

p8130_hw5_xy2395

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11/30/2018

```
knitr::opts_chunk$set(echo = TRUE)
library(tidyverse)
library(faraway)
library(patchwork)
library(funModeling)
library(broom)
library(leaps)
library(modelr)
library(gggridges)
set.seed(1)
```

Dataset Description

state.x77 is a matrix with 50 rows and 8 columns giving the following statistics in the respective columns:

- Population: population estimate as of July 1, 1975.
- Income: per capita income (1974).
- Illiteracy: illiteracy (1970, percent of population).
- Life Exp: life expectancy in years (1969–71).
- Murder: murder and non-negligent manslaughter rate per 100,000 population (1976).
- HS Grad: percent high-school graduates (1970).
- Frost: mean number of days with minimum temperature below freezing (1931–1960) in capital or large city.
- Area: land area in square miles.

```
# Load the data
state_df =
  state.x77 %>%
  as.tibble() %>%
  janitor::clean_names() %>%
  select(life_exp, everything())
```

Part 1: Exploratory Data Analysis

Descriptive Statistics

Build a function to generate descriptive statistics for continuous variables.

```
summary_continuous = function(variable){
  data_frame(
    mean = mean(variable),
    sd = sd(variable),
    median = median(variable),
    maximum = max(variable),
    minimum = min(variable),
```

```

    IQR = IQR(variable)
  )
}

```

Descriptive Statistics for all the variables are shown below. All the variables are continuous.

```

# Generate descriptive statistics
map(state_df, summary_continuous) %>%
  bind_rows() %>%
  mutate(variable = names(state_df)) %>%
  select(variable, everything()) %>%
  knitr::kable(digits = 2,
    caption = "Descriptive statistics of continuous variables")

```

Table 1: Descriptive statistics of continuous variables

variable	mean	sd	median	maximum	minimum	IQR
life_exp	70.88	1.34	70.67	73.6	67.96	1.78
population	4246.42	4464.49	2838.50	21198.0	365.00	3889.00
income	4435.80	614.47	4519.00	6315.0	3098.00	820.75
illiteracy	1.17	0.61	0.95	2.8	0.50	0.95
murder	7.38	3.69	6.85	15.1	1.40	6.32
hs_grad	53.11	8.08	53.25	67.3	37.80	11.10
frost	104.46	51.98	114.50	188.0	0.00	73.50
area	70735.88	85327.30	54277.00	566432.0	1049.00	44177.25

Also show the correlation matrix to check potential correlations. `life_exp` and `murder` are highly correlated. `illiteracy` and `murder` are correlated as well.

```

# generate the correlation matrix
cor(state_df) %>%
  knitr::kable(digits = 2,
    caption = "Correlation matrix for all variables")

```

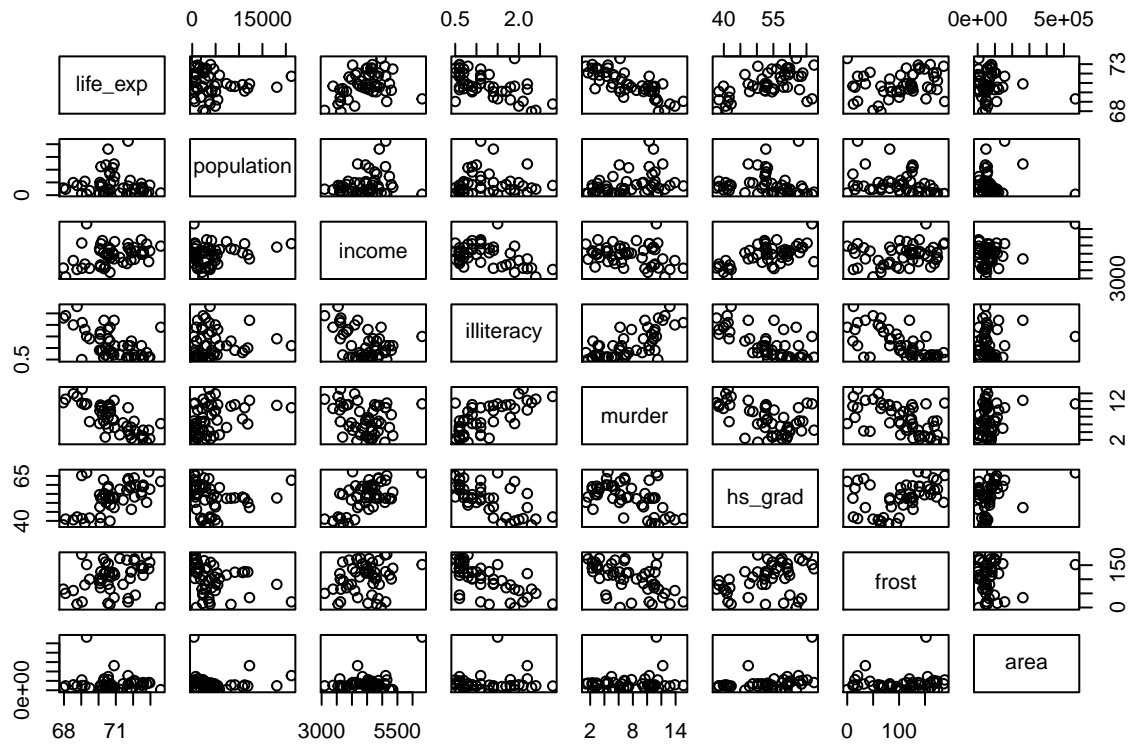
Table 2: Correlation matrix for all variables

	life_exp	population	income	illiteracy	murder	hs_grad	frost	area
life_exp	1.00	-0.07	0.34	-0.59	-0.78	0.58	0.26	-0.11
population	-0.07	1.00	0.21	0.11	0.34	-0.10	-0.33	0.02
income	0.34	0.21	1.00	-0.44	-0.23	0.62	0.23	0.36
illiteracy	-0.59	0.11	-0.44	1.00	0.70	-0.66	-0.67	0.08
murder	-0.78	0.34	-0.23	0.70	1.00	-0.49	-0.54	0.23
hs_grad	0.58	-0.10	0.62	-0.66	-0.49	1.00	0.37	0.33
frost	0.26	-0.33	0.23	-0.67	-0.54	0.37	1.00	0.06
area	-0.11	0.02	0.36	0.08	0.23	0.33	0.06	1.00

