Simple Example

2022-11-23

Goal of this doc

To simulate a really simple case to try to understand the problem with increasing validation sizes with the Wang et al approach. I'm going to dramatically simplify things to try to make it easier to follow.

Load some libraries we will need

Attaching package: 'broom'

```
library(caret)
## Loading required package: ggplot2
## Loading required package: lattice
library(ggplot2)
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
library(magrittr)
library(randomForest)
## randomForest 4.7-1.1
## Type rfNews() to see new features/changes/bug fixes.
## Attaching package: 'randomForest'
## The following object is masked from 'package:dplyr':
##
##
       combine
## The following object is masked from 'package:ggplot2':
##
##
       margin
library(splines)
library(modelr)
library(broom)
```

```
## The following object is masked from 'package:modelr':
##
##
       bootstrap
library(tidyr)
##
## Attaching package: 'tidyr'
## The following object is masked from 'package:magrittr':
##
##
       extract
library(gam)
## Loading required package: foreach
## Loaded gam 1.22
library(patchwork)
library(here)
```

Simulate one very simple case

here() starts at /cloud/project

Let's remove all the bootstrapping and complicated data generation just so we can understand the global behavior of different operations. First I'll just simulate a really simple case:

```
## Set sample sizes with imbalance
n_{train} = 200
n_{\text{test}} = 200
n_valid = 10000
n_tot = n_train + n_test + n_valid
## Make some covariates
x1 = rnorm(n_tot)
x2 = rnorm(n_tot)
x3 = rnorm(n_tot)
err = rnorm(n tot)
## Make an outcome
y = 2*x1 + 3*x2^2 + 0.5*x3^3 + err
## Create an indicator of training, testing, or validation
set_ind = c(rep("train",n_train),
            rep("test",n_test),
            rep("valid",n_valid))
## Build the data set
dat = data.frame(y,x1,x2,x3,set_ind)
```

Fit a prediction model

Ok now lets get the predictions - filter to the training

```
## Build the predictor and make the predictions

train_dat = dat %>%
  filter(set_ind == "train") %>%
  select(y,x1,x2,x3)

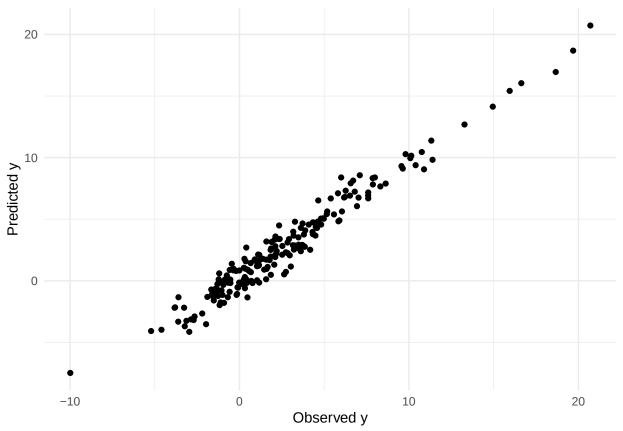
pred_mod = gam(y ~ s(x1) + s(x2) + s(x3))
```

Make the predictions for all the values (note we will only use the predictions in testing and validation data sets):

```
dat$pred = predict(pred_mod,dat)
```

Let's make sure the prediction looks good:

```
## Plot only on the test set
dat %>%
  filter(set_ind=="test") %>%
  ggplot(aes(x=y,y=pred)) +
  geom_point() + theme_minimal() +
  xlab("Observed y") +
  ylab("Predicted y")
```



