

Simulating post selection inference

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Introduction

Much of the current research questions on behavioral and social sciences are investigated using statistical models (Flora, 2017). Models are simplified translations of a reality to mathematical expressions; it aims to express how data were generated. Regression-based models are vastly employed in empirical research and often used to estimate causal effects (Berk, 2010). However, to perform causal modeling, researchers must specify a “correct” model (i.e., an accurate model of the data generating process) prior to data collection and use the obtained data only to estimate regression coefficients¹. In practice, researchers usually only have a vague idea of the right model to answer their research questions, or even if such model can be estimated.

To determine which variables should be included in the model, a common solution is to resort to variable selection algorithms. The drawback is that whenever data-driven variable selection procedures are employed, classical inference guarantees are invalidated due to the model itself becoming stochastic (Berk et al., 2013). It means that if the model selection method evaluates the stochastic component of the data, the model is also considered stochastic.

Framing the problem

Variable selection procedures aren’t in themselves problematic. Nevertheless, when the correct model is unknown prior to data analysis, and the *same* dataset is used for I) variable selection, II) parameter estimation, and III) statistical inferences, the estimated results can be highly biased (Berk et al., 2010). This is so because we add a new source of uncertainty when performing model selection. These procedures discards parameter estimates from the model, and the sampling distribution of the remaining regression parameters estimates gets distorted. In addition, we add another source of uncertainty because the selected model isn’t the same across samples.

Consider a well defined population with its unknown regression parameter values. We draw a random sample and apply a model selection procedure. The “best” model found by the variable selection is sample specific and isn’t guarantee to be the correct model (if we assume that such model in fact exists). Suppose we repeat the process of drawing a random sample and performing model selection 10,000 times. In this example there are six possible candidates, and only one correct model. We can simulate the expected frequency in which each of these models is chosen given a probability of 1/3 for the correct model and 1/9 otherwise. As ?? shows, even if the correct model (in this case, \hat{M}_2) is three times more likely to be selected than the competing models, it is expected to be chosen more frequently but not at the majority of the time. Therefore, the expected selected model is an incorrect one.

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¹see Berk (2010) for further discussion.

Table 1: A table of the first 10 rows of the mtcars data.

Model	Frequency
M1	0.1245
M2	0.3735
M3	0.1287
M4	0.1201
M5	0.1258
M6	0.1274

To understand why the regression parameters estimates might be biased, recall that in a multiple regression we estimate *partial regression coefficients*: in a regression equation, the weight of independent variables are estimated in relation to the other independent variables in the model (Cohen & Cohen, 1983). For a dependent variable, Y , predicted by variables X_1 and X_2 , $B_{Y1.2}$ is the partial regression coefficient for Y on X_1 holding X_2 constant, and $B_{Y2.1}$ is the partial regression coefficient for Y on X_2 holding X_1 constant. This regression equation is written as:

$$\hat{Y} = B_{Y0.12} + B_{Y1.2}X_1 + B_{Y2.1}X_2 + \varepsilon \quad (1)$$

where $B_{Y0.12}$ is the model intercept when X_1 and X_2 are held constant, and ε is the error term.

The regression coefficient for X_i , for $i = \{1, 2\}$, is model dependent. The equations for the regression coefficients for X_1 ($B_{Y1.2}$) and X_2 ($B_{Y2.1}$) illustrate why:

$$B_{Y1.2} = \frac{\rho_{Y1} - \rho_{Y2}\rho_{12}}{(1 - \rho_{12}^2)} \times \frac{\sigma_Y}{\sigma_1} \quad (2)$$

$$B_{Y2.1} = \frac{\rho_{Y2} - \rho_{Y1}\rho_{21}}{(1 - \rho_{21}^2)} \times \frac{\sigma_Y}{\sigma_2} \quad (3)$$

Here ρ stands for the populational correlation coefficient and σ for the populational standard deviation. Unless we have uncorrelated predictors (i.e. $\rho_{12} = 0$), the value for any of the regression coefficients is determined by which other predictors are in the model. If either one is excluded from the model, a different regression coefficient will be estimated: excluding X_2 , for example, would zero all the correlations involving this predictor, leaving,

$$B_{Y1.2} = \frac{\rho_{Y1} - 0 \times 0}{(1 - 0^2)} \times \frac{\sigma_Y}{\sigma_1} = \rho_{Y1} \times \frac{\sigma_Y}{\sigma_1} = B_{Y1} \quad (4)$$

Berk et al. (2010) warns that the sampling distribution of the estimated regression parameters is distorted because estimates made from incorrect models will also be included, resulting in a mixture of distributions. Therefore, the model selection process must be taken into account in the regression estimation whenever it is applied.