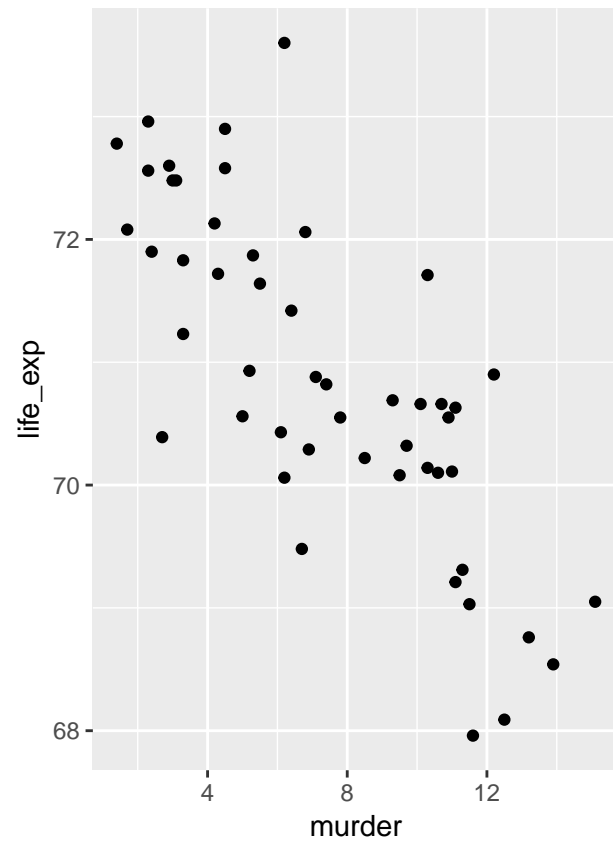
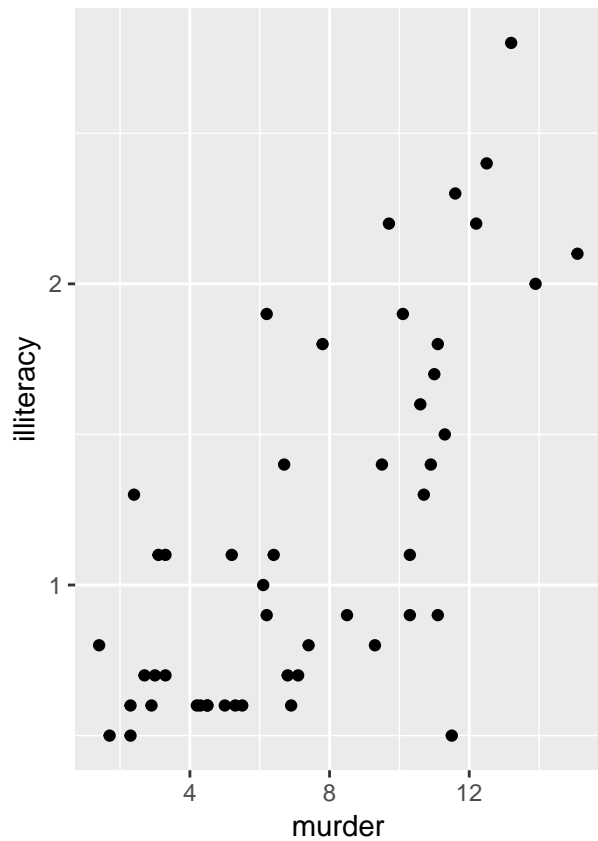
Plots