Let's now do one bootstrap iteration of the Wang et al approach at a time

First we get the relationship model. Here we use a gam (like we do in Sara's code) because we observe that the relationship between the predicted and observed y's doesn't look purely linear.

```
## Filter to the test set
test_dat = dat %>%
    filter(set_ind == "test")

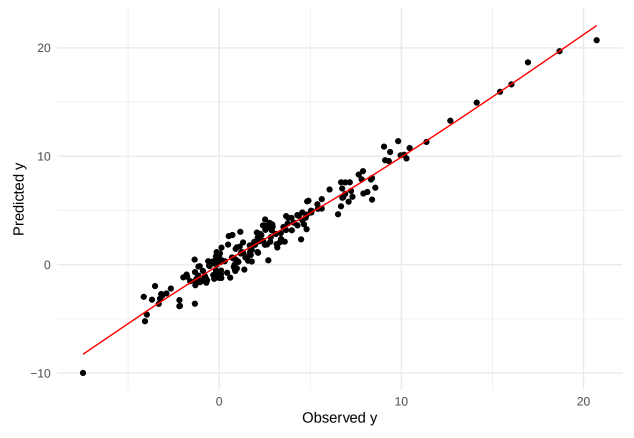
## Fit the relationship model

rel_mod = lm(y ~ ns(pred,df=3),data=test_dat)

## Get the residual variance for use in simulating
sigma2 = sigma(rel_mod)^2
```

Let's make sure the relationship model looks ok:

```
test_dat %>%
  add_predictions(rel_mod,var="fitted") %>%
  arrange(y) %>%
  ggplot(aes(x=pred,y=y)) +
  geom_point() + theme_minimal() +
  xlab("Observed y") +
  ylab("Predicted y") +
  geom_line(aes(x=pred,y=fitted),col="red")
```



Now let's do a step of the simulation for the validation set

I'm going to follow the Wang algorithm directly here.

Step 0. Subset to validation data

```
valid_dat = dat %>%
  filter(set_ind == "valid")
```

Step 1. Sample with replacement from x's

This may not be a strictly necessary step, but we do it in the boostrap iterations so doing this for completeness

```
sim_dat = valid_dat %>% sample_n(size=n_valid,replace=TRUE)
```

Step 2. Sample y's from the relationship model

This is where I could imagine something went wrong in Sara's code (I was having trouble understanding it myself) but basically we need to sample from the regression model for the predicted values:

Step 3. Compute coefficients

Let's compare coefficients and sds from the different models

```
lm(y ~ x1,data=valid_dat) %>% tidy()
## # A tibble: 2 x 5
##
     term
                 estimate std.error statistic p.value
     <chr>
                    <dbl>
                               <dbl>
                                         <dbl>
## 1 (Intercept)
                     2.99
                              0.0485
                                          61.7
                                                      0
## 2 x1
                      2.07
                              0.0481
                                          43.1
                                                      0
## Naive Post-PI
lm(pred ~ x1,data=valid_dat) %>% tidy()
## # A tibble: 2 x 5
##
     term
                 estimate std.error statistic p.value
##
     <chr>>
                    <dbl>
                               <dbl>
                                          <dbl>
                                                  <dbl>
                      2.99
                                          67.8
## 1 (Intercept)
                              0.0441
                                                      0
## 2 x1
                      2.07
                              0.0437
                                          47.5
                                                      0
## Post-PI via simulated data
lm(y_sim ~ x1,data=sim_dat) %>% tidy()
## # A tibble: 2 x 5
##
     term
                 estimate std.error statistic p.value
     <chr>
                    <dbl>
                               <dbl>
                                          <dbl>
                                                  <dbl>
                                          65.5
## 1 (Intercept)
                     2.89
                              0.0441
                                                      0
                      2.09
                              0.0442
                                          47.3
## 2 x1
                                                      0
```