Illustration

We can illustrate this discussion expanding with simulations an analytic example given by Berk et al. (2010). Consider a model for a response variable y with two potential regressors, x and z . Say we are interested in the relationship between y and x while holding z constant, that is, $\hat{\beta}_{yx \cdot z}$. Framing this as a linear regression model we have

$$y_i = \beta_0 + \beta_1 x_i + \beta_2 z_i + \varepsilon_i \quad (5)$$

Now, suppose that we're in a scenario where $\rho_{xz} = 0.5$, both β_1 and β_2 are set to 1 and $\varepsilon \sim N(0, 10)$. We'll use a sample size of 250 subjects, and 1,000 random samples will be drawn from this population. We also calculate *coverage* and *bias* for each regressor. Coverage informs the frequency in which the true coefficient value is captured by the 95% confidence interval (CI) of the estimate. The bias of the estimations is calculated as $\frac{1}{R} \sum (\hat{\theta}_r - \theta)$, where R is the number of repetitions, θ represents a population parameter, and $\hat{\theta}_r$ a sample estimate in each simulation.

Table 2: Coverage and bias for the regression coefficients assuming x and z are always included in the model.

Predictor	Coverage	Bias
x	0.953	0.0036217
z	0.942	-0.0048216

In this first scenario $\beta_{yx \cdot z}$ is estimated assuming x and z are always included in the model. However, we should not expect the same results if by model selection we arrive at a model where z is excluded. As indicated in equation 4, in this case, any correlation where z is involved is equivalent to zero, and β_1 is simply,

$$\beta_{yx} = \rho_{xy} \left(\frac{\sigma_y}{\sigma_x} \right) \quad (6)$$

Note that β_{yx} is not the same as $\beta_{yx \cdot z}$. If we do not have a model specified prior to data collection and analysis, it is not clear which definition of regression parameter for x we're trying to estimate – if $\beta_{yx \cdot z}$, as exemplified on Equation 2, or if β_{yx} from Equation 6. In short, the definition of $\hat{\beta}_1$ depends on the model in which it is placed.

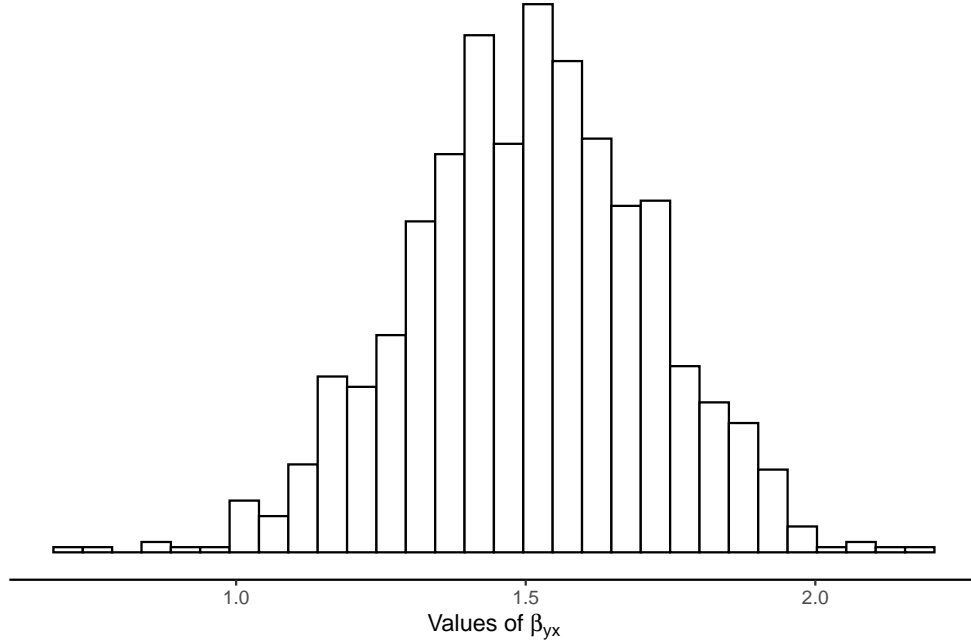


Figure 1: Sampling distributions of the regression coefficient for regressor x in a reduced model where z is excluded.

It is easy to notice a strong bias in the estimates for x 's coefficients when we fit our model as $y \sim x$. Under these conditions, we should expect a coverage of the true coefficient value of x of 0.319 and a bias of 0.3.