First, plot the highly correlated variables.

```
plot(state_df)
```



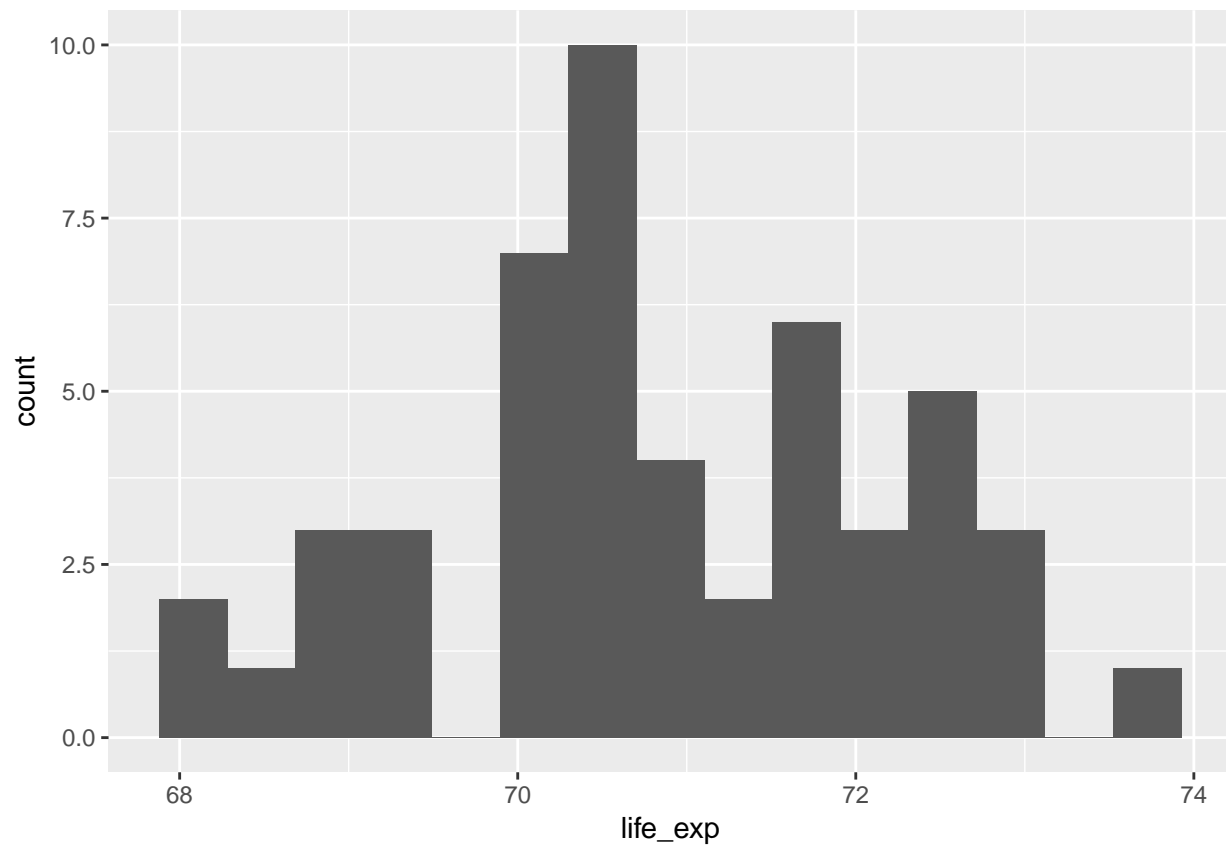
```
murder_illiteracy_points =
state_df %>%
  ggplot(aes(x = murder, y = illiteracy)) +
    geom_point()
murder_lifeexp_points =
state_df %>%
  ggplot(aes(x = murder, y = life_exp)) +
    geom_point()
murder_illiteracy_points + murder_lifeexp_points
```



life_exp and murder are highly correlated. illiteracy and murder are correlated as well.

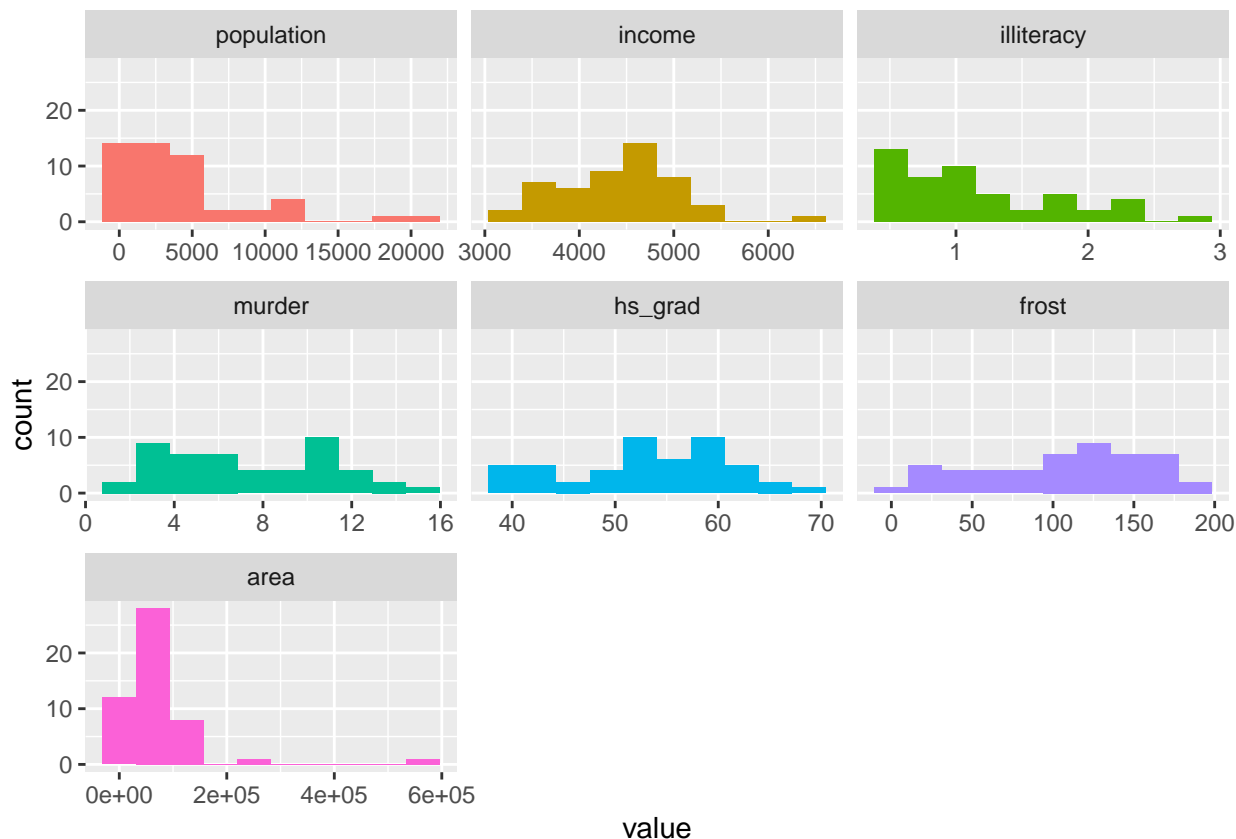
Then check the normality of the outcome, life_exp. Life expectancies of the states are approximately normally distributed .

```
state_df %>%
  ggplot(aes(x = life_exp)) +
  geom_histogram(bins = 15)
```