Let's do the same thing for many simulations just to make sure we understand what's going on here

```
## Set the number of replications
n_rep = 500
```

```
## Set sample sizes with imbalance
n_{train} = 200
n_{test} = 200
n_valid = 2000
n_tot = n_train + n_test + n_valid
beta_wang = beta_naive = beta_oracle = se_wang = se_naive = se_oracle = rep(NA,n_rep)
for(k in 1:n_rep){
  ## Make some covariates
 x1 = rnorm(n_tot)
  x2 = rnorm(n_tot)
 x3 = rnorm(n_tot)
  err = rnorm(n_tot,sd=4)
  ## Make an outcome
  y = 2*x1 + 3*x2^2 + 0.5*x3^3 + err
  ## Create an indicator of training, testing, or validation
  set_ind = c(rep("train",n_train),
            rep("test",n_test),
            rep("valid",n_valid))
  ## Build the data set
  dat = data.frame(y,x1,x2,x3,set_ind)
  ## Fit the model
  train_dat = dat %>%
  filter(set_ind == "train") %>%
  select(y,x1,x2,x3)
  pred_mod = gam(y \sim s(x1) + s(x2) + s(x3))
  ## Add the predictions
  dat$pred = predict(pred_mod,dat)
  ## Filter to the test set
  test dat = dat %>%
   filter(set_ind == "test")
  ## Fit the relationship model
  rel_mod = lm(y ~ ns(pred, df=3), data=test_dat)
  ## Get the residual variance for use in simulating
  sigma2 = sigma(rel_mod)^2
  ## Filter to the validation set
  valid_dat = dat %>%
    filter(set_ind == "valid")
```

```
## Get the simulated data
  sim_dat = valid_dat %>% sample_n(size=n_valid,replace=TRUE)
  ## Sample simulated y's
  y_sim = rnorm(n_valid,
              mean=sim_dat$pred,
              sd = sqrt(sigma2))
  sim_dat = cbind(sim_dat,y_sim)
  ## Fit the models
  lm_oracle = lm(y ~ x1,data=valid_dat)
  lm_naive = lm(pred ~ x1,data = valid_dat)
  lm_wang = lm(y_sim ~ x1,data=sim_dat)
  ## Get the coefficients
  beta_oracle[k] = lm_oracle$coefficients[2]
  beta_naive[k] = lm_naive$coefficients[2]
  beta_wang[k] = lm_wang$coefficients[2]
  ## Get the parametric ses
  se_oracle[k] = summary(lm_oracle)$coefficients[2,2]
  se_naive[k] = summary(lm_naive)$coefficients[2,2]
  se_wang[k] = summary(lm_wang)$coefficients[2,2]
 cat(k)
}
```

1234567891011121314151617181920212223242526272829303132333435363738394041424344454647484950515253545

Let's look at what the coverage looks like

```
results = data.frame(beta_oracle,se_oracle,beta_wang,se_wang,beta_naive,se_naive)

results = results %>%
    mutate(up_oracle = beta_oracle + 1.96*se_oracle,
        low_oracle = beta_oracle - 1.96*se_oracle,
        cov_oracle = ((up_oracle > 2) & (low_oracle < 2)),

    ## Wang
    up_wang = beta_wang + 1.96*se_wang,
    low_wang = beta_wang - 1.96*se_wang,
    cov_wang = ((up_wang > 2) & (low_wang < 2)),

## Naive
    up_naive = beta_naive + 1.96*se_naive,
    low_naive = beta_naive - 1.96*se_naive,
    cov_naive = ((up_naive > 2) & (low_naive < 2)),
)</pre>
```