Conversely, under the correct model bias is negligible and coverage follows the Type I error rate of 5% that we've set for this exercise.

Discarding z from our model has distorted the sampling distribution of x . Thus, as Berk et al. (2010, p. 221) puts it: “when a single model is not specified before the analysis begins, it is not clear what population parameter is the subject of study. And without this clarity, the reasoning behind statistical inference becomes obscure”.

Other issues

The previous example has shown that if two predictors are moderately correlated and one is dropped from the model, our estimation will be biased. But what else should we consider? If we are selecting the variable every time, we should find no issues. Consider the equation for the standard error of the regression coefficient estimated in the example of model selection,

$$SE(\beta_{yx \cdot z}) = \frac{\hat{\sigma}_\varepsilon}{s_x \sqrt{n-1}} \sqrt{\frac{1}{1 - r_{xy}^2}} \quad (7)$$

Here $\hat{\sigma}_\varepsilon$ is an estimate of the residual standard deviation, s_x is the sample standard deviation of x , r_{xy}^2 is the square of the sample correlation between x and z , and n is the sample size. If the standard error for a regression coefficient is large, it means that its distribution will be more dispersed. So, from this equation, we identify as crucial parameters for a wider sampling distribution: larger residual variance, less variance in x , smaller sample size, and, stronger correlation between regressors and the response variable.

In the following sections we'll use simulated data to understand the extent of the problems that arise when model selection, parameters estimation and statistical inferences are undertaken with the same data set. A Shiny Web app was developed to address the issues with model selection outlined in this manuscript. The app runs Monte Carlo simulations based on parameters set by the user via a point-and-click interface. For each specified regression coefficient, the results table offer the average estimate value, its 95% Confidence Interval coverage, the average bias and the average mean squared error. Users can access the app online at marwin.shinyapps.io/model-selection-app/.

Method

For the purposes of this manuscript we use the stepwise approach for model selection with *Akaike Information Criterion* (AIC) as model selection criterion. But it is worth noting that the concerns raised here are irrespective of the chosen model selection method (Berk et al., 2010). The aforementioned app also provide the option to choose BIC or Mallows' Cp as selection method. In addition to bias and coverage, we'll also estimate the average Mean Squared Error (MSE), calculated as $\frac{1}{R} \sum [(\hat{\theta}_r - \theta)^2]$.

Explicar a função de simulação

Results

Noise

To express variability we can use a signal-to-noise ratio (SNR), defined here as:

$$\frac{S}{N} = \mathbf{b} \Sigma \mathbf{b} \sigma^{-2} \quad (8)$$

where, \mathbf{b} is a $(p + 1)$ vector of regression coefficients, Σ is the covariance matrix of predictors and σ^2 is the error term variance. Note that “model selection bias also occurs when an explanatory variable has a weak relationship with the response variable. The relationship is real, but small. Therefore, it is rarely selected

as significant” (Lukacs et al., 2009, p. 118). We can experiment with a range of values for the SNR to see how it affects the estimates. We’ll use most parameters set in our first simulation exercise and SNR values ranging from 0.1 to 2. One thousand (1,000) simulations will be run for each of those values.

Table 3: Values for estimate, coverage, bias, and MSE for both predictors with varying SNR values.

SNR	N	Predictor	Estimate	Coverage	Bias	MSE
0.01	250	x1	2.2990007	0.283	1.2990007	2.9643164
0.01	250	x2	2.2570703	0.283	1.2570703	3.0839713
0.10	250	x1	1.1426579	0.813	0.1426579	0.1434384
0.10	250	x2	1.1233778	0.786	0.1233778	0.1458715
0.50	250	x1	0.9962085	0.950	-0.0037915	0.0336320
0.50	250	x2	1.0018315	0.950	0.0018315	0.0330238
1.00	250	x1	1.0039931	0.952	0.0039931	0.0156801
1.00	250	x2	0.9953709	0.970	-0.0046291	0.0147253
2.00	250	x1	1.0025169	0.942	0.0025169	0.0083488
2.00	250	x2	1.0037286	0.944	0.0037286	0.0082008

As expected, as the noise ratio gets larger, the selected model includes the true coefficient at lower frequencies. Because model selection interferes with the true model composition, coefficient estimates deviate from its true value. In this particular setting, with $\text{SNR} \geq 0.5$, coverage frequency approximates 95% and the estimates get closer to their true value. Measures of bias and MSE are also useful to display how our uncertainty gets smaller with less variability in the data.