Also show the distribution of other variables.

```
state_df %>% select(-life_exp) %>%  
plot_num()
```



Part 2: Model Building Using Automatic Procedures

Backward Elimination

Fit a model using all the predictors.

```
fit_all = lm(life_exp ~ ., data = state_df)
summary(fit_all) %>%
  tidy %>%
  arrange(p.value)
```

```
## # A tibble: 8 x 5
##   term      estimate std.error statistic  p.value
##   <chr>      <dbl>     <dbl>     <dbl>    <dbl>
## 1 (Intercept) 7.09e+1 1.75      40.6    2.51e-35
## 2 murder     -3.01e-1 0.0466    -6.46   8.68e- 8
## 3 hs_grad      4.89e-2 0.0233     2.10   4.20e- 2
## 4 frost      -5.74e-3 0.00314   -1.82   7.52e- 2
## 5 population   5.18e-5 0.0000292 1.77    8.32e- 2
## 6 illiteracy   3.38e-2 0.366     0.0923 9.27e- 1
## 7 income      -2.18e-5 0.000244  -0.0892 9.29e- 1
## 8 area        -7.38e-8 0.00000167 -0.0443 9.65e- 1
```

Do the backward elimination. Based on the result above including all the predictors, **area** has the highest p-value $0.9649 > 0.05$, and is thus eliminated in the first step.

```
# no area
step1 = update(fit_all, . ~ . -area)
summary(step1) %>%
  tidy %>%
  arrange(p.value)
```

```
## # A tibble: 7 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>    <dbl>    <dbl>
## 1 (Intercept)  71.0      1.39     51.2    3.69e-40
## 2 murder      -0.302    0.0433    -6.96    1.45e- 8
## 3 hs_grad      0.0485    0.0207     2.35    2.37e- 2
## 4 frost       -0.00578   0.00297    -1.94    5.84e- 2
## 5 population    0.0000519 0.0000288    1.80    7.85e- 2
## 6 income       -0.0000244 0.000234    -0.104   9.17e- 1
## 7 illiteracy    0.0285    0.342     0.0833  9.34e- 1
```

Delete illiteracy with the highest p-value $0.9340 > 0.05$.

```
# no illiteracy
step2 = update(step1, . ~ . -illiteracy)
summary(step2) %>%
  tidy %>%
  arrange(p.value)
```

```
## # A tibble: 6 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>    <dbl>    <dbl>
## 1 (Intercept)  71.1      1.03     69.1    1.66e-46
## 2 murder      -0.300    0.0370    -8.10    2.91e-10
## 3 hs_grad      0.0478    0.0186     2.57    1.37e- 2
## 4 frost       -0.00591   0.00247    -2.39    2.10e- 2
## 5 population    0.0000511 0.0000271    1.89    6.57e- 2
## 6 income       -0.0000248 0.000232    -0.107   9.15e- 1
```

Delete income with the highest p-value $0.9153 > 0.05$.

```
# no income
step3 = update(step2, . ~ . -income)
summary(step3) %>%
  tidy %>%
  arrange(p.value)
```

```
## # A tibble: 5 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>    <dbl>    <dbl>
## 1 (Intercept)  71.0      0.953     74.5    8.61e-49
## 2 murder      -0.300    0.0366    -8.20    1.77e-10
## 3 hs_grad      0.0466    0.0148     3.14    2.97e- 3
## 4 frost       -0.00594   0.00242    -2.46    1.80e- 2
## 5 population    0.0000501 0.0000251    2.00    5.20e- 2
```

Delete population with the highest p-value $0.05201 > 0.05$.

```
# no population
step4 = update(step3, . ~ . -population)
summary(step4) %>% glance
```

```
## # A tibble: 1 x 6
##   r.squared adj.r.squared sigma statistic p.value    df
## *      <dbl>         <dbl> <dbl>      <dbl>    <dbl> <int>
## 1    0.713           0.694 0.743      38.0 1.63e-12     4
```

```
summary(step4) %>%
  tidy %>%
  arrange(p.value)
```

```
## # A tibble: 4 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>      <dbl>    <dbl>
## 1 (Intercept)  71.0      0.983      72.2 5.25e-49
## 2 murder      -0.283    0.0367     -7.71 8.04e-10
## 3 hs_grad      0.0499    0.0152      3.29 1.95e- 3
## 4 frost       -0.00691   0.00245     -2.82 6.99e- 3
```

Now each predictor remaining in the model has a p-value < 0.05 , so we stop here. The model we obtained by stepwise elimination is `life_exp ~ murder + hs_grad + frost`.