Make some plots

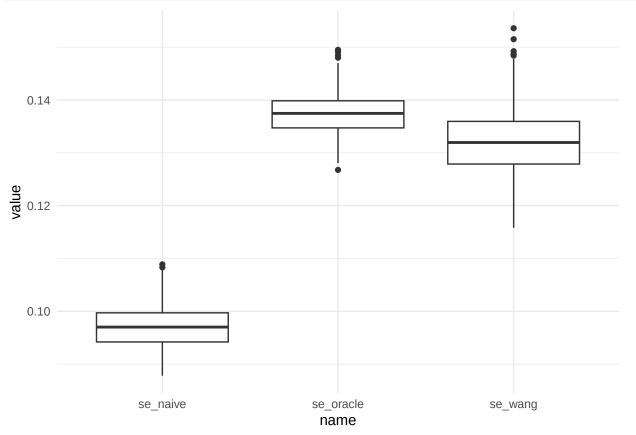
Coverage is improved from naive, but not as good as oracle

```
p1 = results %>%
arrange(beta_naive) %>%
```

```
ggplot(aes(y = 1:n_rep)) + geom_point(aes(x=beta_naive,y=1:n_rep)) +
  geom_linerange(aes(xmin=low_naive, xmax=up_naive,col=cov_naive)) +
  theme_minimal() +
  geom_vline(xintercept=2) +
  xlim(-1,5)
p2 = results %>%
  arrange(beta_wang) %>%
  ggplot(aes(y = 1:n_rep)) + geom_point(aes(x=beta_wang,y=1:n_rep)) +
  geom_linerange(aes(xmin=low_wang, xmax=up_wang,col=cov_wang)) +
  theme_minimal() +
  geom_vline(xintercept=2) +
  xlim(-1,5)
p3 = results \%
  arrange(beta_oracle) %>%
  ggplot(aes(y = 1:n_rep)) + geom_point(aes(x=beta_oracle,y=1:n_rep)) +
  geom_linerange(aes(xmin=low_oracle, xmax=up_oracle,col=cov_oracle)) +
  geom_vline(xintercept=2) +
  theme_minimal() +
  xlim(-1,5)
p1 + p2 + p3
                                                              500
  500
                                500
  400
                                400
                                                              400
  300
                                300
                                                              300
                cov_naive
                                              cov_wang
                                                                            cov_oracle
                    FALSE
                                                  FALSE
                                                                               FALSE
                    TRUE
                                                  TRUE
                                                                               TRUE
  200
                                200
                                                              200
  100
                                100
                                                              100
    0
                                  0
                                                                0
                                     0 2 4
                                                                  0 2 4
       0 2 4
    beta_naive
                                  beta_wang
                                                               beta_oracle
```

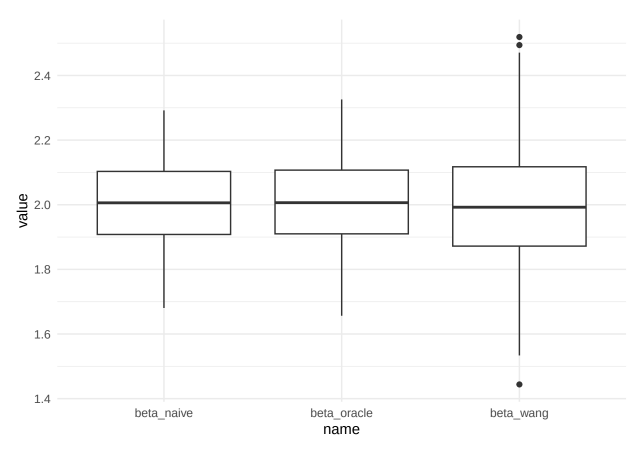
Standard errors are closer

```
results %>%
  select(se_naive,se_wang,se_oracle) %>%
  pivot_longer(1:3) %>%
  ggplot(aes(x=name,y=value)) +
  geom_boxplot() +
  theme_minimal()
```

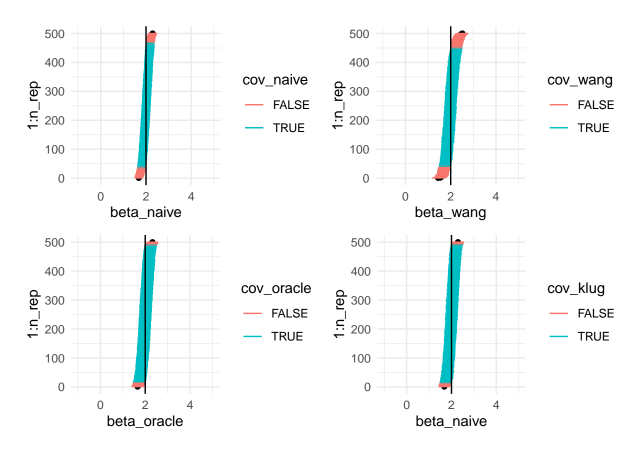


There are too many "big" values of the estimate

```
results %>%
  select(beta_naive,beta_wang,beta_oracle) %>%
  pivot_longer(1:3) %>%
  ggplot(aes(x=name,y=value)) +
  geom_boxplot() +
  theme_minimal()
```



What if we just replace the estimate with the naive estimate, but use the Wang SE (klugey version of what the bootstrap would do)



Look at coverage

0.86

1

Looks like the klug works in this one random case :).

0.818

0.956