Sample size

Once we know Equation 7 it is expected that larger samples produce smaller standard errors for the regression coefficients. We can confirm this intuition using simulated data again, but this time varying sample sizes.

Table 4: Values for estimate, coverage, bias, and MSE for both predictors with varying sample sizes.

SNR	N	Predictor	Estimate	Coverage	Bias	MSE
0.5	10	x1	1.9262739	0.396	0.9262739	1.9822681
0.5	10	x2	1.7230094	0.403	0.7230094	1.7787062
0.5	50	x1	1.1539985	0.798	0.1539985	0.1508404
0.5	50	x2	1.1451933	0.770	0.1451933	0.1593684
0.5	100	x1	1.0323179	0.937	0.0323179	0.0789976
0.5	100	x2	1.0277209	0.929	0.0277209	0.0771598
0.5	200	x1	0.9988481	0.956	-0.0011519	0.0399842
0.5	200	x2	1.0046174	0.957	0.0046174	0.0392427
0.5	500	x1	1.0056553	0.943	0.0056553	0.0163749
0.5	500	x2	0.9970542	0.949	-0.0029458	0.0160090

Candidate predictors

So far we’ve seen that with 2 predictors, each with true coefficient values of 1 and a correlation of 0.5, we get more precise estimates when $\text{SNR} \geq 0.5$ and $n \geq 100$. The number of candidate predictor variables and its covariance matrix are two important aspects not addressed yet. To demonstrate their influence we can run simulations with varying values for each, where each case will have a different combination of number of predictors and correlation between them. For simplicity, all the off-diagonal elements the correlation matrix

of predictors will be equal, meaning that every predictor in the full model is correlated with the others by the same degree.

To keep the computing load at reasonable levels, we'll limit the length of vectors of p and ρ to 10 values each. The grid for ρ ranges from 0.1 to 0.9 and we'll increase p from 2 to 10. We'll set $\text{SNR} = 0.5$, $n = 100$ and set all true coefficient values to 1. Again, we're using AIC to select the best model, and performing 1,000 simulation replicates.

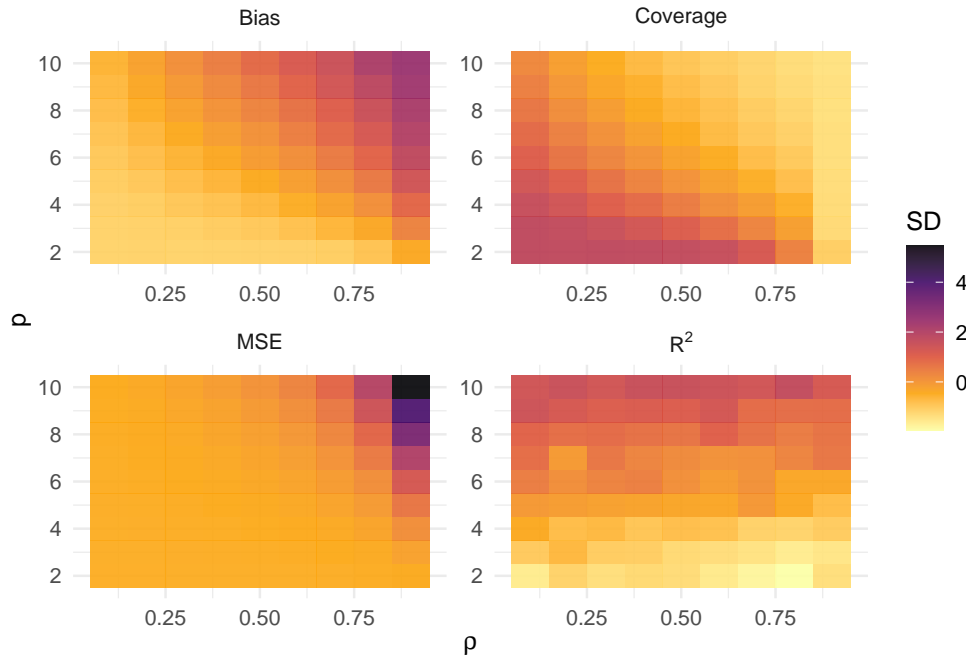


Figure 2: Standardized measures of Bias, Coverage, Mean Square Error (MSE) and R^2 , across each simulation of pairs of number of predictors and correlation values

The horizontal axis values of Figure 2 were standardized to keep all four plots in the same scale. In this heat map, darker colors represent larger values (greater standard deviation). The pattern for Bias and MSE is similar: performing model selection with a great number of highly correlated predictors is likely to produced highly biased estimates. As expected, Coverage follows an inverted pattern from Bias and MSE: highly biased estimates fall out of the 95% CI coverage.

