log(nrow(Mroz)) to stepAIC()) selects 5 of the 7 original
predictors:
R> m.mroz.sel <- stepAIC(m.mroz, k=log(nrow(Mroz)),</pre>
                       trace=FALSE)
R> summary(m.mroz.sel)
Call:
glm(formula = lfp ~ k5 + age + wc + lwg + inc, family = binomial,
    data = Mroz)
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) 2.90193 0.54290 5.345 9.03e-08
           -1.43180 0.19320 -7.411 1.25e-13
k5
           -0.05853 0.01142 -5.127 2.94e-07
age
            0.87237
                       0.20639 4.227 2.37e-05
wcyes
            0.61568
                       0.15014 4.101 4.12e-05
lwg
           -0.03367
                       0.00780 -4.317 1.58e-05
inc
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 1029.75 on 752 degrees of freedom
Residual deviance: 906.46 on 747 degrees of freedom
AIC: 918.46
Number of Fisher Scoring iterations: 3
R> BayesRule(Mroz$lfp == "yes",
           predict(m.mroz.sel, type="response"))
[1] 0.3187251
attr(,"casewise loss")
```

[1] "y != round(yhat)"

Bayes rule applied to the selected model misclassifies 32% of the cases in the Mroz data. Cross-validating the selected model produces a similar, slightly larger, estimate of misclassification, about 33%:

```
R> cv(m.mroz.sel, criterion=BayesRule, seed=345266)

R RNG seed set to 345266

10-Fold Cross Validation
method: exact
criterion: BayesRule
cross-validation criterion = 0.3306773
bias-adjusted cross-validation criterion = 0.3333245
95% CI for bias-adjusted CV criterion = (0.2996998, 0.3669493)
full-sample criterion = 0.3187251
```

Is this estimate of predictive performance optimistic?

R> m.mroz.sel.cv <- cvSelect(selectStepAIC, Mroz,</pre>

We proceed to apply the model-selection procedure by cross-validation, producing more or less the same result:

```
# seed=6681,
# criterion=BayesRule,
# model=m.mroz,
# AIC=FALSE)

R RNG seed set to 6681

R> m.mroz.sel.cv

10-Fold Cross Validation
cross-validation criterion = 0.3306773
bias-adjusted cross-validation criterion = 0.3345203
95% CI for bias-adjusted CV criterion = (0.3008955, 0.368145)
full-sample criterion = 0.3187251
```

Setting AIC=FALSE in the call to cvSelect() uses the BIC rather than the AIC as the model-selection criterion. As it turns out, exactly the same predictors are selected when each of the 10 folds are omitted, and the several coefficient estimates are very similar, as we show using compareFolds():

R> compareFolds(m.mroz.sel.cv)

```
(Intercept) age inc k5 lwg wcyes
Fold 1 2.5014 -0.0454 -0.0388 -1.3613 0.5653 0.85
Fold 2 3.0789 -0.0659 -0.0306 -1.5335 0.6923 0.79
```

```
3.0141 -0.0595 -0.0305 -1.3994
Fold 3
                                              0.5428
                                                      0.86
Fold 4
             2.7251 -0.0543 -0.0354 -1.4474
                                              0.6298
                                                       1.09
Fold 5
             2.7617 -0.0566 -0.0320 -1.4752
                                              0.6324
                                                      0.74
Fold 6
             3.0234 -0.0621 -0.0348 -1.4537
                                              0.6618
                                                      0.94
Fold 7
             2.9615 -0.0600 -0.0351 -1.4127
                                              0.5835
                                                      0.97
             2.9598 -0.0603 -0.0329 -1.3865
Fold 8
                                              0.6210
                                                      0.69
Fold 9
             3.2481 -0.0650 -0.0381 -1.4138
                                              0.6093
                                                      0.94
Fold 10
             2.7724 -0.0569 -0.0295 -1.4503
                                              0.6347
                                                      0.85
```

In this example, therefore, we appear to obtain a realistic estimate of model performance directly from the selected model, because there is little added uncertainty induced by model selection.

5.3. Cross-validating choice of transformations in regression

The cv package also provides a cvSelect() procedure, selectTrans(), for choosing transformations of the predictors and the response in regression.

Some background: As Weisberg (2014, Sec. 8.2) explains, there are technical advantages to having (numeric) predictors in linear regression analysis that are themselves linearly related. If the predictors aren't linearly related, then the relationships between them can often be straightened by power transformations. Transformations can be selected after graphical examination of the data, or by analytic methods. Once the relationships between the predictors are linearized, it can be advantageous similarly to transform the response variable towards normality.

Selecting transformations analytically raises the possibility of automating the process, as would be required for cross-validation. One could, in principle, apply graphical methods to select transformations for each fold, but because a data analyst couldn't forget the choices made for previous folds, the process wouldn't really be applied independently to the folds.

To illustrate, we adapt an example appearing in several places in Fox and Weisberg (2019) (for example in Chapter 3 on transforming data), using data on the prestige and other characteristics of 102 Canadian occupations circa 1970. The data are in the Prestige data frame in the carData package:

```
R> data("Prestige", package="carData")
R> head(Prestige)
```

	education	income	women	prestige	census	type
<pre>gov.administrators</pre>	13.11	12351	11.16	68.8	1113	prof
general.managers	12.26	25879	4.02	69.1	1130	prof
accountants	12.77	9271	15.70	63.4	1171	prof
purchasing.officers	11.42	8865	9.11	56.8	1175	prof
chemists	14.62	8403	11.68	73.5	2111	prof
physicists	15.64	11030	5.13	77.6	2113	prof

R> summary(Prestige)

```
education
                                                        prestige
                      income
                                       women
Min.
       : 6.380
                         : 611
                                          : 0.000
                                                             :14.80
                  Min.
                                   Min.
                                                     Min.
1st Qu.: 8.445
                  1st Qu.: 4106
                                   1st Qu.: 3.592
                                                     1st Qu.:35.23
Median :10.540
                  Median: 5930
                                   Median :13.600
                                                     Median :43.60
Mean
       :10.738
                  Mean
                          : 6798
                                   Mean
                                          :28.979
                                                     Mean
                                                             :46.83
3rd Qu.:12.648
                  3rd Qu.: 8187
                                   3rd Qu.:52.203
                                                     3rd Qu.:59.27
Max.
       :15.970
                          :25879
                                   Max.
                                          :97.510
                                                     Max.
                                                             :87.20
                  Max.
    census
                  type
Min.
       :1113
                bc :44
1st Qu.:3120
                prof:31
Median:5135
                wc :23
Mean
       :5402
                NA's: 4
3rd Qu.:8312
       :9517
Max.
```

The variables in the Prestige data set are:

- education: average years of education for incumbents in the occupation, from the 1971 Canadian Census.
- income: average dollars of annual income for the occupation, from the Census.
- women: percentage of occupational incumbents who were women, also from the Census.
- prestige: the average prestige rating of the occupation on a 0–100 "thermometer" scale, in a Canadian social survey conducted around the same time.
- type, type of occupation, and census, the Census occupational code, which are not used in our example.

The object of a regression analysis for the Prestige data (and their original purpose) is to predict occupational prestige from the other variables in the data set.

A scatterplot matrix (using the scatterplotMatrix() function in the car package) of the numeric variables in the data reveals that the distributions of income and women are positively skewed, and that some of the relationships among the three predictors, and between the predictors and the response (i.e., prestige), are nonlinear:

The powerTransform() function in the car package transforms variables towards multivariate normality by a generalization of Box and Cox's maximum-likelihood-like approach (Box and Cox 1964). Several "families" of power transformations can be used, including the original Box-Cox family, simple powers (and roots), and two adaptations of the Box-Cox family to data that may include negative values and zeros: the Box-Cox-with-negatives family and the Yeo-Johnson family; see Weisberg (2014, Chap. 8), and Fox and Weisberg (2019, Chap. 3) for details. Because women has some zero values, we use the Yeo-Johnson family:

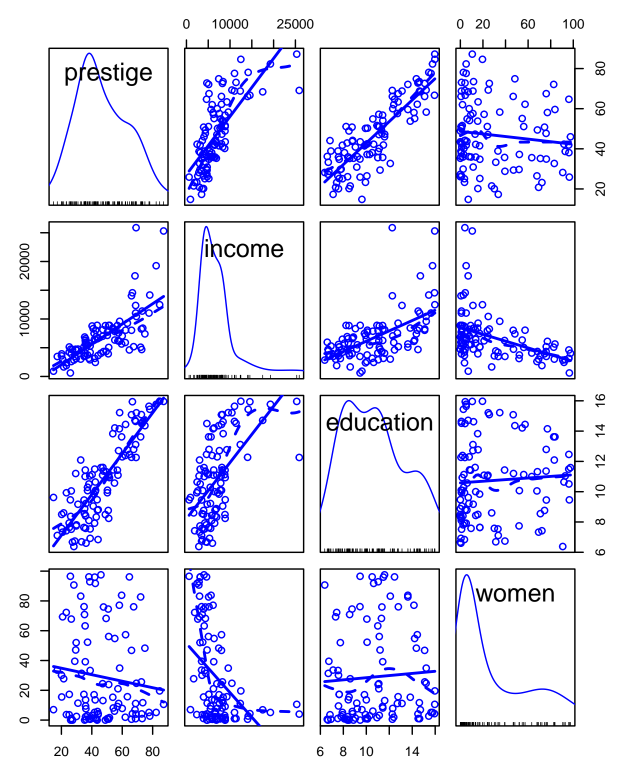


Figure 15: Scatterplot matrix for the 'Prestige' data.

yjPower Transformations to Multinormality

	Est Power	Rounded Pwr	Wald Lwr Bnd	Wald Upr Bnd
income	0.2678	0.33	0.1051	0.4304
${\tt education}$	0.5162	1.00	-0.2822	1.3145
women	0.1630	0.16	0.0112	0.3149

Likelihood ratio test that all transformation parameters are equal to 0

LRT df pval

LR test, lambda = $(0\ 0\ 0)\ 15.73879\ 3\ 0.0012827$

We thus have evidence of the desirability of transforming income (by the 1/3 power) and women (by the 0.16 power—which is close to the "0" power, i.e., the log transformation), but not education. Applying the "rounded" power transformations makes the predictors better-behaved:

```
R> P <- Prestige[, c("prestige", "income", "education", "women")]
R> (lambdas <- trans$roundlam)

income education    women
0.3300000 1.0000000 0.1630182

R> names(lambdas) <- c("income", "education", "women")
R> for (var in c("income", "education", "women")){
+ P[, var] <- yjPower(P[, var], lambda=lambdas[var])
+ }
R> summary(P)
```

```
prestige
                   income
                                 education
                                                   women
Min.
      :14.80
               Min.
                      :22.15
                               Min.
                                     : 6.380
                                               Min.
                                                      :0.000
1st Qu.:35.23
               1st Qu.:44.17
                               1st Qu.: 8.445
                                               1st Qu.:1.731
Median :43.60
               Median :50.26
                               Median :10.540
                                               Median :3.362
     :46.83
               Mean
                      :50.76
                                    :10.738
                                                     :3.502
Mean
                               Mean
                                               Mean
3rd Qu.:59.27
               3rd Qu.:56.24
                                               3rd Qu.:5.591
                               3rd Qu.:12.648
Max.
      :87.20
               Max.
                      :83.62
                               Max.
                                     :15.970
                                               Max.
                                                     :6.830
```

```
R> scatterplotMatrix(~ prestige + income + education + women,
+ data=P, smooth=list(spread=FALSE))
```

Comparing the MSE for the regressions with the original and transformed predictors shows a advantage to the latter:

```
R> m.pres <- lm(prestige ~ income + education + women, data=Prestige)
R> m.pres.trans <- lm(prestige ~ income + education + women, data=P)
R> mse(Prestige$prestige, fitted(m.pres))
```

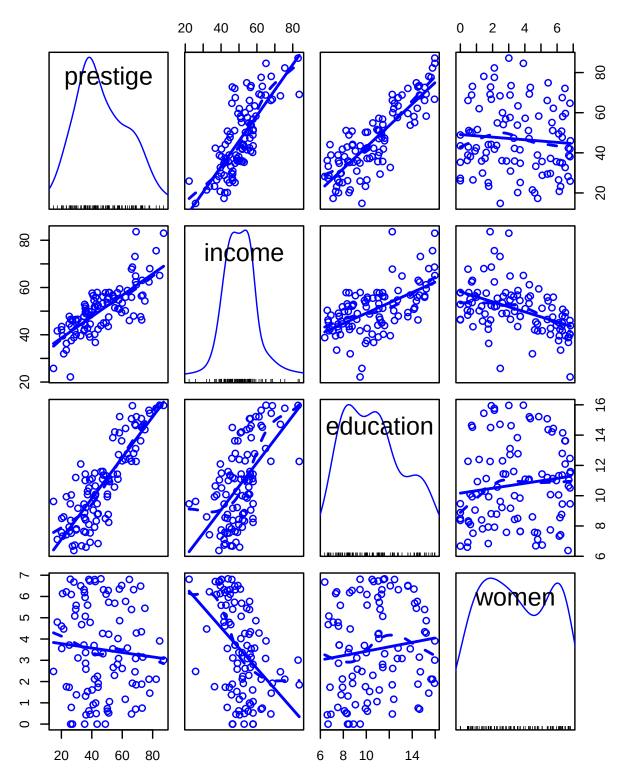


Figure 16: Scatterplot matrix for the 'Prestige' data with the predictors transformed.

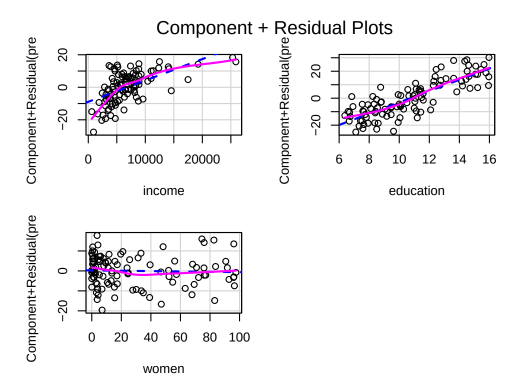


Figure 17: Component+residual plots for the 'Prestige' regression with the original predictors.