Forward Elimination

Fit simple linear regressions for all variables and look for the variable with the lowest p-value. Shown below is a summary of each variable fitted in the simple linear regression as the only predictor.

```
variables_list = names(state_df)

map(variables_list, ~lm(substitute(life_exp ~ i, list(i = as.name(.x))), data = state_df)) %>%
  map_df(., tidy) %>%
  filter(term != "(Intercept)") %>%
  arrange(p.value)
```

```
## # A tibble: 7 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>      <dbl>    <dbl>
## 1 murder      -0.284    0.0328     -8.66 2.26e-11
## 2 illiteracy -1.30     0.257     -5.04 6.97e- 6
## 3 hs_grad      0.0968    0.0195      4.96 9.20e- 6
## 4 income       0.000743 0.000297      2.51 1.56e- 2
## 5 frost        0.00677 0.00360      1.88 6.60e- 2
## 6 area        -0.00000169 0.00000226   -0.748 4.58e- 1
## 7 population -0.0000205 0.0000433   -0.473 6.39e- 1
```

So we first enter the one with the lowest p-value $2.26e-11 < 0.05$: murder.

```
forward1 = lm(life_exp ~ murder, data = state_df)
tidy(forward1)
```

```
## # A tibble: 2 x 5
##   term          estimate std.error statistic  p.value
##   <chr>          <dbl>    <dbl>      <dbl>    <dbl>
## 1 (Intercept)  73.0      0.270     270. 4.72e-78
## 2 murder      -0.284    0.0328     -8.66 2.26e-11
```

Add another one predictor to the simple linear regression model. Do this for all the rest predictors. Shown below is a summary of p-value for each variable added separately to the model as an additional predictor.


```
map(.x = variables_list, ~update(forward1, substitute(. ~ . + i, list(i = as.name(.x))))) %>%
  map_df(., tidy) %>%
  filter(term != "(Intercept)", term != "murder") %>%
  arrange(p.value)
```

```
## # A tibble: 6 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>   <dbl>
## 1 hs_grad      0.0439     0.0161      2.72 0.00909
## 2 population  0.0000683  0.0000274    2.49 0.0164
## 3 frost      -0.00578    0.00266    -2.17 0.0352
## 4 income      0.000370    0.000197    1.88 0.0666
## 5 area        0.00000118 0.00000146    0.806 0.424
## 6 illiteracy -0.172      0.281      -0.613 0.543
```

Enter the one with the lowest p-value 0.00909: hs_grad.

```
forward2 <- update(forward1, . ~ . + hs_grad)
tidy(forward2)
```

```
## # A tibble: 3 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>   <dbl>
## 1 (Intercept)  70.3      1.02     69.2 5.91e-49
## 2 murder      -0.237     0.0353    -6.72 2.18e- 8
## 3 hs_grad      0.0439     0.0161      2.72 9.09e- 3
```

Add another one predictor to the model. Do this for all the rest predictors. Shown below is a summary of p-value for each variable added separately to the model as an additional predictor.

```
map(.x = variables_list, ~update(forward2, substitute(. ~ . + i, list(i = as.name(.x))))) %>%
  map_df(., tidy) %>%
  filter(term != "(Intercept)", term != "murder", term != 'hs_grad') %>%
  arrange(p.value)
```

```
## # A tibble: 5 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>   <dbl>
## 1 frost      -0.00691    0.00245    -2.82 0.00699
## 2 population  0.0000625  0.0000259    2.41 0.0199
## 3 illiteracy  0.254      0.305      0.833 0.409
## 4 area      -0.00000106 0.00000162   -0.658 0.514
## 5 income      0.0000953  0.000239     0.398 0.692
```

Enter the one with the lowest p-value 0.00699: frost.

```
forward3 <- update(forward2, . ~ . + frost)
tidy(forward3)
```

```
## # A tibble: 4 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>   <dbl>
## 1 (Intercept)  71.0      0.983     72.2 5.25e-49
## 2 murder      -0.283     0.0367    -7.71 8.04e-10
## 3 hs_grad      0.0499     0.0152     3.29 1.95e- 3
## 4 frost      -0.00691    0.00245    -2.82 6.99e- 3
```

Add another one predictor to the model. Do this for all the rest predictors. Shown below is a summary of

p-value for each variable added separately to the model as an additional predictor.

```
map(.x = variables_list, ~update(forward3, substitute(. ~ . + i, list(i = as.name(.x))))) %>%
  map_df(., tidy) %>%
  filter(term != "(Intercept)", term != "murder", term != 'hs_grad', term != 'frost') %>%
  arrange(p.value)
```

```
## # A tibble: 4 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>    <dbl>
## 1 population  0.0000501  0.0000251     2.00  0.0520
## 2 income      0.000127  0.000223     0.571  0.571
## 3 illiteracy -0.182      0.328     -0.554  0.582
## 4 area       -0.000000329 0.00000154    -0.214  0.832
```

P-value of all new added variables are larger than 0.05, which means that they are not significant predictor, so we stop here.

```
forward_fit = lm(life_exp ~ murder + hs_grad + frost, data = state_df)
summary(forward_fit) %>% tidy()
```

```
## # A tibble: 4 x 5
##   term          estimate std.error statistic p.value
##   <chr>          <dbl>    <dbl>    <dbl>    <dbl>
## 1 (Intercept)  71.0      0.983     72.2  5.25e-49
## 2 murder      -0.283    0.0367    -7.71  8.04e-10
## 3 hs_grad       0.0499   0.0152     3.29  1.95e- 3
## 4 frost       -0.00691  0.00245    -2.82  6.99e- 3
```

```
summary(forward_fit) %>% glance()
```

```
## # A tibble: 1 x 6
##   r.squared adj.r.squared sigma statistic p.value    df
##   *      <dbl>         <dbl> <dbl>    <dbl>    <dbl> <int>
## 1    0.713      0.694 0.743     38.0  1.63e-12     4
```

The model we obtained by forward elimination is $\text{life_exp} \sim \text{murder} + \text{hs_grad} + \text{frost}$.