This example illustrates that our best case scenario is to have few, weakly correlated predictors. Of course, this is a setting rarely seen in empirical research – what's the point of going through model selection if you have only a handful of variables? Of notice here is that bias in estimates are likely to get bigger if model selection is naively performed with little prior information about the data generating system, to filter out candidate models. We have seen that sample size, SNR, number of predictors, and correlation between parameters with themselves and with the response variable, are characteristics of the model that each in their own way contributes to producing biased estimates.

Visualizing distributions

We've discussed earlier how dropping a predictor from the model can distort the coefficient sampling distribution of the remaining ones when they're not orthogonal. However, even when the preferred (or "correct") model is selected, there is no guarantee about obtaining sound regression coefficient estimates. In this final example, we'll show that if by chance the "correct" model is achieved by model selection, the

sampling distributions of the resulting regression coefficients might be different whether we condition on arriving at the correct model or on the correct model being known in advance.

Consider this example from Berk et al. (2010). As with the other examples, we'll implement forward stepwise regression using the AIC as a fit criterion. The full regression model takes the form of

$$y_i = \beta_0 + \beta_1 w_i + \beta_2 x_i + \beta_3 z_i + \varepsilon_i \quad (9)$$

where $\beta_0 = 3.0$, $\beta_1 = 0.0$, $\beta_2 = 1.0$, and $\beta_3 = 2.0$. The variances and covariance are set as: $\sigma_\varepsilon^2 = 10.0$, $\sigma_w^2 = 5.0$, $\sigma_x^2 = 6.0$, $\sigma_z^2 = 7.0$, $\sigma_{w,x} = 4.0$, $\sigma_{w,z} = 5.0$, and $\sigma_{x,z} = 5.0$. The sample size is 200.

In their article Berk et al. (2010) uses the term *preferred* model instead of *correct* model. They do so because a model that excludes W can also be called correct the same as one like Equation 9, as long as $\beta_1 = 0$ is allowed. To be consistent with the original text, we'll use *preferred* to refer to the model with W excluded. This model is preferred because it generates the same conditional expectations for the response using up one less degree of freedom. The plots show the regression estimates t -values. "A distribution of t -values is more informative than a distribution of regression coefficients because it takes the regression coefficients and their standard errors into account" (Berk et al., 2010, p. 266).



Figure 3: Stepwise regression sampling distributions of the regression coefficient t -values for regressors X and Z . Red density plot is conditional on the preferred model being known. The blue density plot is conditional on the regressor being included in a model

In both Figures 3 and 4 the red density plot represents the regression estimates t -values distribution when no model selection is performed. In Figure 3 the blue density plot shows the distributions when either X or Z are included in the model. For Figure 4 the blue distribution refers to distributions of either X or Z t -values when the preferred model is selected.

The contrast between red and blue curves is apparent. The difference that is the most striking is in t -values distributions post-model-selection for regressor Z when conditioned on the regressor being included in a model. This curve displays a bimodal distribution and highly biased mean and standard deviation, as summarized below.



Figure 4: Stepwise regression sampling distributions of the regression coefficient t^* -values for regressors X and Z. Red density plot is conditional on the preferred model being known. The blue density plot is conditional on the preferred model being selected

Table 5: Mean and standard deviation for coefficients for x and z given four different model conditions.

Model	M_x	σ_x	M_z	σ_z
Full model	2.236	1.028	4.731	1.066
Preferred model	2.554	0.772	4.502	0.925
X included	2.562	0.807	4.502	0.925
Z included	2.554	0.772	5.615	2.459

It is especially telling observing those plots that the assumed underlying distribution can be very different from what is obtained. Statistical inference performed in such scenarios would be misleading. Figure 4 confirms that conditioning on arriving at the preferred model does not guarantee trustable estimates.

Conclusion

Model selection methods are routine in research on the social and behavioral sciences, and commonly taught in applied statistics courses and textbooks. However, little is mentioned about the biased estimates obtained when such procedures are carried out. With simulated data we have identified specific characteristics of the data generating model that can potentially increase the bias in estimates obtained through variable selection. We can conclude that post-model-selection sampling distribution can deviate greatly from the assumed underlying distribution, even when the best representative model of the data generation process has been selected.

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