```
[1] 59.15265
attr(,"casewise loss")
[1] "(y - yhat)^2"

R> mse(P$prestige, fitted(m.pres.trans))
[1] 50.60016
attr(,"casewise loss")
[1] "(y - yhat)^2"
```

Similarly, component+residual plots for the two regressions, produced by the crPlots() function in the car package, suggest that the partial relationship of prestige to income is more nearly linear in the transformed data, but the transformation of women fails to capture what appears to be a slight quadratic partial relationship; the partial relationship of prestige to education is close to linear in both regressions:

```
R> crPlots(m.pres)
```

R> crPlots(m.pres.trans)

Having transformed the predictors towards multinormality, we now consider whether there's evidence for transforming the response (using powerTransform() for Box and Cox's original method), and we discover that there's not:

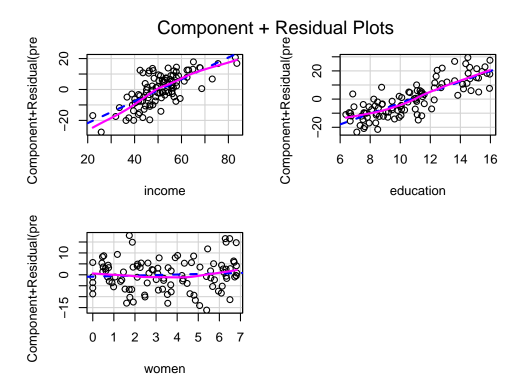


Figure 18: Component+residual plots for the 'Prestige' regression with transformed predictors.

R> summary(powerTransform(m.pres.trans))

bcPower Transformation to Normality
Est Power Rounded Pwr Wald Lwr Bnd Wald Upr Bnd
Y1 1.0194 1 0.6773 1.3615

Likelihood ratio test that transformation parameter is equal to 0 (log transformation)

LRT df pval LR test, lambda = (0) 32.2174 1 1.3785e-08

Likelihood ratio test that no transformation is needed

LRT df pval LR test, lambda = (1) 0.01238421 1 0.91139

The selectTrans() function in the cv package automates the process of selecting predictor and response transformations. The function takes a data set and "working" model as arguments, along with the candidate predictors and response for transformation, and the transformation family to employ. If the predictors argument is missing then only the response is transformed, and if the response argument is missing, only the supplied predictors are transformed. The default family for transforming the predictors is "bcPower"—the original Box-Cox family—as is the default family.y for transforming the response; here we specify family="yjPower because of the zeros in women. selectTrans() returns the result

of applying a lack-of-fit criterion to the model after the selected transformation is applied, with the default criterion=mse:

```
R> selectTrans(data=Prestige, model=m.pres,
              predictors=c("income", "education", "women"),
              response="prestige", family="yjPower")
[1] 50.60016
attr(,"casewise loss")
[1] "(y - yhat)^2"
selectTrans() also takes an optional indices argument, making it suitable for doing com-
putations on a subset of the data (i.e., a CV fold), and hence for use with cvSelect() (see
?selectTrans for details):
R> cvs <- cvSelect(selectTrans, data=Prestige, model=m.pres, seed=1463,
                  predictors=c("income", "education", "women"),
                  response="prestige",
                  family="yjPower")
R RNG seed set to 1463
R> cvs
10-Fold Cross Validation
cross-validation criterion = 54.4871
bias-adjusted cross-validation criterion = 54.30824
full-sample criterion = 50.60016
R> cv(m.pres, seed=1463) # untransformed model with same folds
R RNG seed set to 1463
10-Fold Cross Validation
method: Woodbury
criterion: mse
cross-validation criterion = 63.2926
bias-adjusted cross-validation criterion = 63.07251
full-sample criterion = 59.15265
R> compareFolds(cvs)
        lam.education lam.income lam.women lambda
Fold 1
                1.000
                            0.330
                                      0.330
```

1

1

0.169

0.330

Fold 2

Fold 3

1.000

1.000

0.330

0.330

Fold 4	1.000	0.330	0.330	1
Fold 5	1.000	0.330	0.000	1
Fold 6	1.000	0.330	0.330	1
Fold 7	1.000	0.330	0.330	1
Fold 8	1.000	0.330	0.000	1
Fold 9	1.000	0.330	0.000	1
Fold 10	1.000	0.330	0.000	1

The results suggest that the predictive power of the transformed regression is reliably greater than that of the untransformed regression (though in both case, the cross-validated MSE is considerably higher than the MSE computed for the whole data). Examining the selected transformations for each fold reveals that the predictor education and the response prestige are never transformed; that the 1/3 power is selected for income in all of the folds; and that the transformation selected for women varies narrowly across the folds between the 0th power (i.e., log) and the 1/3 power.

5.4. Selecting both transformations and predictors¹⁴

As we mentioned, Hastie et al. (2009, Sec. 7.10.2: "The Wrong and Right Way to Do Cross-validation") explain that honest cross-validation has to take account of model specification and selection. Statistical modeling is at least partly a craft, and one could imagine applying that craft to successive partial data sets, each with a fold removed. The resulting procedure would be tedious, though possibly worth the effort, but it would also be difficult to realize in practice: After all, we can hardly erase our memory of statistical modeling choices between analyzing partial data sets.

Alternatively, if we're able to automate the process of model selection, then we can more realistically apply CV mechanically. That's what we did in the preceding two sections, first for predictor selection and then for selection of transformations in regression. In this section, we consider the case where we both select variable transformations and then proceed to select predictors. It's insufficient to apply these steps sequentially, first, for example, using cvSelect() with selectTrans() and then with selectStepAIC(); rather we should apply the whole model-selection procedure with each fold omitted. The selectTransAndStepAIC() function, also supplied by the cv package, does exactly that.

To illustrate this process, we return to the Auto data set:

R> summary(Auto)

mpg	cylinders	displacement	horsepower	weight
Min. : 9.00	Min. :3.000	Min. : 68.0	Min. : 46.0	Min. :1613
1st Qu.:17.00	1st Qu.:4.000	1st Qu.:105.0	1st Qu.: 75.0	1st Qu.:2225
Median :22.75	Median :4.000	Median :151.0	Median : 93.5	Median :2804
Mean :23.45	Mean :5.472	Mean :194.4	Mean :104.5	Mean :2978
3rd Qu.:29.00	3rd Qu.:8.000	3rd Qu.:275.8	3rd Qu.:126.0	3rd Qu.:3615
Max. :46.60	Max. :8.000	Max. :455.0	Max. :230.0	Max. :5140

¹⁴The presentation in the section benefits from an email conversation with Bill Venables, who of course isn't responsible for the use to which we've put his insightful remarks.

name

: 5

:365

```
origin
  acceleration
                     year
 Min. : 8.00
                Min.
                       :70.00
                               Min.
                                     :1.000
                                               amc matador
 1st Qu.:13.78
                1st Qu.:73.00
                                1st Qu.:1.000
                                               ford pinto
 Median :15.50
               Median :76.00
                               Median :1.000
                                               toyota corolla
                     :75.98
 Mean
      :15.54
                Mean
                               Mean
                                     :1.577
                                               amc gremlin
 3rd Qu.:17.02
                3rd Qu.:79.00
                                3rd Qu.:2.000
                                               amc hornet
 Max.
       :24.80 Max. :82.00
                                Max. :3.000
                                               chevrolet chevette:
                                               (Other)
R> xtabs(~ year, data=Auto)
year
70 71 72 73 74 75 76 77 78 79 80 81 82
29 27 28 40 26 30 34 28 36 29 27 28 30
R> xtabs(~ origin, data=Auto)
origin
  1
     2
         3
245 68 79
R> xtabs(~ cylinders, data=Auto)
cylinders
```

3

4 199

4

5

6 3 83 103

We previously used the Auto here in a preliminary example where we employed CV to inform the selection of the order of a polynomial regression of mpg on horsepower. Here, we consider more generally the problem of predicting mpg from the other variables in the Auto data. We begin with a bit of data management, and then examine the pairwise relationships among the numeric variables in the data set:

```
R> Auto$cylinders <- factor(Auto$cylinders,</pre>
                            labels=c("3.4", "3.4", "5.6", "5.6", "8"))
R> Auto$year <- as.factor(Auto$year)</pre>
R> Auto$origin <- factor(Auto$origin,
                         labels=c("America", "Europe", "Japan"))
R> rownames(Auto) <- make.names(Auto$name, unique=TRUE)</pre>
R> Auto$name <- NULL
R> scatterplotMatrix(~ mpg + displacement + horsepower + weight + acceleration,
                     smooth=list(spread=FALSE), data=Auto)
```

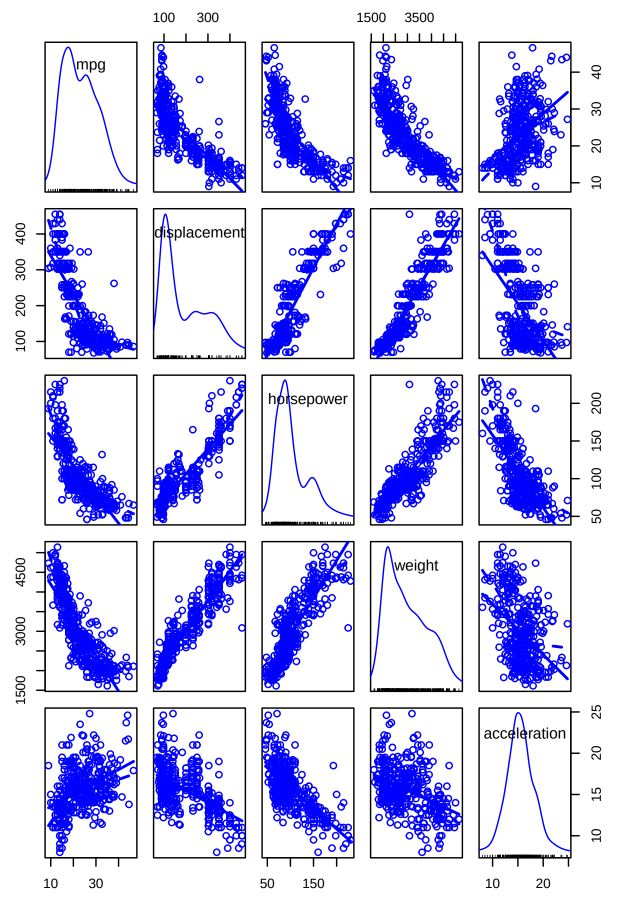


Figure 19: Scatterplot matrix for the numeric variables in the 'Auto' data

A comment before we proceed: origin is clearly categorical and so converting it to a factor is natural, but we could imagine treating cylinders and year as numeric predictors. There are, however, only 5 distinct values of cylinders (ranging from 3 to 8), but cars with 3 or 5 cylinders are rare. and none of the cars has 7 cylinders. There are similarly only 13 distinct years between 1970 and 1982 in the data, and the relationship between mpg and year is difficult to characterize. It's apparent that most these variables are positively skewed and that many of the pairwise relationships among them are nonlinear.

We begin with a "working model" that specifies linear partial relationships of the response to the numeric predictors:

1.5251 10.9504

```
R> m.auto <- lm(mpg ~ ., data = Auto)
R> summary(m.auto)

Call:
lm(formula = mpg ~ ., data = Auto)

Residuals:
    Min    1Q Median    3Q Max
```

Coefficients:

-9.0064 -1.7450 -0.0917

```
Estimate Std. Error t value Pr(>|t|)
             37.0341323
                          1.9693933
                                     18.805
                                             < 2e-16
(Intercept)
cylinders5.6 -2.6029412
                          0.6552000
                                     -3.973 8.54e-05
cylinders8
                                     -0.497 0.619335
             -0.5824578
                          1.1714516
displacement
              0.0174253
                          0.0067340
                                      2.588 0.010043
horsepower
             -0.0413534
                          0.0133786
                                     -3.091 0.002145
weight
             -0.0055479
                          0.0006323
                                     -8.774
                                            < 2e-16
acceleration 0.0615272
                          0.0883132
                                      0.697 0.486431
year71
              0.9680584
                          0.8373899
                                      1.156 0.248408
                          0.8251155
                                     -0.729 0.466517
year72
             -0.6014345
year73
             -0.6876890
                          0.7402723
                                     -0.929 0.353510
                          0.8765000
                                      1.569 0.117408
              1.3755758
year74
              0.9299288
                          0.8590716
                                      1.082 0.279742
year75
              1.5598929
                          0.8225051
                                      1.897 0.058669
year76
year77
              2.9094161
                          0.8417285
                                      3.456 0.000611
              3.1751976
                          0.7989396
                                      3.974 8.48e-05
year78
              5.0192987
                          0.8457587
                                      5.935 6.76e-09
year79
year80
              9.0997634
                          0.8972933
                                     10.141
                                              < 2e-16
year81
              6.6886597
                          0.8852181
                                      7.556 3.28e-13
year82
              8.0711248
                          0.8706683
                                      9.270
                                             < 2e-16
```

¹⁵Of course, making the decision to treat **year** as a factor on this basis could be construed as cheating in the current context, which illustrates the difficulty of automating the whole model-selection process. It's rarely desirable, in our opinion, to forgo exploration of the data to ensure the purity of model validation. We believe, however, that it's still useful to automate as much of the process as we can to obtain a more realistic, if still biased, estimate of the predictive power of a model.

