title: "Biostat 561: Homework 9"

author: "Instructor: Amy Willis, Biostatistics, UW"

date: "29 May, 2019"

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Homework 9 due June 5, 3:15pm

No office hours Mon 3 June due to travel; special office hours Tuesday 4 June, 2:30pm

Link to Homework 9 submission: https://classroom.github.com/a/iAetmO52

In this homework, we will continue to explore the coverage of confidence intervals for  $\beta_1$  under the model  $Y_i = \beta_0 + \beta_1 x_i + \epsilon_i$ , contrasting confidence intervals created using model-based standard errors  $(var(\hat{\beta}) = \hat{\sigma}^2(X^TX)^{-1})$  and confidence intervals created using White's heteroskedasticity-robust "sandwich" standard errors  $(var(\hat{\beta}) = (X^TX)^{-1}X^Tdiag(e_i^2)X(X^TX)^{-1})$ .)

We will simulate from the model

$$Y_i \mid X_i, u_i = \beta_1 X_i + \epsilon_i$$
, where  $X_i \stackrel{i.i.d.}{\sim} N(0,1)$ ,  $u_i \stackrel{i.i.d.}{\sim} N(0,1)$ , independent of the  $X_i$ ,  $\epsilon_i = f(X_i) \times u_i$ .

## Question 1: R scripts to assess coverage from the command line

Adapt the R scripts from class to compare the coverage of 95% confidence intervals for  $\beta$  using both model-based and heteroskedasticity-consistent standard errors. Your script should take arguments n (the sample size),  $\beta$  (the true value of  $\beta$  to simulate under), and option (which has value either 1 or 2 for the form of f(x), where  $f_1(x) = 1 + x^2$  and  $f_2(x) = 2 - x^2$ ), seed (the starting seed) and reps (the number of replicates to use). Use the package sandwich to compute the heteroskedasticity-consistent standard errors.

Show the output of

Rscript qtn1\_response.R n=100 beta=1 option=1 seed=123 reps=1000

and

Rscript qtn1 response.R n=100 beta=1 option=2 seed=123 reps=1000

What coverage do you find based on each method?

How long did your scripts take to run?

## Question 2: Running R scripts on the clusters

Time to use the cluster! Use the examples from lecture to guide your script-writing in this question.

We want to investigate the effect of n on the coverage of the two procedures for the two data-generating mechanisms  $f_1(x)$  and  $f_2(x)$ . We want to run 5000 simulations per n and per option, and look at  $n \in \{50, 100, 150, \ldots, 500\}$ .

Adapt your response to question 1 to write an R script with one argument, reps (the number of replicates to use) that runs reps replicates for option 1 and 2, and all  $n \in \{50, 100, 150, \dots, 500\}$ . Confirm it works for small reps. Call this script qtn2\_response.R.

Write a shell script called call\_qtn2.sh that calls qtn2\_response.R. qtn2.sh should be similar to call\_sim\_robust\_se.sh, seen in class, and have arguments sim-name, nreps-total and nreps-per-job.

Adapt submit\_sim\_robust\_se.sh to create run\_qtn2.sh to run your simulation study on the cluster. Check it works locally for a small simulation.

Connect to bayes.biostat.washington.edu using your favourite ssh client (see Homework 8). Run  $run_qtn2.sh$  on the bayes cluster to run 5000 reps per option and n. Split each simulation into either 5 or 10 jobs (i.e., not 1 job and not 5000 jobs). Perform this batch submission either using a loop in your shell script, or by using a job array.

Present your results graphically and/or tabularly in a way that best illustrates your findings. Comment on what you see.

Optional but recommended for 533 students: Explain why you get the results you see.

Upload your script(s) to your github repository along with a pdf containing a write up of your results.