Stepwise Regreession

```
step(fit_all, direction = 'backward') %>%
  summary()
```

```
## Start:  AIC=-22.18
## life_exp ~ population + income + illiteracy + murder + hs_grad +
##   frost + area
##
##           Df Sum of Sq  RSS    AIC
## - area      1    0.0011 23.298 -24.182
## - income     1    0.0044 23.302 -24.175
## - illiteracy 1    0.0047 23.302 -24.174
## <none>                23.297 -22.185
## - population 1    1.7472 25.044 -20.569
## - frost      1    1.8466 25.144 -20.371
## - hs_grad    1    2.4413 25.738 -19.202
## - murder     1   23.1411 46.438  10.305
```

```

##
## Step: AIC=-24.18
## life_exp ~ population + income + illiteracy + murder + hs_grad +
##      frost
##
##           Df Sum of Sq    RSS    AIC
## - illiteracy 1      0.0038 23.302 -26.174
## - income     1      0.0059 23.304 -26.170
## <none>                        23.298 -24.182
## - population 1      1.7599 25.058 -22.541
## - frost      1      2.0488 25.347 -21.968
## - hs_grad    1      2.9804 26.279 -20.163
## - murder     1     26.2721 49.570  11.569
##
## Step: AIC=-26.17
## life_exp ~ population + income + murder + hs_grad + frost
##
##           Df Sum of Sq    RSS    AIC
## - income     1      0.006 23.308 -28.161
## <none>                        23.302 -26.174
## - population 1      1.887 25.189 -24.280
## - frost      1      3.037 26.339 -22.048
## - hs_grad    1      3.495 26.797 -21.187
## - murder     1     34.739 58.041  17.456
##
## Step: AIC=-28.16
## life_exp ~ population + murder + hs_grad + frost
##
##           Df Sum of Sq    RSS    AIC
## <none>                        23.308 -28.161
## - population 1      2.064 25.372 -25.920
## - frost      1      3.122 26.430 -23.877
## - hs_grad    1      5.112 28.420 -20.246
## - murder     1     34.816 58.124  15.528
##
## Call:
## lm(formula = life_exp ~ population + murder + hs_grad + frost,
##     data = state_df)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -1.47095 -0.53464 -0.03701  0.57621  1.50683
##
## Coefficients:
##              Estimate Std. Error t value Pr(>|t|)
## (Intercept)  7.103e+01  9.529e-01  74.542 < 2e-16 ***
## population    5.014e-05  2.512e-05   1.996  0.05201 .
## murder       -3.001e-01  3.661e-02  -8.199 1.77e-10 ***
## hs_grad       4.658e-02  1.483e-02   3.142  0.00297 **
## frost        -5.943e-03  2.421e-03  -2.455  0.01802 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##

```

```
## Residual standard error: 0.7197 on 45 degrees of freedom
## Multiple R-squared:  0.736, Adjusted R-squared:  0.7126
## F-statistic: 31.37 on 4 and 45 DF,  p-value: 1.696e-12
```

The model we obtained by stepwise regression is `life_exp ~ population + murder + hs_grad + frost`.

Questions

a) Do the procedures generate the same model?

Forward elimination and backward elimination generated the same model: `life_exp ~ murder + hs_grad + frost`. However, stepwise regression generated a larger model with one more predictor, `population`.

b) Is there any variable a close call? What was your decision: keep or discard? Provide arguments for your choice. (Note: this question might have more or less relevance depending on the ‘subset’ you choose).

The variable `population` is a close call, with $p\text{-value} = 0.052 \sim 0.05$. I would keep it, because its $p\text{-value}$ is quite close to 0.05. This model has a better AIC than a smaller model. Also, adding ‘`population`’ contributes to the goodness of fit by increasing the adjusted R^2 from 0.6939 to 0.7126.

c) Is there any association between ‘Illiteracy’ and ‘HS graduation rate’? Does your ‘subset’ contain both?

The Pearson correlation coefficient between ‘Illiteracy’ and ‘HS graduation rate’ is -0.66, indicating a moderate association. My subset only contains one of them. ‘Illiteracy’ is not included.