```
originEurope 2.0466642 0.5171236 3.958 9.07e-05
originJapan 2.1448874 0.5077169 4.225 3.02e-05
```

Residual standard error: 2.924 on 371 degrees of freedom Multiple R-squared: 0.8668, Adjusted R-squared: 0.8596 F-statistic: 120.7 on 20 and 371 DF, p-value: < 2.2e-16

R> Anova(m.auto)

Anova Table (Type II tests)

Response: mpg

```
Sum Sq Df F value
                                Pr(>F)
cylinders
             292.3
                    2 17.0915 7.935e-08
displacement
            57.3 1 6.6959 0.010043
horsepower
             81.7 1 9.5544 0.002145
weight
             658.3
                    1 76.9801 < 2.2e-16
acceleration 4.2 1 0.4854 0.486431
year
            3016.8 12 29.3987 < 2.2e-16
origin
            190.3 2 11.1287 2.024e-05
            3172.5 371
Residuals
```

R> crPlots(m.auto)

The component+residual plots, created with the crPlots() function in the previously loaded car package, clearly reveal the inadequacy of the model.

We proceed to transform the numeric predictors towards multi-normality:

```
R> num.predictors <- c("displacement", "horsepower", "weight", "acceleration")
R> tr.x <- powerTransform(Auto[, num.predictors])
R> summary(tr.x)
```

bcPower Transformations to Multinormality

	Est Power	Rounded	Pwr	Wald	Lwr	Bnd	Wald	Upr	Bnd
displacement	-0.0509		0		-0.2	2082		0.3	1065
horsepower	-0.1249		0		-0.2	2693		0.0	0194
weight	-0.0870		0		-0.2	2948		0.3	1208
acceleration	0.3061		0		-0.0	0255		0.6	6376

Likelihood ratio test that transformation parameters are equal to 0 (all log transformations)

```
LRT df pval LR test, lambda = (0\ 0\ 0\ 0)\ 4.872911\ 4\ 0.30059
```

Likelihood ratio test that no transformations are needed $$\operatorname{LRT}$\ df $$\operatorname{pval}$$

```
LR test, lambda = (1 1 1 1) 390.0777 4 < 2.22e-16
```

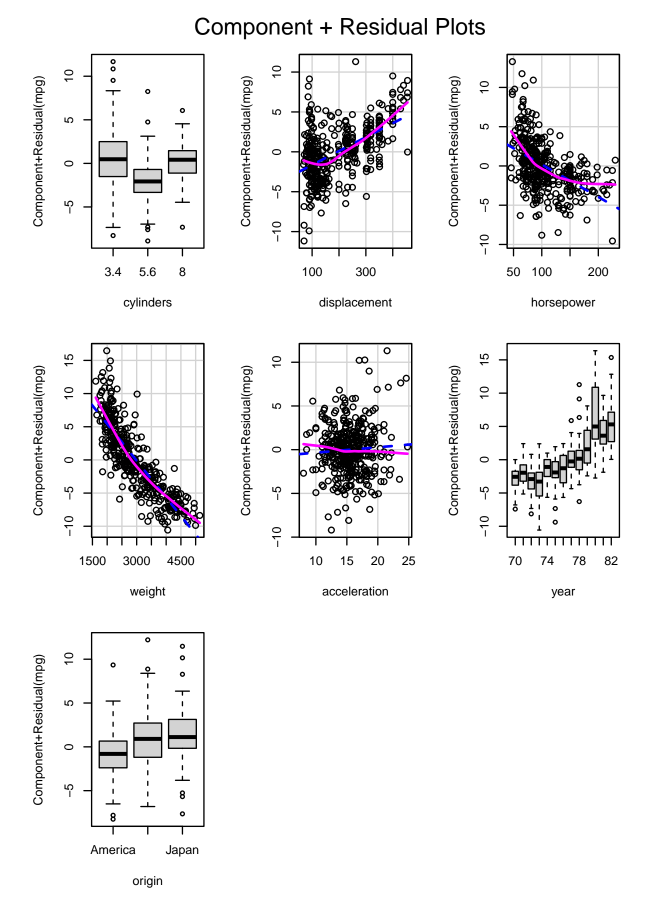


Figure 20: Component+residual plots for the working model fit to the 'Auto' data

We then apply the (rounded) transformations—all, as it turns out, logs—to the data and re-estimate the model:

```
R> A <- Auto
R> powers <- tr.x$roundlam
R> for (pred in num.predictors){
    A[, pred] <- bcPower(A[, pred], lambda=powers[pred])</pre>
+ }
R> head(A)
                           mpg cylinders displacement horsepower
chevrolet.chevelle.malibu 18
                                       8
                                             5.726848
                                                         4.867534 8.161660
buick.skylark.320
                            15
                                       8
                                             5.857933
                                                         5.105945 8.214194
plymouth.satellite
                            18
                                       8
                                             5.762051
                                                         5.010635 8.142063
amc.rebel.sst
                                       8
                                                         5.010635 8.141190
                            16
                                             5.717028
ford.torino
                            17
                                       8
                                             5.710427
                                                         4.941642 8.145840
ford.galaxie.500
                            15
                                       8
                                             6.061457
                                                         5.288267 8.375860
                           acceleration year origin
chevrolet.chevelle.malibu
                               2.484907
                                          70 America
buick.skylark.320
                                          70 America
                               2.442347
plymouth.satellite
                               2.397895
                                          70 America
amc.rebel.sst
                               2.484907
                                          70 America
                                          70 America
ford.torino
                               2.351375
ford.galaxie.500
                               2.302585
                                         70 America
R> m <- update(m.auto, data=A)</pre>
Finally, we perform Box-Cox regression to transform the response (also obtaining a log trans-
```

formation):

```
R> summary(powerTransform(m))
```

```
bcPower Transformation to Normality
   Est Power Rounded Pwr Wald Lwr Bnd Wald Upr Bnd
Y1
      0.0024
                       0
                              -0.1607
                                            0.1654
Likelihood ratio test that transformation parameter is equal to 0
 (log transformation)
                               LRT df
                                         pval
LR test, lambda = (0) 0.0008015428 1 0.97741
Likelihood ratio test that no transformation is needed
                           LRT df
LR test, lambda = (1) 124.1307 1 < 2.22e-16
```

```
R> m <- update(m, log(mpg) ~ .)</pre>
R> summary(m)
```

Call:

```
lm(formula = log(mpg) ~ cylinders + displacement + horsepower +
    weight + acceleration + year + origin, data = A)
```

Residuals:

Coefficients:

	${\tt Estimate}$	Std. Error	t value	Pr(> t)
(Intercept)	8.89652	0.35822	24.835	< 2e-16
${\tt cylinders 5.6}$	-0.06355	0.02574	-2.469	0.0140
cylinders8	-0.07691	0.03900	-1.972	0.0493
${\tt displacement}$	0.02799	0.05146	0.544	0.5868
horsepower	-0.29010	0.05631	-5.152	4.20e-07
weight	-0.54274	0.08193	-6.624	1.23e-10
${\tt acceleration}$	-0.14214	0.05630	-2.525	0.0120
year71	0.02505	0.02891	0.866	0.3869
year72	-0.01680	0.02894	-0.580	0.5620
year73	-0.04257	0.02602	-1.636	0.1026
year74	0.04932	0.03041	1.622	0.1056
year75	0.04715	0.02959	1.594	0.1118
year76	0.07087	0.02845	2.491	0.0132
year77	0.13241	0.02927	4.523	8.21e-06
year78	0.14472	0.02777	5.211	3.13e-07
year79	0.23354	0.02921	7.994	1.67e-14
year80	0.32381	0.03170	10.216	< 2e-16
year81	0.25655	0.03094	8.291	2.10e-15
year82	0.30756	0.03036	10.131	< 2e-16
$\verb"originEurope"$	0.04921	0.01955	2.518	0.0122
originJapan	0.04409	0.01947	2.265	0.0241

Residual standard error: 0.1043 on 371 degrees of freedom Multiple R-squared: 0.9108, Adjusted R-squared: 0.906 F-statistic: 189.4 on 20 and 371 DF, p-value: < 2.2e-16

R> Anova(m)

Anova Table (Type II tests)

Response: log(mpg)

```
Sum Sq Df F value Pr(>F)
cylinders 0.0663 2 3.0521 0.04845
displacement 0.0032 1 0.2959 0.58679
horsepower 0.2885 1 26.5420 4.198e-07
weight 0.4769 1 43.8805 1.229e-10
```

```
acceleration 0.0693 1 6.3745 0.01199
year 4.4521 12 34.1339 < 2.2e-16
origin 0.0807 2 3.7128 0.02532
Residuals 4.0325 371
```

The transformed numeric variables are much better-behaved:

```
R> scatterplotMatrix(~ log(mpg) + displacement + horsepower + weight
+ acceleration,
+ smooth=list(spread=FALSE), data=A)
```

And the partial relationships in the model fit to the transformed data are much more nearly linear:

```
R> crPlots(m)
```

Having transformed both the numeric predictors and the response, we proceed to use the stepAIC() function in the MASS package to perform predictor selection, employing the BIC model-selection criterion (by setting the k argument of stepAIC() to log(n)):

```
R> m.step <- stepAIC(m, k=log(nrow(A)), trace=FALSE)
R> summary(m.step)
```

Call:

```
lm(formula = log(mpg) ~ horsepower + weight + acceleration +
    year + origin, data = A)
```

Residuals:

```
Min 1Q Median 3Q Max -0.35230 -0.05682 0.00677 0.06741 0.35861
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
             9.434594
                        0.261529 36.075 < 2e-16
            -0.276254
horsepower
                        0.056143 -4.921 1.30e-06
weight
            -0.609071
                        0.056003 -10.876 < 2e-16
acceleration -0.131380
                        0.053195 -2.470 0.01397
                                  0.967 0.33412
year71
             0.027984
                        0.028936
            -0.007111
year72
                        0.028446 -0.250 0.80274
            -0.039529
                        0.026014 -1.520 0.12947
year73
year74
             0.052752
                        0.029986
                                 1.759 0.07936
year75
             0.053199
                        0.029280
                                  1.817 0.07004
                                  2.634 0.00878
             0.074317
                        0.028212
year76
                                  4.777 2.56e-06
year77
             0.137931
                        0.028875
             0.145876
                        0.027529
                                  5.299 1.99e-07
year78
```

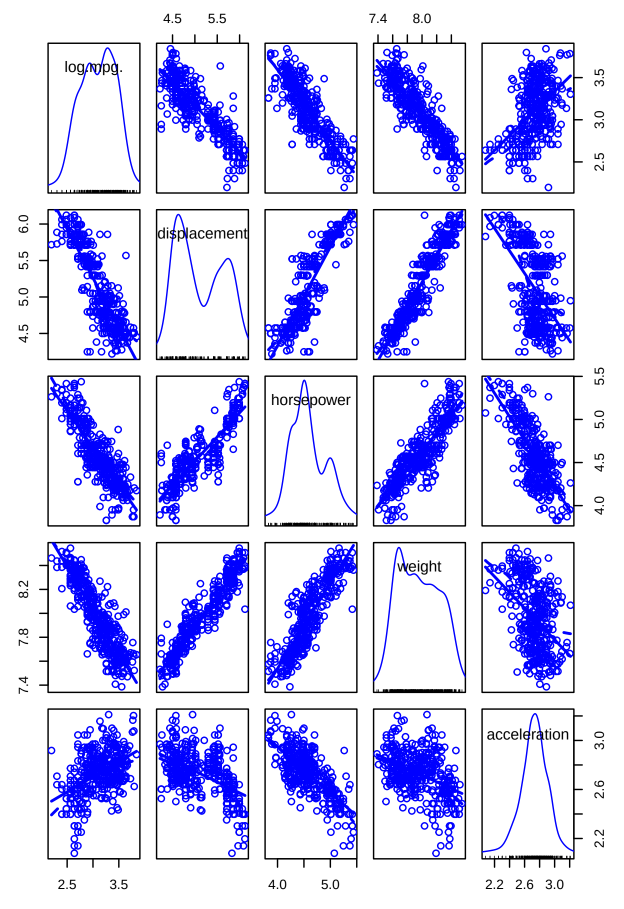


Figure 21: Scatterplot matrix for the transformed numeric variables in the 'Auto' data

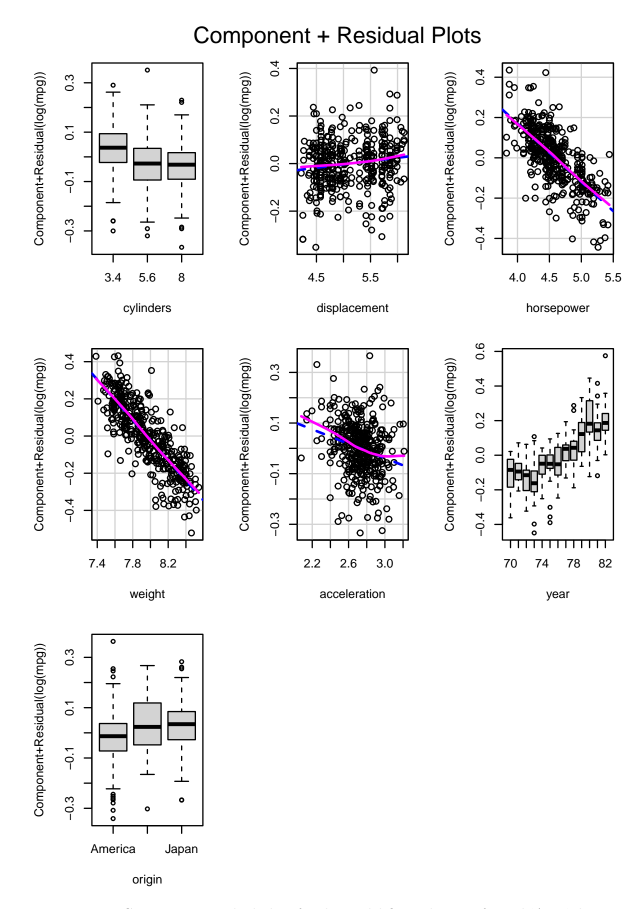


Figure 22: Component+residual plots for the model fit to the transformed 'Auto' data

```
0.029080
                                    8.117 6.99e-15
year79
              0.236036
              0.335274
                         0.031148 10.764 < 2e-16
year80
year81
              0.262872
                         0.030555
                                    8.603 < 2e-16
year82
              0.323391
                         0.029608
                                   10.922 < 2e-16
originEurope
             0.055818
                         0.016785
                                    3.326 0.00097
                                    2.492 0.01314
originJapan
              0.043554
                         0.017479
```

Residual standard error: 0.1049 on 374 degrees of freedom Multiple R-squared: 0.909, Adjusted R-squared: 0.9049 F-statistic: 219.8 on 17 and 374 DF, p-value: < 2.2e-16

R> Anova(m.step)

Anova Table (Type II tests)

```
Response: log(mpg)
```

```
Sum Sq Df
                         F value
                                    Pr(>F)
                      1 24.2120 1.296e-06
horsepower
             0.2663
             1.3010
                      1 118.2805 < 2.2e-16
weight
acceleration 0.0671
                      1
                          6.0998 0.013965
                         36.0549 < 2.2e-16
year
             4.7589
                     12
origin
             0.1366
                      2
                          6.2090 0.002225
Residuals
             4.1137 374
```

The selected model includes three of the numeric predictors, horsepower, weight, and acceleration, along with the factors year and origin. We can calculate the MSE for this model, but we expect that the result will be optimistic because we used the whole data to help specify the model

```
R> mse(Auto$mpg, exp(fitted(m.step)))
```

```
[1] 6.512144
attr(,"casewise loss")
[1] "(y - yhat)^2"
```

This is considerably smaller than the MSE for the original working model:

```
R> mse(Auto$mpg, fitted(m.auto))
```

```
[1] 8.093171
attr(,"casewise loss")
[1] "(y - yhat)^2"
```

A perhaps subtle point is that we compute the MSE for the selected model on the original mpg response scale rather than the log scale, so as to make the selected model comparable to the working model. That's slightly uncomfortable given the skewed distribution of mpg. An alternative is to use the median absolute error instead of the mean-squared error, employing the medAbsErr() function from the cv package:

```
R> medAbsErr(Auto$mpg, exp(fitted(m.step)))
[1] 1.339604
```

R> medAbsErr(Auto\$mpg, fitted(m.auto))

[1] 1.666121

Now let's use cvSelect() with selectTransAndStepAIC() to automate and cross-validate the whole model-specification process:

R> num.predictors

```
[1] "displacement" "horsepower" "weight" "acceleration"
```

R RNG seed set to 76692

R> cvs

10-Fold Cross Validation cross-validation criterion = 1.495075 full-sample criterion = 1.339604

R> compareFolds(cvs)

	(Intercept)	horsepower lar	m.acceleration	lam.displacement	lam.horsepower
Fold 1	9.71384	-0.17408	0.50000	0.00000	0.00000
Fold 2	9.21713	-0.31480	0.00000	0.00000	0.00000
Fold 3	9.61824	-0.19248	0.00000	0.00000	0.00000
Fold 4	8.69910	-0.25523	0.50000	0.00000	0.00000
Fold 5	9.14403	-0.14934	0.00000	0.00000	0.00000
Fold 6	9.63481	-0.16739	0.50000	0.00000	0.00000
Fold 7	9.98933	-0.36847	0.00000	0.00000	-0.15447
Fold 8	9.06301	-0.29721	0.00000	0.00000	0.00000
Fold 9	8.88315	-0.22684	0.00000	0.00000	0.00000
Fold 10	9.61727	-0.17086	0.00000	0.00000	0.00000
	lam.weight	lambda weig	ght year71	year72 year73	year74
Fold 1	0.00000	0.00000 -0.746	636 0.03764 -0	.00327 -0.02477	0.05606
Fold 2	0.00000	0.00000 -0.47	728 0.02173 -0	.01488 -0.03770	0.04312
Fold 3	0.00000	0.00000 -0.720	0.01128 -0	.02569 -0.03872	0.05187
Fold 4	0.00000	0.00000 -0.538	846 0.02153 -0	.02922 -0.05181	0.04136
Fold 5	0.00000	0.00000 -0.690	0.02531 -0	.01062 -0.04625	0.05039

```
Fold 6
                     0.00000 - 0.74049
                                        0.02456 0.00759 -0.03412
           0.00000
                                                                    0.06266
Fold 7
           0.00000
                     0.00000 -0.72843
                                        0.02532 -0.01271 -0.04144
                                                                    0.04568
Fold 8
           0.00000
                     0.00000 - 0.46392
                                        0.02702 -0.02041 -0.05605
                                                                    0.04437
Fold 9
           0.00000
                     0.00000 - 0.47136
                                        0.00860 -0.03620 -0.04835
                                                                    0.01906
                                        0.02937 -0.00899 -0.03814
Fold 10
           0.00000 0.00000 -0.73550
                                                                    0.05408
                                                year79
                                                          year80
                                                                   year81
          year75
                    year76
                             year77
                                       year78
                                                                             year82
Fold 1
         0.07080
                   0.07250
                            0.14420
                                      0.14281
                                               0.23266
                                                         0.35127
                                                                  0.25635
                                                                            0.30546
Fold 2
         0.04031
                                      0.14917
                                               0.21871
                                                                  0.26196
                                                                            0.30943
                   0.06718
                            0.13094
                                                         0.33192
Fold 3
         0.03837
                   0.06399
                            0.11593
                                      0.12601
                                               0.20499
                                                         0.32821
                                                                  0.24478
                                                                            0.29204
Fold 4
         0.04072
                   0.05537
                            0.12292
                                      0.14083
                                               0.22878
                                                         0.32947
                                                                  0.25140
                                                                            0.27248
Fold 5
         0.05596
                   0.07044
                            0.13356
                                      0.14724
                                               0.24675
                                                         0.33331
                                                                  0.26938
                                                                            0.32594
Fold 6
         0.06940
                  0.07769
                            0.14211
                                      0.14647
                                               0.23532
                                                         0.34761
                                                                  0.26737
                                                                            0.33062
Fold 7
         0.03614
                  0.07385
                            0.12976
                                      0.14040
                                               0.23976
                                                                  0.27652
                                                                            0.30659
                                                         0.33998
Fold 8
         0.06573
                            0.13158
                                      0.13987
                                               0.23011
                                                         0.32880
                  0.08135
                                                                  0.25886
                                                                            0.30538
Fold 9
         0.03018
                  0.05846
                            0.10536
                                      0.11722
                                               0.20665
                                                         0.31533
                                                                  0.23352
                                                                            0.29375
Fold 10
         0.04881
                   0.07862
                            0.14101
                                      0.14313
                                               0.23258
                                                         0.35649
                                                                  0.26214
                                                                            0.32421
        acceleration displacement cylinders5.6 cylinders8 originEurope
Fold 1
Fold 2
            -0.18909
                          -0.09197
Fold 3
                                        -0.09080
Fold 4
            -0.03484
                                                   -0.10909
Fold 5
                                                                  0.06261
Fold 6
Fold 7
Fold 8
            -0.17676
                          -0.10542
Fold 9
            -0.14514
                          -0.13452
Fold 10
        originJapan
Fold 1
Fold 2
Fold 3
Fold 4
Fold 5
               0.04
Fold 6
Fold 7
Fold 8
Fold 9
Fold 10
```

Here, as for selectTrans(), the predictors and response arguments specify candidate variables for transformation, and AIC=FALSE uses the BIC for model selection. The starting model, m.auto, is the working model fit to the Auto data. The CV criterion isn't bias-adjusted because median absolute error isn't a mean of casewise error components.

Some noteworthy points:

• selectTransStepAIC() automatically computes CV cost criteria, here the median absolute error, on the untransformed response scale.

- The estimate of the median absolute error that we obtain by cross-validating the whole model-specification process is a little larger than the median absolute error computed for the model we fit to the Auto data separately selecting transformations of the predictors and the response and then selecting predictors for the whole data set.
- When we look at the transformations and predictors selected with each of the 10 folds omitted (i.e., the output of compareFolds()), we see that there is little uncertainty in choosing variable transformations (the lam.*s for the xs and lambda for y in the output), but considerably more uncertainty in subsequently selecting predictors: horsepower, weight, and year are always included among the selected predictors; acceleration and displacement are each included respectively in 4 and 3 of 10 selected models; and cylinders and origin are each included in only 1 of 10 models. Recall that when we selected predictors for the full data, we obtained a model with horsepower, weight, acceleration, year, and origin.

6. Parallel computations

The CV functions in the **cv** package are all capable of performing parallel computations by setting the **ncores** argument (specifying the number of computer cores to be used) to a number > 1 (which is the default). Parallel computation can be advantageous for large problems, reducing the execution time of the program.

To illustrate, let's time model selection in Mroz's logistic regression, repeating the computation as performed previously and then doing it in parallel using 2 cores:

```
R> system.time(m.mroz.sel.cv <- cvSelect(selectStepAIC, Mroz,</pre>
                              seed=6681,
                              criterion=BayesRule,
+
                              model=m.mroz,
                              AIC=FALSE))
R RNG seed set to 6681
   user
         system elapsed
  0.249
          0.004
                   0.253
R> system.time(m.mroz.sel.cv.p <- cvSelect(selectStepAIC, Mroz,</pre>
                              seed=6681,
                              criterion=BayesRule,
                              model=m.mroz,
                              AIC=FALSE,
                              ncores=2))
R RNG seed set to 6681
         system elapsed
  0.035
          0.004
                   0.949
```

```
R> all.equal(m.mroz.sel.cv, m.mroz.sel.cv.p)
```

[1] TRUE

In this small problem, the parallel computation is actually *slower*, because there is an overhead cost to parallelization, but we can see that it produces the same result as before.

7. Extending the cv package

The cv package is designed to be extensible in several directions. In this vignette, we discuss three kinds of extensions, ordered by increasing general complexity: (1) adding a cross-validation cost criterion; (2) adding a model class that's not directly accommodated by the cv() default method or by another directly inherited method, with separate consideration of mixed-effects models; and (3) adding a new model-selection procedure suitable for use with selectModel().

7.1. Adding a cost criterion

A cost criterion suitable for use with cv() or cvSelect() should take two arguments, y (the observed response vector) and yhat (a vector of fitted or predicted response values), and return a numeric index of lack of fit. The cv package supplies several such criteria: mse(y, yhat), which returns the mean-squared prediction error for a numeric response; rmse(y, yhat), which returns the (square-)root mean-squared error; medAbsErr(y, yhat), which returns the median absolute error; and BayesRule(y, yhat) (and its non-error-checking version, BayesRule2(y, yhat)), suitable for use with a binary regression model, where y is the binary response coded 0 for a "failure" or 1 for a "success"; where yhat is the predicted probability of success; and where the proportion of incorrectly classified cases is returned.

To illustrate using a different prediction cost criterion, we'll base a cost criterion on the area under the receiver operating characteristic ("ROC") curve for a logistic regression. The ROC curve is a graphical representation of the classification power of a binary regression model, and the area under the ROC curve ("AUC"), which varies from 0 to 1, is a common summary measure based on the ROC (see "Receiver operating characteristic" 2023). The Metrics package (Hamner and Frasco 2018) includes a variety of measures useful for model selection, including an auc() function. We convert the AUC into a cost measure by taking its complement:

```
R> AUCcomp <- function(y, yhat) 1 - Metrics::auc(y, yhat)
```

We then apply AUCcomp() to the Mroz logistic regression discussed in the main cv package vignette, which we reproduce here, using the Mroz data frame from the carData package (Fox and Weisberg 2019):

```
R> data("Mroz", package="carData")
R> m.mroz <- glm(lfp ~ ., data=Mroz, family=binomial)
R> summary(m.mroz)
```

```
Call:
glm(formula = lfp ~ ., family = binomial, data = Mroz)
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
(Intercept) 3.182140
                        0.644375 4.938 7.88e-07
k5
            -1.462913
                        0.197001 -7.426 1.12e-13
k618
            -0.064571
                        0.068001 -0.950 0.342337
            -0.062871
                        0.012783 -4.918 8.73e-07
age
             0.807274
                                   3.510 0.000448
wcyes
                        0.229980
hcyes
             0.111734
                        0.206040
                                   0.542 0.587618
                        0.150818 4.009 6.09e-05
             0.604693
lwg
            -0.034446
                        0.008208 -4.196 2.71e-05
inc
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 1029.75 on 752 degrees of freedom
Residual deviance: 905.27 on 745 degrees of freedom
AIC: 921.27
Number of Fisher Scoring iterations: 4
R> AUCcomp(with(Mroz, as.numeric(lfp == "yes")), fitted(m.mroz))
[1] 0.2636161
Cross-validating this cost measure is straightforward:
R> library("cv")
R> cv(m.mroz, criterion=AUCcomp, seed=3639)
R RNG seed set to 3639
10-Fold Cross Validation
method: exact
criterion: AUCcomp
cross-validation criterion = 0.2747088
full-sample criterion = 0.2636161
```

As expected, the cross-validated complement to the AUC is somewhat less optimistic than the criterion computed from the model fit to the whole data set.

As we explain in the vignette "Cross-validation of regression models," the cv() function differentiates between CV criteria that are averages of casewise components and criteria that are not. Computation of bias corrections and confidence intervals is limited to the former. We show in the appendix to this vignette that the AUC, and hence its complement, cannot be expressed as averages of casewise components.

cv() looks for a "casewise loss" attribute of the value returned by a CV criterion function. If this attribute exists, then the criterion is treated as the mean of casewise components, and cv() uses the unexported function getLossFn() to construct a function that returns the casewise components of the criterion.

We illustrate with the mse():

```
R> mse
[1] 23.94366 18.98477 18.94499 18.87633 18.42697 18.24065 18.07817 18.06613
[9] 18.02697 18.00953

R> cv:::getLossFn(mse(rnorm(100), rnorm(100)))

function (y, yhat)
{
        (y - yhat)^2
}

<environment: 0x12bdf3978>
```

For this scheme to work, the "casewise loss" attribute must be a character string (or vector of character strings), here "(y - yhat)^2", that evaluates to an expression that is a function of y and yhat, and that computes the vector of casewise components of the CV criterion.

7.2. Adding a model class not covered by the default cv() method

Independently sampled cases

Suppose that we want to cross-validate a multinomial logistic regression model fit by the multinom() function in the **nnet** package (Venables and Ripley 2002). We borrow an example from Fox (2016, Sec. 14.2.1), with data from the British Election Panel Study on vote choice in the 2001 British election. Data for the example are in the BEPS data frame in the **carData** package:

```
R> data("BEPS", package="carData")
R> head(BEPS)
```

		vo	te age	economic.cond.nation	nal e	economic.cond.household	Blair
1	Liberal	Democr	rat 43		3	3	4
2		Labo	our 36		4	4	4
3		Labo	our 35		4	4	5
4		Labo	our 24		4	2	2
5		Labo	our 41		2	2	1
6		Labo	our 47		3	4	4
	Hague K	ennedy	Europe	political.knowledge	gend	ler	
1	1	4	2	2	fema	ale	
2	4	4	5	2	ma	ale	

${\tt male}$	2	3	3	2	3
${\tt female}$	0	4	3	1	4
${\tt male}$	2	6	4	1	5
male	2	4	2	4	6

The polytomous (multi-category) response variable is vote, a factor with levels "Conservative", "Labour", and "Liberal Democrat". The predictors of vote are:

• age, in years;

gender

Blair

economic.cond.national
economic.cond.household

- econ.cond.national and econ.cond.household, the respondent's ratings of the state of the economy, on 1 to 5 scales.
- Blair, Hague, and Kennedy, ratings of the leaders of the Labour, Conservative, and Liberal Democratic parties, on 1 to 5 scales.
- Europe, an 11-point scale on attitude towards European integration, with high scores representing "Euro-skepticism."
- political.knowledge, knowledge of the parties' positions on European integration, with scores from 0 to 3.
- gender, "female" or "male".

The model fit to the data includes an interaction between Europe and political.knowledge; the other predictors enter the model additively:

```
R> library("nnet")
R> m.beps <- multinom(vote ~ age + gender + economic.cond.national +</pre>
                         economic.cond.household + Blair + Hague + Kennedy +
                         Europe*political.knowledge, data=BEPS)
# weights: 36 (22 variable)
initial value 1675.383740
iter 10 value 1240.047788
iter 20 value 1163.199642
iter 30 value 1116.519687
final value 1116.519666
converged
R> car::Anova(m.beps)
Analysis of Deviance Table (Type II tests)
Response: vote
                           LR Chisq Df Pr(>Chisq)
age
                             13.872 2 0.0009721
```

0.453 2 0.7972568

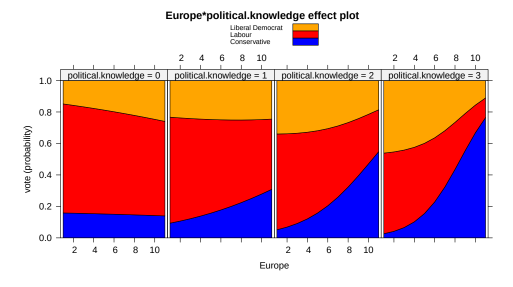
5.652 2 0.0592570 135.369 2 < 2.2e-16

30.604 2 2.262e-07

Hague	166.770	2	< 2.2e-16
Kennedy	68.878	2	1.105e-15
Europe	78.033	2	< 2.2e-16
political.knowledge	55.568	2	8.582e-13
Europe:political.knowledge	50.804	2	9.291e-12

Most of the predictors, including the Europe \times political.knowledge interaction, are associated with very small p-values; the Anova() function is from the **car** package (Fox and Weisberg 2019).

Here's an "effect plot", using the the **effects** package (Fox and Weisberg 2019) to visualize the Europe × political.knowledge interaction in a "stacked-area" graph:



To cross-validate this multinomial-logit model we need an appropriate cost criterion. None of the criteria supplied by the **cv** package—for example, neither mse(), which is appropriate for a numeric response, nor BayesRule(), which is appropriate for a binary response—will do. One possibility is to adapt Bayes rule to a polytomous response:

R> head(BEPS\$vote)

```
[1] Liberal Democrat Labour Labour Labour [5] Labour Labour Levels: Conservative Labour Liberal Democrat

R> yhat <- predict(m.beps, type="class")
R> head(yhat)
```

```
[1] Labour
                                                          Labour
                      Labour
                                        Labour
[5] Liberal Democrat Labour
Levels: Conservative Labour Liberal Democrat
R> BayesRuleMulti <- function(y, yhat){</pre>
    result <- mean(y != yhat)
    attr(result, "casewise loss") <- "y != yhat"
    result
+ }
R.>
R> BayesRuleMulti(BEPS$vote, yhat)
[1] 0.3186885
attr(,"casewise loss")
[1] "y != yhat"
```

The predict() method for "multinom" models called with argument type="class" reports the Bayes-rule prediction for each case—that is, the response category with the highest predicted probability. Our BayesRuleMulti() function calculates the proportion of misclassified cases. Because this value is the mean of casewise components, we attach a "casewise loss" attribute to the result (as explained in the preceding section).

The marginal proportions for the response categories are

and so the marginal Bayes-rule prediction, that everyone will vote Labour, produces an error rate of 1-0.47213=0.52787. The multinomial-logit model appears to do substantially better than that, but does its performance hold up to cross-validation?

We check first whether the default cv() method works "out-of-the-box" for the "multinom" model:

```
R> cv(m.beps, seed=3465, criterion=BayesRuleMulti)
```

Error in GetResponse.default(model): non-vector response

The default method of GetResponse() (a function supplied by the cv package—see ?GetResponse) fails for a "multinom" object. A straightforward solution is to supply a GetResponse.multinom() method that returns the factor response (using the get_response() function from the insight package, Lüdecke, Waggoner, and Makowski 2019),

```
R> GetResponse.multinom <- function(model, ...) {
+ insight::get_response(model)
+ }
R>
R> head(GetResponse(m.beps))
```

```
[1] Liberal Democrat Labour
                                                          Labour
                                        Labour
[5] Labour
                      Labour
Levels: Conservative Labour Liberal Democrat
and to try again:
R> cv(m.beps, seed=3465, criterion=BayesRuleMulti)
R RNG seed set to 3465
# weights: 36 (22 variable)
initial value 1507.296060
iter 10 value 1134.575036
iter 20 value 1037.413231
iter 30 value 1007.705242
iter 30 value 1007.705235
iter 30 value 1007.705235
final value 1007.705235
converged
Error in match.arg(type): 'arg' should be one of "class", "probs"
A traceback() (not shown) reveals that the problem is that the default method of cv()
calls the "multinom" method for predict() with the argument type="response", when the
correct argument should be type="class". We therefore must write a "multinom" method
for cv(), but that proves to be very simple:
R> cv.multinom <- function (model, data, criterion=BayesRuleMulti, k, reps,
                            seed, ...){
    NextMethod(type="class", criterion=criterion)
+ }
That is, we simply call the default cv() method with the type argument properly set. In
addition to supplying the correct type argument, our method sets the default criterion for
the cv.multinom() method to BayesRuleMulti.
Then:
R> m.beps <- update(m.beps, trace=FALSE)</pre>
R > cv(m.beps, seed=3465)
R RNG seed set to 3465
10-Fold Cross Validation
cross-validation criterion = 0.3245902
bias-adjusted cross-validation criterion = 0.3236756
95% CI for bias-adjusted CV criterion = (0.300168, 0.3471831)
full-sample criterion = 0.3186885
```

Prior to invoking cv(), we called update() with trace=FALSE to suppress the iteration history reported by default by multinom()—it would be tedious to see the iteration history for each fold. The cross-validated polytomous Bayes-rule criterion confirms that the fitted model does substantially better than the marginal Bayes-rule prediction that everyone votes for Labour.

Mixed-effects models

Adding a cv() method for a mixed-model class is somewhat more complicated. We provide the cvMixed() function to facilitate this process, and to see how that works, consider the "lme" method from the cv package:

```
R> cv:::cv.lme
function(model, data = insight::get_data(model), criterion = mse,
                   k, reps = 1, seed, ncores = 1, clusterVariables, ...){
  cvMixed(
    model,
    package="nlme",
    data=data,
    criterion=criterion,
    k=k,
    reps=reps,
    seed=seed,
    ncores=ncores,
    clusterVariables=clusterVariables,
    predict.clusters.args=list(object=model,
                                newdata=data,
                                level=0),
    predict.cases.args=list(object=model,
                             newdata=data,
                             level=1),
    ...)
}
<bytecode: 0x117501408>
<environment: namespace:cv>
```

Notice that cv.lme() sets up a call to cvMixed(), which does the computational work. Most of the arguments of cvMixed() are familiar:

- model is the mixed-model object, here of class "lme".
- package is the name of the package in which the mixed-modeling function used to fit the model, here lme(), resides—i.e., "nlme"; cvMixed() uses this argument to retrieve the package namespace.
- data is the data set to which the model is fit, by default extracted by the get_data() function in the insight package.

- criterion is the CV criterion, defaulting to the mse() function.
- k is the number of CV folds, defaulting to "loo" for CV by clusters and 10 for CV by cases.
- reps is the number of times the CV process is repeated, defaulting to 1.
- seed is the seed for R's random-number generator, defaulting to a randomly selected (and saved) value.
- ncores is the number of cores to use for parallel computation; if 1, the default, then the computation isn't parallelized.
- clusterVariables is a character vector of the names of variables defining clusters; if missing, then CV is based on cases rather than clusters.

The remaining two arguments are unfamiliar:

- predict.clusters.args is a named list of arguments to be passed to the predict() function to obtain predictions for the full data set from a model fit to a subset of the data for cluster-based CV. The first two arguments should be object and newdata. It is typically necessary to tell cvMixed() how to base predictions only on fixed effects; in the case of "lme" models, this is done by setting level = 0.
- Similarly, predict.cases.args is a named list of arguments to be passed to predict() for case-based CV. Setting level = 1 includes random effects in the predictions.

Finally, any additional arguments, absorbed by ..., are passed to update() when the model is refit with each fold omitted. cvMixed() returns an object of class "cv".

Now imagine that we want to support a new class of mixed-effects models. To be concrete, we illustrate with the glmmPQL() function in the MASS package (Venables and Ripley 2002), which fits generalized-linear mixed-effects models by penalized quasi-likelihood. Not coincidentally, the arguments of glmmPQL() are similar to those of lme() (with an additional family argument), because the former iteratively invokes the latter; so cv.glmmPQL() should resemble cv.lme().

As it turns out, neither the default method for GetResponse() nor insight::get_data() work for "glmmPQL" objects. These objects include a "data" element, however, and so we can simply extract this element as the default for the data argument of our cv.glmmPQL() method.

To get the response variable is more complicated: We refit the fixed part of the model as a GLM with only the regression constant on the right-hand side, and extract the response from that; because all we need is the response variable, we limit the number of GLM iterations to 1 and suppress warning messages about non-convergence:

¹⁶This example is somewhat artificial in that glmmPQL() has largely been superseded by computationally superior functions, such the glmer() function in the lme4 package. There is, however, one situation in which glmmPQL() might prove useful: to specify serial dependency in case-level errors within clusters for longitudinal data, which is not currently supported by glmer().