Part 3: Criterion-based Procedures

```
# Leaps function provides all-subsets analysis
```

```
# Printing the 2 best models of each size, using the Cp criterion:
```

```
leaps(x = state_df %>% select(-life_exp), y = state_df[[1]], nbest = 1, method = "Cp")
```

```
## $which
```

```
##      1      2      3      4      5      6      7
## 1 FALSE FALSE FALSE TRUE FALSE FALSE FALSE
## 2 FALSE FALSE FALSE TRUE  TRUE FALSE FALSE
## 3 FALSE FALSE FALSE TRUE  TRUE  TRUE FALSE
## 4  TRUE FALSE FALSE TRUE  TRUE  TRUE FALSE
## 5  TRUE  TRUE FALSE TRUE  TRUE  TRUE FALSE
## 6  TRUE  TRUE  TRUE TRUE  TRUE  TRUE FALSE
## 7  TRUE  TRUE  TRUE TRUE  TRUE  TRUE  TRUE
##
```

```
## $label
```

```
## [1] "(Intercept)" "1"          "2"          "3"          "4"
## [6] "5"              "6"          "7"
```

```
##
```

```
## $size
```

```
## [1] 2 3 4 5 6 7 8
```

```
##
```

```
## $Cp
```

```
## [1] 16.126760  9.669894  3.739878  2.019659  4.008737  6.001959  8.000000
```

```
# Printing the 2 best models of each size, using the adjusted R^2 criterion:
```

```
leaps(x = state_df %>% select(-life_exp), y = state_df[[1]], nbest = 1, method = "adjr2")
```

```

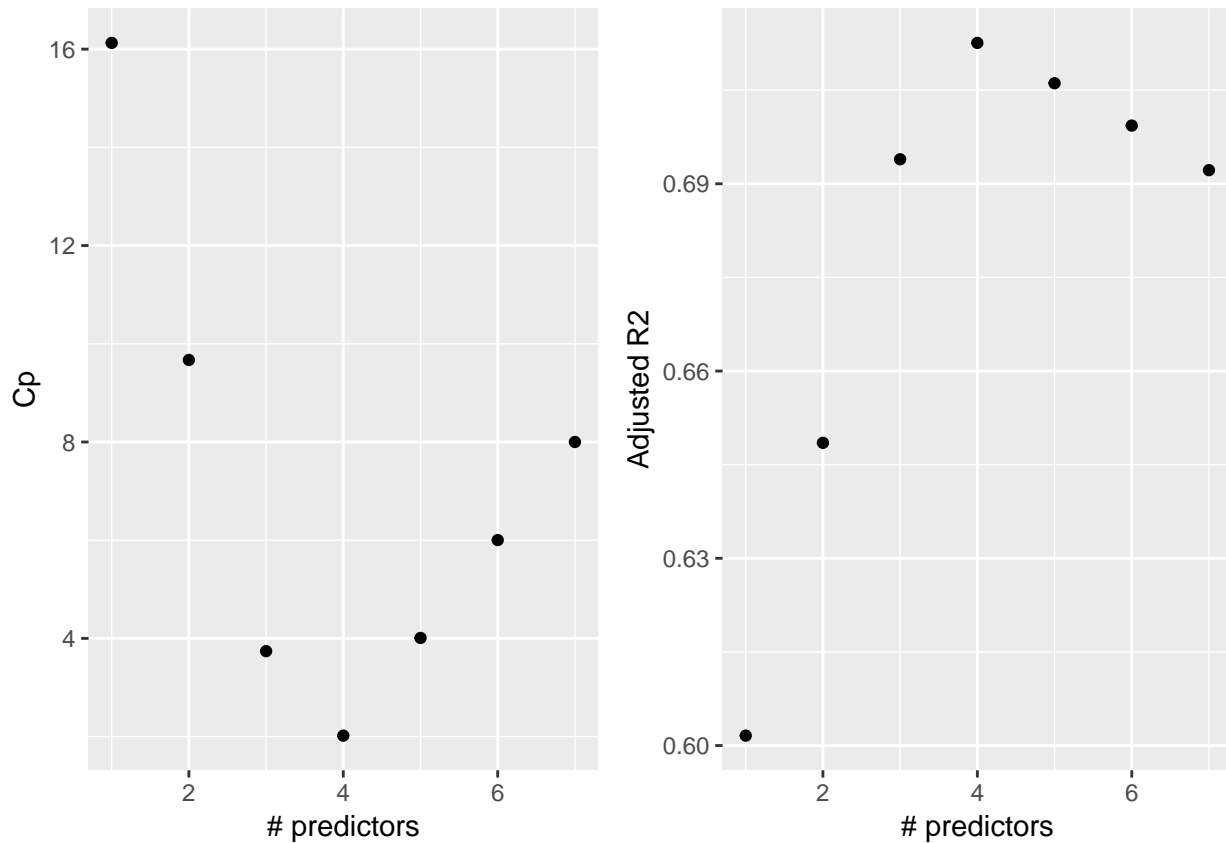
## $which
##      1      2      3      4      5      6      7
## 1 FALSE FALSE FALSE TRUE FALSE FALSE FALSE
## 2 FALSE FALSE FALSE TRUE  TRUE FALSE FALSE
## 3 FALSE FALSE FALSE TRUE  TRUE  TRUE FALSE
## 4  TRUE FALSE FALSE TRUE  TRUE  TRUE FALSE
## 5  TRUE  TRUE FALSE TRUE  TRUE  TRUE FALSE
## 6  TRUE  TRUE  TRUE TRUE  TRUE  TRUE FALSE
## 7  TRUE  TRUE  TRUE TRUE  TRUE  TRUE  TRUE
##
## $label
## [1] "(Intercept)" "1"          "2"          "3"          "4"
## [6] "5"              "6"              "7"
##
## $size
## [1] 2 3 4 5 6 7 8
##
## $adjr2
## [1] 0.6015893 0.6484991 0.6939230 0.7125690 0.7061129 0.6993268 0.6921823

# Summary of models for each size (one model per size)
b = regsubsets(life_exp ~ ., data = state_df)
rs <- summary(b)

# Plots of Cp and Adj-R2 as functions of parameters
plot_cp =
  tibble(x = 1:7, y = rs$cp) %>%
  ggplot(aes(x = x, y = y)) +
    geom_point() +
    labs(x = "# predictors", y = "Cp")
plot_adjr2 =
  tibble(x = 1:7, y = rs$adjr2) %>%
  ggplot(aes(x = x, y = y)) +
    geom_point() +
    labs(x = "# predictors", y = "Adjusted R2")

plot_cp + plot_adjr2