```
R> GetResponse.glmmPQL <- function(model, ...){</pre>
    f <- formula(model)</pre>
    f[[3]] <- 1 # regression constant only on RHS
   model <- suppressWarnings(glm(f, data=model$data, family=model$family,
                                   control=list(maxit=1)))
    cv::GetResponse(model)
+ }
Writing the cv() method is then straightforward:
R> cv.glmmPQL <- function(model, data = model$data, criterion = mse,
                        k, reps = 1, seed, ncores = 1, clusterVariables, ...){
    cvMixed(
      model,
      package="MASS",
      data=data,
      criterion=criterion,
      k=k,
      reps=reps,
      seed=seed,
      ncores=ncores,
      clusterVariables=clusterVariables,
      predict.clusters.args=list(object=model,
                                  newdata=data,
                                  level=0,
                                  type="response"),
      predict.cases.args=list(object=model,
                               newdata=data,
                               level=1,
                               type="response"),
      verbose=FALSE,
      ...)
+ }
```

We set the argument verbose=FALSE to suppress glmmPQL()'s iteration counter when cvMixed() calls update().

Let's apply our newly minted method to a logistic regression with a random intercept in an example that appears in <code>?glmmPQL</code>:

```
iteration 3
iteration 4
iteration 5
iteration 6
R> summary(m.pql)
Linear mixed-effects model fit by maximum likelihood
  Data: bacteria
  AIC BIC logLik
  NA NA
Random effects:
 Formula: ~1 | ID
        (Intercept) Residual
StdDev: 1.410637 0.7800511
Variance function:
 Structure: fixed weights
 Formula: ~invwt
Fixed effects: y ~ trt + I(week > 2)
                    Value Std.Error DF t-value p-value
(Intercept)
               3.412014 0.5185033 169 6.580506 0.0000
                -1.247355 0.6440635 47 -1.936696 0.0588
trtdrug
trtdrug+
               -0.754327 0.6453978 47 -1.168779 0.2484
I(week > 2)TRUE -1.607257 0.3583379 169 -4.485311 0.0000
 Correlation:
                (Intr) trtdrg trtdr+
                -0.598
trtdrug
                -0.571 0.460
trtdrug+
I(week > 2)TRUE -0.537 0.047 -0.001
Standardized Within-Group Residuals:
                   Q1
                             Med
                                         QЗ
                                                   Max
-5.1985361 0.1572336 0.3513075 0.4949482 1.7448845
Number of Observations: 220
Number of Groups: 50
We compare this result to that obtained from glmer() in the lme4 package:
R> library("lme4")
R> m.glmer \leftarrow glmer(y \sim trt + I(week > 2) + (1 | ID),
                 family = binomial, data = bacteria)
R> summary(m.glmer)
```

Generalized linear mixed model fit by maximum likelihood (Laplace Approximation) [glmerMod] Family: binomial (logit) Formula: $y \sim trt + I(week > 2) + (1 | ID)$ Data: bacteria AIC BIC logLik deviance df.resid 202.3 219.2 -96.1 192.3 215 Scaled residuals: Min 1Q Median ЗQ Max -4.5615 0.1359 0.3022 0.4217 1.1276 Random effects: Groups Name Variance Std.Dev. (Intercept) 1.543 1.242 Number of obs: 220, groups: ID, 50 Fixed effects: Estimate Std. Error z value Pr(>|z|)(Intercept) 3.5479 0.6958 5.099 3.41e-07 trtdrug -1.3667 0.6770 -2.019 0.043517 -0.7826 0.6831 -1.146 0.251925 trtdrug+ I(week > 2)TRUE -1.59850.4759 -3.359 0.000783 Correlation of Fixed Effects: (Intr) trtdrg trtdr+ trtdrug -0.593trtdrug+ -0.537 0.487 I(wk>2)TRUE -0.656 0.126 0.064 # comparison of fixed effects: R> car::compareCoefs(m.pql, m.glmer) Warning in car::compareCoefs(m.pql, m.glmer): models to be compared are of different classes Calls: 1: glmmPQL(fixed = y ~ trt + I(week > 2), random = ~1 | ID, family = binomial, data = bacteria) 2: $glmer(formula = y \sim trt + I(week > 2) + (1 | ID), data = bacteria,$ family = binomial) Model 1 Model 2 (Intercept) 3.412 3.548 SE 0.514 0.696

```
trtdrug -1.247 -1.367

SE 0.638 0.677

trtdrug+ -0.754 -0.783

SE 0.640 0.683

I(week > 2)TRUE -1.607 -1.598

SE 0.355 0.476
```

The two sets of estimates are similar, but not identical

Finally, we try out our cv.glmmPQL() method, cross-validating both by clusters and by cases,

R> cv(m.pql, clusterVariables="ID", criterion=BayesRule)

n-Fold Cross Validation based on 50 {ID} clusters cross-validation criterion = 0.1954545 bias-adjusted cross-validation criterion = 0.1954545 full-sample criterion = 0.1954545

R> cv(m.pql, data=bacteria, criterion=BayesRule, seed=1490)

R RNG seed set to 1490

10-Fold Cross Validation cross-validation criterion = 0.2090909 bias-adjusted cross-validation criterion = 0.2072727 full-sample criterion = 0.1454545

and again compare to glmer():

R> cv(m.glmer, clusterVariables="ID", criterion=BayesRule)

n-Fold Cross Validation based on 50 {ID} clusters cross-validation criterion = 0.1954545 bias-adjusted cross-validation criterion = 0.1954545 full-sample criterion = 0.1954545

R> cv(m.glmer, data=bacteria, criterion=BayesRule, seed=1490)

R RNG seed set to 1490

10-Fold Cross Validation cross-validation criterion = 0.1954545 bias-adjusted cross-validation criterion = 0.1936364 full-sample criterion = 0.15

7.3. Adding a model-selection procedure

The selectStepAIC() function supplied by the cv package, which is based on the stepAIC() function from the nnet package (Venables and Ripley 2002) for stepwise model selection, is suitable for the procedure argument of cvSelect(). The use of selectStepAIC() is illustrated in the principal vignette for the package.

We'll employ selectStepAIC() as a "template" for writing a CV model-selection procedure. To see the code for this function, type cv::selectStepAIC at the R command prompt, or examine the sources for the cv package at https://github.com/gmonette/cv (the code for selectStepAIC() is in https://github.com/gmonette/cv/blob/main/R/cvSelect.R).

Another approach to model selection is all-subsets regression. The regsubsets() function in the leaps package (Lumley and Miller 2020) implements an efficient algorithm for selecting the best-fitting linear least-squares regressions for subsets of predictors of all sizes, from 1 through the maximum number of candidate predictors.¹⁷ To illustrate the use of regsubsets(), we employ the swiss data frame supplied by the leaps package:

```
R> library("leaps")
R> head(swiss)
```

	Fertility	Agriculture	Examination	Education	Catholic
Courtelary	80.2	17.0	15	12	9.96
Delemont	83.1	45.1	6	9	84.84
Franches-Mnt	92.5	39.7	5	5	93.40
Moutier	85.8	36.5	12	7	33.77
Neuveville	76.9	43.5	17	15	5.16
Porrentruy	76.1	35.3	9	7	90.57
	Infant.Mon	ctality			
Courtelary		22.2			
Delemont		22.2			
Franches-Mnt		20.2			
Moutier		20.3			
Neuveville		20.6			
Porrentruy		26.6			

R> nrow(swiss)

[1] 47

The data set includes the following variables, for each of 47 French-speaking Swiss provinces circa 1888:

• Fertility: A standardized fertility measure.

¹⁷The regsubsets() function computes several measures of model predictive performance, including the R^2 and R^2 adjusted for degrees of freedom, the residual sums of squares, Mallows's C_p , and the BIC. Several of these are suitable for comparing models with differing numbers of coefficients—we use the BIC below—but all necessarily agree when comparing models with the *same* number of coefficients.

- Agriculture: The percentage of the male population engaged in agriculture.
- Examination: The percentage of draftees into the Swiss army receiving the highest grade on an examination.
- Education: The percentage of draftees with more than a primary-school education.
- Catholic: The percentage of the population who were Catholic.
- Infant.Mortality: The infant-mortality rate, expressed as the percentage of live births surviving less than a year.

Following Lumley and Miller (2020), we treat Fertility as the response and the other variables as predictors in a linear least-squares regression:

```
R> m.swiss <- lm(Fertility ~ ., data=swiss)
R> summary(m.swiss)
```

Call:

lm(formula = Fertility ~ ., data = swiss)

Residuals:

```
Min 1Q Median 3Q Max -15.2743 -5.2617 0.5032 4.1198 15.3213
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                66.91518 10.70604 6.250 1.91e-07
Agriculture
                -0.17211
                          0.07030 -2.448 0.01873
Examination
                -0.25801
                           0.25388 -1.016 0.31546
Education
                -0.87094
                           0.18303 -4.758 2.43e-05
Catholic
                0.10412
                           0.03526
                                    2.953 0.00519
Infant.Mortality 1.07705
                           0.38172
                                    2.822 0.00734
```

Residual standard error: 7.165 on 41 degrees of freedom Multiple R-squared: 0.7067, Adjusted R-squared: 0.671 F-statistic: 19.76 on 5 and 41 DF, p-value: 5.594e-10

```
R > cv(m.swiss, seed=8433)
```

R RNG seed set to 8433

```
10-Fold Cross Validation
```

method: Woodbury
criterion: mse

cross-validation criterion = 59.68315

bias-adjusted cross-validation criterion = 58.84614

full-sample criterion = 44.78815

Thus, the RMSE for the model fit to the complete data is considerably smaller than the CV estimate of the RMSE. Can we do better by selecting a subset of the predictors, taking account of the additional uncertainty induced by model selection?

First, let's apply best-subset selection to the complete data set:

```
R> swiss.sub <- regsubsets(Fertility ~ ., data=swiss)</pre>
R> summary(swiss.sub)
Subset selection object
Call: regsubsets.formula(Fertility ~ ., data = swiss)
5 Variables
             (and intercept)
                 Forced in Forced out
Agriculture
                     FALSE
                                 FALSE
Examination
                     FALSE
                                 FALSE
Education
                     FALSE
                                 FALSE
Catholic
                      FALSE
                                 FALSE
Infant.Mortality
                     FALSE
                                 FALSE
1 subsets of each size up to 5
Selection Algorithm: exhaustive
         Agriculture Examination Education Catholic Infant. Mortality
                                                      11 11
                      11 11
                                            11 11
  (1)""
                                  "*"
1
                      11 11
                                  "*"
                                             "*"
                                                      11 11
  (1)""
                      11 11
3 (1)""
                                  "*"
                                             "*"
                                                      "*"
4 (1) "*"
                      11 11
                                  "*"
                                             "*"
                                                      "*"
5 (1) "*"
                      "*"
                                  "*"
                                             "*"
                                                      "*"
R> (bics <- summary(swiss.sub)$bic)</pre>
[1] -19.60287 -28.61139 -35.65643 -37.23388 -34.55301
R> which.min(bics)
[1] 4
R> car::subsets(swiss.sub, legend="topright")
```

The graph, produced by the subsets() function in the car package, shows that the model with the smallest BIC is the best model with 4 predictors, including Agriculture, Education, Catholic, and Infant.Mortality, but not Examination:

```
R> summary(m.best)
Call:
```

lm(formula = Fertility ~ Agriculture + Education + Catholic +

R> m.best <- update(m.swiss, . ~ . - Examination)</pre>

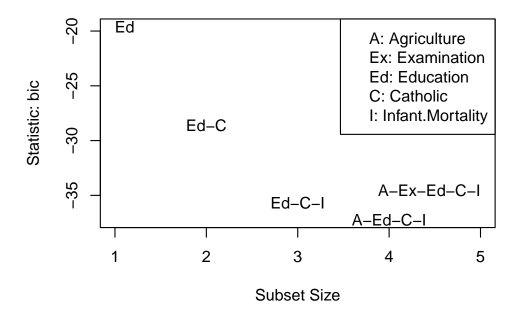


Figure 23: Selecting the best model of each size.

Infant.Mortality, data = swiss)

Residuals:

Min	1Q	Median	3Q	Max
-14.6765	-6.0522	0.7514	3.1664	16.1422

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	62.10131	9.60489	6.466	8.49e-08
Agriculture	-0.15462	0.06819	-2.267	0.02857
Education	-0.98026	0.14814	-6.617	5.14e-08
Catholic	0.12467	0.02889	4.315	9.50e-05
Infant Mortality	1 07844	0.38187	2 824	0 00722

Residual standard error: 7.168 on 42 degrees of freedom Multiple R-squared: 0.6993, Adjusted R-squared: 0.6707 F-statistic: 24.42 on 4 and 42 DF, p-value: 1.717e-10

R> cv(m.best, seed=8433) # use same folds as before

R RNG seed set to 8433

10-Fold Cross Validation

method: Woodbury
criterion: mse

cross-validation criterion = 58.46664

bias-adjusted cross-validation criterion = 57.7779

full-sample criterion = 45.91637

The RMSE for the selected model is (of course) slightly higher than for the full model fit previously, but the cross-validated RMSE is a bit lower; as we explain in the main vignette, however, it isn't kosher to select and cross-validate a model on the same data.

Here's a function named selectSubsets(), meant to be used with cvSelect(), suitable for cross-validating the model-selection process:

```
R> selectSubsets <- function(data=insight::get_data(model),</pre>
                              model,
                              indices,
                              criterion=mse,
                              save.coef=TRUE, ...){
    if (inherits(model, "lm", which=TRUE) != 1)
      stop("selectSubsets is appropriate only for 'lm', models")
+
    y <- GetResponse(model)</pre>
    formula <- formula(model)</pre>
    X <- model.matrix(model)</pre>
    if (missing(indices)) {
      # select the best model from the full data by BIC
      sel <- leaps::regsubsets(formula, data=data, ...)</pre>
      bics <- summary(sel)$bic</pre>
      best <- coef(sel, 1:length(bics))[[which.min(bics)]]</pre>
      x.names <- names(best)</pre>
      # fit the best model; intercept is already in X, hence - 1:
      m.best \leftarrow lm(y \sim X[, x.names] - 1)
      fit.all <- predict(m.best, newdata=data)</pre>
      return(criterion(y, fit.all)) # return the CV criterion
    }
+
    # select the best model omitting the i-th fold (given by indices)
    sel.i <- leaps::regsubsets(formula, data[-indices, ], ...)</pre>
    bics.i <- summary(sel.i)$bic</pre>
    best.i <- coef(sel.i, 1:length(bics.i))[[which.min(bics.i)]]</pre>
    x.names.i <- names(best.i)</pre>
    m.best.i <- lm(y[-indices] ~ X[-indices, x.names.i] - 1)</pre>
                 # predict() doesn't work here:
    fit.all.i <- as.vector(X[, x.names.i] %*% coef(m.best.i))</pre>
    fit.i <- fit.all.i[indices]</pre>
    # return the fitted values for i-th fold, CV criterion for all cases,
        and the regression coefficients
    list(fit.i=fit.i, # fitted values for i-th fold
         crit.all.i=criterion(y, fit.all.i), # CV crit for all cases
         coefficients = if (save.coef){ # regression coefficients
            coefs <- coef(m.best.i)</pre>
```

A slightly tricky point is that because of scoping issues, predict() doesn't work with the model fit omitting the *i*th fold, and so the fitted values for all cases are computed directly as $\hat{\mathbf{y}}_{-i} = \mathbf{X}\mathbf{b}_{-i}$, where **X** is the model-matrix for all of the cases, and \mathbf{b}_{-i} is the vector of least-squares coefficients for the selected model with the *i*th fold omitted.

Additionally, the command $lm(y[-indices] \sim X[-indices, x.names.i] - 1)$, which is the selected model with the *i*th fold deleted, produces awkward coefficient names like "X[-indices, x.names.i]Infant.Mortality". Purely for aesthetic reasons, the command sub("X\\[-indices, x.names.i\\]", "", names(coefs)) fixes these awkward names, removing the extraneous text, "X[-indices, x.names.i]".

Applying selectSubsets() to the full data produces the full-data cross-validated RMSE (which we obtained previously): %%{r test-selectSubsets} %%selectSubsets(model=m.swiss) %% Similarly, applying the function to an imaginary "fold" of 5 cases returns the RMSE for the cases in the fold, based on the model selected and fit to the cases omitting the fold; the RMSE for all of the cases, based on the same model; and the coefficients of the selected model, which includes 4 or the 5 predictors (and the intercept): %%{r test-selectSubsets-fold} %%selectSubsets(model=m.swiss, indices=seq(5, 45, by=10)) %%

Then, using selectSubsets() in cross-validation, we get:

Cross-validation shows that model selection exacts a penalty in RMSE. Examining the models selected for the 10 folds reveals that there is some uncertainty in identifying the predictors in the "best" model, with Agriculture sometimes appearing and sometimes not:

```
R> compareFolds(cv.swiss)
```

		(Intercept)	${\tt Catholic}$	Education	<pre>Infant.Mortality</pre>	Agriculture
Fold	1	59.0852	0.1397	-1.0203	1.2985	-0.17
Fold	2	67.0335	0.1367	-1.0499	0.9413	-0.20
Fold	3	55.0453	0.1221	-0.8757	1.3541	-0.15
Fold	4	62.5543	0.1236	-0.9719	1.0679	-0.16
Fold	5	50.4643	0.1057	-0.7863	1.2144	
Fold	6	68.0289	0.1195	-1.0073	0.8294	-0.17
Fold	7	66.5219	0.1357	-1.0827	0.9523	-0.19
Fold	8	46.3507	0.0776	-0.7637	1.4463	
Fold	9	62.2632	0.1230	-1.0067	1.1000	-0.17
Fold	10	52.5112	0.1005	-0.7232	1.0809	

8. Computational notes

8.1. Efficient computations for linear and generalized linear models

The most straightforward way to implement cross-validation in R for statistical modeling functions that are written in the canonical manner is to use update() to refit the model with each fold removed. This is the approach taken in the default method for cv(), and it is appropriate if the cases are independently sampled. Refitting the model in this manner for each fold is generally feasible when the number of folds in modest, but can be prohibitively costly for leave-one-out cross-validation when the number of cases is large.

The "lm" and "glm" methods for cv() take advantage of computational efficiencies by avoiding refitting the model with each fold removed. Consider, in particular, the weighted linear model $\mathbf{y}_{n\times 1} = \mathbf{X}_{n\times p}\boldsymbol{\beta}_{p\times 1} + \boldsymbol{\varepsilon}_{n\times 1}$, where $\boldsymbol{\varepsilon} \sim \mathbf{N}_n\left(\mathbf{0},\sigma^2\mathbf{W}_{n\times n}^{-1}\right)$. Here, \mathbf{y} is the response vector, \mathbf{X} the model matrix, and $\boldsymbol{\varepsilon}$ the error vector, each for n cases, and $\boldsymbol{\beta}$ is the vector of p population regression coefficients. The errors are assumed to be multivariately normally distributed with 0 means and covariance matrix $\sigma^2\mathbf{W}^{-1}$, where $\mathbf{W} = \operatorname{diag}(w_i)$ is a diagonal matrix of inverse-variance weights. For the linear model with constant error variance, the weight matrix is taken to be $\mathbf{W} = \mathbf{I}_n$, the order-n identity matrix.

The weighted-least-squares (WLS) estimator of β is (see, e.g., Fox 2016, Sec. 12.2.2) ¹⁸

$$\mathbf{b}_{\mathrm{WLS}} = \left(\mathbf{X}^T \mathbf{W} \mathbf{X}\right)^{-1} \mathbf{X}^T \mathbf{W} \mathbf{y}$$

Fitted values are then $\hat{\mathbf{y}} = \mathbf{X}\mathbf{b}_{\text{WLS}}$.

The LOO fitted value for the *i*th case can be efficiently computed by $\hat{y}_{-i} = y_i - e_i/(1 - h_i)$ where $h_i = \mathbf{x}_i^T \left(\mathbf{X}^T \mathbf{W} \mathbf{X}\right)^{-1} \mathbf{x}_i$ (the so-called "hatvalue"). Here, \mathbf{x}_i^T is the *i*th row of \mathbf{X} , and \mathbf{x}_i is the *i*th row written as a column vector. This approach can break down when one or more hatvalues are equal to 1, in which case the formula for \hat{y}_{-i} requires division by 0.

¹⁸This is a definitional formula, which assumes that the model matrix \mathbf{X} is of full column rank, and which can be subject to numerical instability when \mathbf{X} is ill-conditioned. $\mathtt{lm}()$ uses the singular-value decomposition of the model matrix to obtain computationally more stable results.

To compute cross-validated fitted values when the folds contain more than one case, we make use of the Woodbury matrix identify ("Woodbury matrix identity" 2023),

$$\left(\mathbf{A}_{m\times m} + \mathbf{U}_{m\times k}\mathbf{C}_{k\times k}\mathbf{V}_{k\times m}\right)^{-1} = \mathbf{A}^{-1} - \mathbf{A}^{-1}\mathbf{U}\left(\mathbf{C}^{-1} + \mathbf{V}\mathbf{A}^{-1}\mathbf{U}\right)^{-1}\mathbf{V}\mathbf{A}^{-1}$$

where \mathbf{A} is a nonsingular order-n matrix. We apply this result by letting

$$\mathbf{A} = \mathbf{X}^T \mathbf{W} \mathbf{X}$$
$$\mathbf{U} = \mathbf{X}_{\mathbf{j}}^T$$
$$\mathbf{V} = -\mathbf{X}_{\mathbf{j}}$$
$$\mathbf{C} = \mathbf{W}_{\mathbf{i}}$$

where the subscript $\mathbf{j} = (i_{j1}, \dots, i_{jm})^T$ represents the vector of indices for the cases in the jth fold, $j = 1, \dots, k$. The negative sign in $\mathbf{V} = -\mathbf{X}_{\mathbf{j}}$ reflects the *removal*, rather than addition, of the cases in \mathbf{j} .

Applying the Woodbury identity isn't quite as fast as using the hat values, but it is generally much faster than refitting the model. A disadvantage of the Woodbury identity, however, is that it entails explicit matrix inversion and thus may be numerically unstable. The inverse of $\mathbf{A} = \mathbf{X}^T \mathbf{W} \mathbf{X}$ is available directly in the "lm" object, but the second term on the right-hand side of the Woodbury identity requires a matrix inversion with each fold deleted. (In contrast, the inverse of each $\mathbf{C} = \mathbf{W}_{\mathbf{j}}$ is straightforward because \mathbf{W} is diagonal.)

The Woodbury identity also requires that the model matrix be of full rank. We impose that restriction in our code by removing redundant regressors from the model matrix for all of the cases, but that doesn't preclude rank deficiency from surfacing when a fold is removed. Rank deficiency of \mathbf{X} doesn't disqualify cross-validation because all we need are fitted values under the estimated model.

glm() computes the maximum-likelihood estimates for a generalized linear model by iterated weighted least squares (see, e.g., Fox and Weisberg 2019, Sec. 6.12). The last iteration is therefore just a WLS fit of the "working response" on the model matrix using "working weights." Both the working weights and the working response at convergence are available from the information in the object returned by glm().

We then treat re-estimation of the model with a case or cases deleted as a WLS problem, using the hatvalues or the Woodbury matrix identity. The resulting fitted values for the deleted fold aren't exact—that is, except for the Gaussian family, the result isn't identical to what we would obtain by literally refitting the model—but in our (limited) experience, the approximation is very good, especially for LOO CV, which is when we would be most tempted to use it. Nevertheless, because these results are approximate, the default for the "glm" cv() method is to perform the exact computation, which entails refitting the model with each fold omitted.

8.2. Computation of the bias-corrected CV criterion and confidence intervals

Let $CV(\mathbf{y}, \hat{\mathbf{y}})$ represent a cross-validation cost criterion, such as mean-squared error, computed for all of the n values of the response \mathbf{y} based on fitted values $\hat{\mathbf{y}}$ from the model fit to

all of the data. We require that $CV(\mathbf{y}, \widehat{\mathbf{y}})$ is the mean of casewise components, that is, $CV(\mathbf{y}, \widehat{\mathbf{y}}) = \frac{1}{n} \sum_{i=1}^{n} cv(y_i, \widehat{y}_i)$. For example, $MSE(\mathbf{y}, \widehat{\mathbf{y}}) = \frac{1}{n} \sum_{i=1}^{n} (y_i - \widehat{y}_i)^2$.

We divide the *n* cases into *k* folds of approximately $n_j \approx n/k$ cases each, where $n = \sum n_j$. As above, let **j** denote the indices of the cases in the *j*th fold.

Now define $CV_j = CV(\mathbf{y}, \hat{\mathbf{y}}^{(j)})$. The superscript (j) on $\hat{\mathbf{y}}^{(j)}$ represents fitted values computed for all of the cases from the model with fold j omitted. Let $\hat{\mathbf{y}}^{(-i)}$ represent the vector of fitted values for all n cases where the fitted value for the ith case is computed from the model fit with the fold including the ith case omitted (i.e., fold j for which $i \in \mathbf{j}$).

Then the cross-validation criterion is just $CV = CV(\mathbf{y}, \widehat{\mathbf{y}}^{(-i)})$. Following Davison and Hinkley (1997, pp. 293–295), the bias-adjusted cross-validation criterion is

$$CV_{adj} = CV + CV(\mathbf{y}, \hat{\mathbf{y}}) - \frac{1}{n} \sum_{j=1}^{k} n_j CV_j$$

We compute the standard error of CV as

$$SE(CV) = \frac{1}{\sqrt{n}} \sqrt{\frac{\sum_{i=1}^{n} \left[cv(y_i, \hat{y}_i^{(-i)}) - CV \right]^2}{n-1}}$$

that is, as the standard deviation of the casewise components of CV divided by the square-root of the number of cases.

We then use SE(CV) to construct a $100 \times (1 - \alpha)\%$ confidence interval around the *adjusted* CV estimate of error:

$$\left[\mathrm{CV}_{\mathrm{adj}} - z_{1-\alpha/2} \mathrm{SE}(\mathrm{CV}), \mathrm{CV}_{\mathrm{adj}} + z_{1-\alpha/2} \mathrm{SE}(\mathrm{CV})\right]$$

where $z_{1-\alpha/2}$ is the $1-\alpha/2$ quantile of the standard-normal distribution (e.g, $z \approx 1.96$ for a 95% confidence interval, for which $1-\alpha/2 = .975$).

Bates et al. (2023) show that the coverage of this confidence interval is poor for small samples, and they suggest a much more computationally intensive procedure, called nested cross-validation, to compute better estimates of error and confidence intervals with better coverage for small samples. We may implement Bates et al.'s approach in a later release of the \mathbf{cv} package. At present we use the confidence interval above for sufficiently large n, which, based on Bates et al.'s results, we take by default to be $n \geq 400$.

9. Why the complement of AUC isn't a casewise CV criterion

Consider calculating AUC for folds in which a validation set contains n_v observations. To calculate AUC in the validation set, we need the vector of prediction criteria, $\hat{\mathbf{y}}_{v_{(n_v \times 1)}} = (\hat{y}_1, ..., \hat{y}_{n_v})^T$, and the vector of observed responses in the validation set, $\mathbf{y}_{v_{(n_v \times 1)}} = (y_1, ..., y_{n_v})^T$ with $y_i \in \{0, 1\}, i = 1, ..., n_v$.

To construct the ROC curve, only the ordering of the values in $\hat{\mathbf{y}}_v$ is relevant. Thus, assuming that there are no ties, and reordering observations if necessary, we can set $\hat{\mathbf{y}}_v = (1, 2, \dots, n_v)^T$.

¹⁹Arlot and Celisse (2010) term the casewise loss, $cv(y_i, \hat{y}_i)$, the "contrast function."

If the AUC can be expressed as the casewise mean or sum of a function $cv(\hat{y}_i, y_i)$, where $cv: \{1, 2, ..., n_v\} \times \{0, 1\} \rightarrow [0, 1]$, then

$$\sum_{i=1}^{n_v} \operatorname{cv}(\widehat{y}_i, y_i) = \operatorname{AUC}(\widehat{\mathbf{y}}_v, \mathbf{y}_v)$$
(1)

must hold for all 2^{n_v} possible values of $\mathbf{y}_v = (y_1, ..., y_{n_v})^T$. If all ys have the same value, either 1 or 0, then the definition of AUC is ambiguous. AUC could be considered undefined, or it could be set to 0 if all ys are 0 and to 1 if all ys are 1. If AUC is considered to be undefined in these cases, we have $2^{n_v} - 2$ admissible values for \mathbf{y}_v .

Thus, equation (1) produces either 2^{n_v} or $2^{n_v} - 2$ constraints. Although there are only $2n_v$ possible values for the $\operatorname{cv}(\cdot)$ function, equation (1) could, nevertheless, have consistent solutions. We therefore need to determine whether there is a value of n_v for which (1) has no consistent solution for all admissible values of \mathbf{y}_v . In that eventuality, we will have shown that AUC cannot, in general, be expressed through a casewise sum.

If $n_v = 3$, we show below that (1) has no consistent solution if we include all possibilities for \mathbf{y}_v , but does if we exclude cases where all ys have the same value. If $n_v = 4$, we show that there are no consistent solutions in either case.

The following R function computes AUC from $\hat{\mathbf{y}}_v$ and \mathbf{y}_v , accommodating the cases where \mathbf{y}_v is all 0s or all 1s:

```
R> AUC <- function(y, yhat = seq_along(y)) {
+    s <- sum(y)
+    if (s == 0) return(0)
+    if (s == length(y)) return(1)
+    Metrics::auc(y, yhat)
+ }</pre>
```

We then define a function to generate all possible \mathbf{y}_v s of length n_v as rows of the matrix $\mathbf{Y}_{(2^{n_v} \times n_v)}$:

```
R> Ymat <- function(n_v, exclude_identical = FALSE) {  + \text{ stopifnot}(n_v > 0 \&\& \text{ round}(n_v) == n_v) \quad \# n_v \text{ must be a positive integer } \\ + \text{ ret } <- \text{ sapply}(0:(2^n_v - 1), \\ + \quad \text{ function}(x) \text{ as.integer}(\text{intToBits}(x)) )[1:n_v,] \\ + \text{ ret } <- \text{ if } (\text{is.matrix}(\text{ret})) \text{ t}(\text{ret}) \text{ else matrix}(\text{ret}) \\ + \text{ colnames}(\text{ret}) <- \text{ paste0}("y", 1:\text{ncol}(\text{ret})) \\ + \text{ if } (\text{exclude\_identical}) \text{ ret}[-c(1, \text{ nrow}(\text{ret})),] \text{ else ret} \\ + \} \\ \text{For } n_v = 3, \\ R> \text{ Ymat}(3) \\ \text{y1 y2 y3} \\ \text{[1,]} \quad 0 \quad 0 \quad 0
```

If we exclude \mathbf{y}_v s with identical values, then

R> Ymat(3, exclude_identical = TRUE)

Here is Y with corresponding values of AUC:

R> cbind(Ymat(3), AUC = apply(Ymat(3), 1, AUC))

The values of $\operatorname{cv}(\widehat{y}_i, y_i)$ that express AUC as a sum of casewise values are solutions of equation (1), which can be written as solutions of the following system of 2^{n_v} linear simultaneous equations in $2n_v$ unknowns:

$$(\mathbf{U} - \mathbf{Y})\mathbf{c}_0 + \mathbf{Y}\mathbf{c}_1 = [\mathbf{U} - \mathbf{Y}, \mathbf{Y}]\begin{bmatrix} \mathbf{c}_0 \\ \mathbf{c}_1 \end{bmatrix} = AUC(\widehat{\mathbf{Y}}, \mathbf{Y})$$
 (2)

where $\mathbf{U}_{(2^{n_v} \times n_v)}$ is a matrix of 1s conformable with \mathbf{Y} ; $\mathbf{c}_0 = [\operatorname{cv}(1,0), c(2,0), ..., \operatorname{cv}(n_v,0)]^T$; $\mathbf{c}_1 = [\operatorname{cv}(1,1), c(2,1), ..., \operatorname{cv}(n_v,1)]^T$; $[\mathbf{U} - \mathbf{Y}, \mathbf{Y}]_{(2^{n_v} \times 2n_v)}$ and $\begin{bmatrix} \mathbf{c}_0 \\ \mathbf{c}_1 \end{bmatrix}_{(2n_v \times 1)}$ are partitioned matrix

trices; and $\hat{\mathbf{Y}}_{(2^{n_v} \times n_v)}$ is a matrix each of whose rows consists of the integers 1 to n_v .

We can test whether equation (2) has a solution for any given n_v by trying to solve it as a least-squares problem, considering whether the residuals of the associated linear model are all 0, using the "design matrix" $[\mathbf{U} - \mathbf{Y}, \mathbf{Y}]$ to predict the "outcome" $\mathrm{AUC}(\widehat{\mathbf{Y}}, \mathbf{Y})_{(2^{n_v} \times 1)}$:

```
R> resids <- function(n_v, exclude_identical = FALSE,
                     tol = sqrt(.Machine$double.eps)) {
    Y <- Ymat(n_v, exclude_identical = exclude_identical)
   AUC <- apply(Y, 1, AUC)
   X \leftarrow cbind(1-Y, Y)
    opts <- options(warn = -1)
   on.exit(options(opts))
  fit <- lsfit(X, AUC, intercept = FALSE)</pre>
   ret <- max(abs(residuals(fit)))</pre>
  if(ret < tol){</pre>
     ret <- 0
    solution <- coef(fit)</pre>
    rep(0:1, each = n_v), ")")
      attr(ret, "solution") <- zapsmall(solution)</pre>
    }
    ret
+ }
The case n_v = 3, excluding identical ys, has a solution:
R> resids(3, exclude_identical = TRUE)
[1] 0
attr(, "solution")
c(1,0) c(2,0) c(3,0) c(1,1) c(2,1) c(3,1)
          0.0 -0.5 0.5 0.0 0.0
But, if identical ys are included, the equation is not consistent:
R> resids(3, exclude_identical = FALSE)
[1] 0.125
For n_v = 4, there are no solutions in either case:
R> resids(4, exclude_identical = TRUE)
[1] 0.08333333
R> resids(4, exclude_identical = FALSE)
[1] 0.25
```

Consequently, the widely employed AUC measure of fit for binary regression cannot in general be used for a casewise cross-validation criterion.

References

- Arlot S, Celisse A (2010). "A survey of cross-validation procedures for model selection." Statistics Surveys, 4, 40 79. URL https://doi.org/10.1214/09-SS054.
- Bates D, Mächler M, Bolker B, Walker S (2015). "Fitting Linear Mixed-Effects Models Using lme4." *Journal of Statistical Software*, **67**(1), 1–48.
- Bates S, Hastie T, Tibshirani R (2023). "Cross-validation: What does it estimate and how well does it do it?" Journal of the American Statistical Association, in press. URL https://doi.org/10.1080/01621459.2023.2197686.
- Box GEP, Cox DR (1964). "An analysis of transformations." Journal of the Royal Statistical Society, Series B, 26, 211–252.
- Canty A, Ripley BD (2022). boot: Bootstrap R (S-Plus) Functions. R package version 1.3-28.1.
- Davison AC, Hinkley DV (1997). Bootstrap Methods and Their Applications. Cambridge University Press, Cambridge.
- Diggle PJ, Liang KY, Zeger SL (1994). Analysis of longitudinal data. Oxford University Press, Oxford.
- Fox J (2016). Applied Regression Analysis and Generalized Linear Models. Second edition edition. Sage, Thousand Oaks CA.
- Fox J, Weisberg S (2019). An R Companion to Applied Regression. Third edition edition. Sage, Thousand Oaks CA.
- Hamner B, Frasco M (2018). *Metrics: Evaluation Metrics for Machine Learning*. R package version 0.1.4, URL https://CRAN.R-project.org/package=Metrics.
- Harrell Jr F (2015). Regression Modeling Strategies. Second edition edition. Springer, New York.
- Hastie T, Tibshirani R, Friedman J (2009). The Elements of Statistical Learning: Data Mining, Inference, and Prediction. Second edition edition. Springer, New York. URL https://hastie.su.domains/ElemStatLearn/printings/ESLII_print12_toc.pdf.
- James G, Witten D, Hastie T, Tibshirani R (2021). An Introduction to Statistical Learning with Applications in R. Second edition edition. Springer, New York.
- Lumley T, Miller A (2020). leaps: Regression Subset Selection. R package version 3.1, URL https://CRAN.R-project.org/package=leaps.
- Lüdecke D, Waggoner P, Makowski D (2019). "insight: A Unified Interface to Access Information from Model Objects in R." *Journal of Open Source Software*, **4**(38), 1412.
- Mersmann O (2023). microbenchmark: Accurate Timing Functions. R package version 1.4.10, URL https://CRAN.R-project.org/package=microbenchmark.
- Pinheiro JC, Bates DM (2000). Mixed-Effects Models in S and S-PLUS. Springer, New York.
- Raudenbush SW, Bryk AS (2002). *Hierarchical Linear Models: Applications and data analysis methods*. Second edition edition. Sage, Thousand Oaks CA.

- "Receiver operating characteristic" (2023). "Receiver operating characteristic—Wikipedia, The Free Encyclopedia." [Online; accessed 25-August-2023], URL https://en.wikipedia.org/wiki/Receiver_operating_characteristic.
- Sarkar D (2008). Lattice: Multivariate Data Visualization with R. Springer, New York. ISBN 978-0-387-75968-5. URL http://lmdvr.r-forge.r-project.org.
- Sarkar D, Andrews F (2022). latticeExtra: Extra Graphical Utilities Based on lattice. R package version 0.6-30, URL https://CRAN.R-project.org/package=latticeExtra.
- StataCorp LLC (2023). Stata multilevel mixed-effects reference manual, release 18. Stata Press, College Station TX. URL https://www.stata.com/manuals/me.pdf.
- Vehtari A (2023). "Cross-validation FAQ." URL https://users.aalto.fi/~ave/CV-FAQ. html.
- Venables WN, Ripley BD (2002). *Modern Applied Statistics with S.* Fourth edition edition. Springer, New York.
- Weisberg S (2014). Applied Linear Regression. Second edition edition. Wiley, Hoboken NJ.
- Wickham H, François R, Henry L, Müller K, Vaughan D (2023). dplyr: A Grammar of Data Manipulation. R package version 1.1.2, URL https://CRAN.R-project.org/package=dplyr.
- "Woodbury matrix identity" (2023). "Woodbury matrix identity—Wikipedia, The Free Encyclopedia." [Online; accessed 15-August-2023], URL https://en.wikipedia.org/wiki/Woodbury matrix identity.

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