```



rs

```
## Subset selection object
## Call: regsubsets.formula(life_exp ~ ., data = state_df)
## 7 Variables (and intercept)
##           Forced in Forced out
## population      FALSE      FALSE
## income          FALSE      FALSE
## illiteracy       FALSE      FALSE
## murder          FALSE      FALSE
## hs_grad         FALSE      FALSE
## frost           FALSE      FALSE
## area            FALSE      FALSE
## 1 subsets of each size up to 7
## Selection Algorithm: exhaustive
##           population income illiteracy murder hs_grad frost area
## 1  ( 1 ) " "          " "          " "      "*"    " "    " "
## 2  ( 1 ) " "          " "          " "      "*"    "*"    " "
## 3  ( 1 ) " "          " "          " "      "*"    "*"    "*"
## 4  ( 1 ) "*"          " "          " "      "*"    "*"    "*"
## 5  ( 1 ) "*"          "*"          " "      "*"    "*"    "*"
## 6  ( 1 ) "*"          "*"          "*"      "*"    "*"    "*"
## 7  ( 1 ) "*"          "*"          "*"      "*"    "*"    "*"

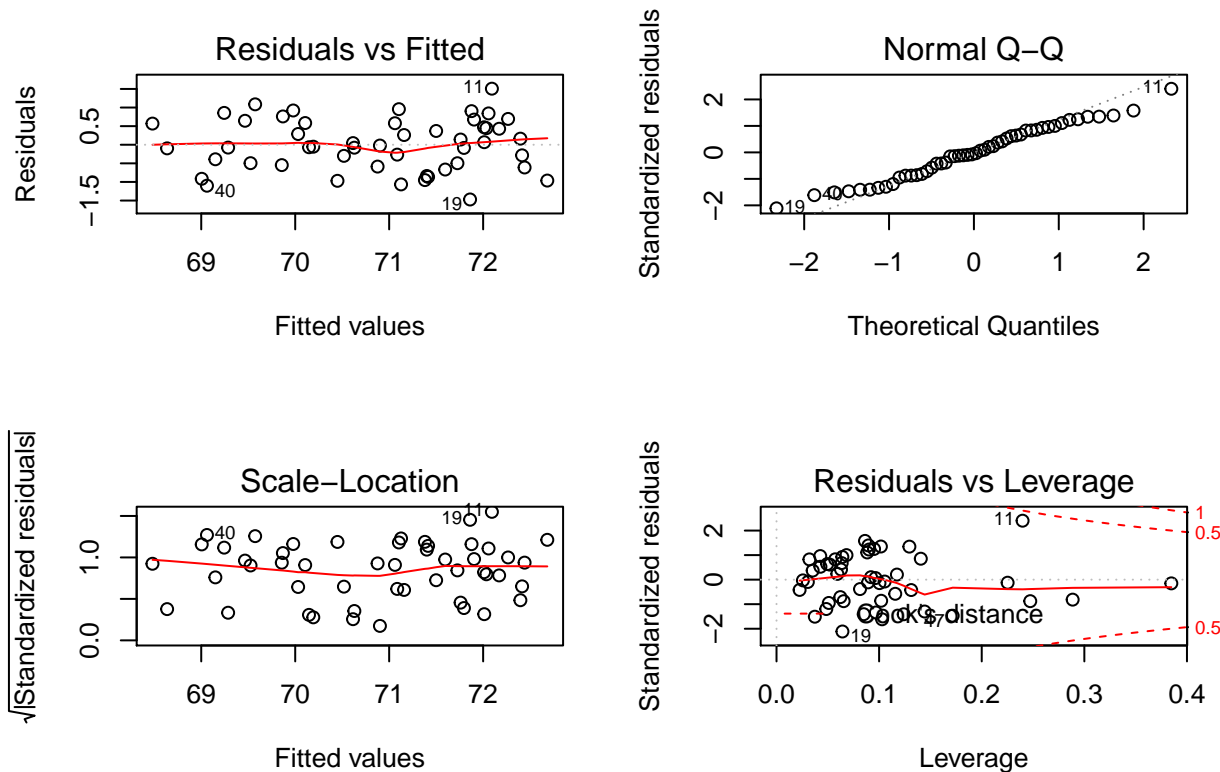
```

Based on the Cp and adjusted R2 criterion, I would choose the 4-predictor model. The best 4-predictor model is $\text{life_exp} \sim \text{population} + \text{murder} + \text{hs_grad} + \text{frost}$. It has the highest adjusted R2 and the lowest Cp value.

Part 4: Diagnostics

The models chosen from Part 2 and 3 are the same: `life_exp ~ population + murder + hs_grad + frost`, so we will use this one as our final model.

```
final_fit = lm(life_exp ~ population + murder + hs_grad + frost, data = state_df)
par(mfrow = c(2,2))
plot(final_fit)
```



```
influence.measures(final_fit)
```

```
## Influence measures of
## lm(formula = life_exp ~ population + murder + hs_grad + frost, data = state_df) :
##
##      dfb.1_ dfb.pplt dfb.mrdr dfb.hs_g dfb.frst dffit cov.r
## 1  0.035492 -1.31e-01 1.81e-01 -0.040552 -0.093627 0.34385 1.200
## 2  0.410334 1.83e-01 -4.02e-01 -0.349223 -0.164049 -0.50117 1.364
## 3  0.015219 2.09e-01 5.00e-02 -0.236749 0.457644 -0.53734 1.081
## 4  0.329890 -1.72e-01 2.46e-03 -0.317837 -0.087569 0.49350 0.921
## 5  0.035541 -9.14e-02 4.05e-03 -0.044178 0.026988 -0.11867 1.814
## 6 -0.316742 -5.38e-02 2.18e-01 0.272452 0.219005 0.44171 0.986
## 7  0.051252 1.77e-02 -9.17e-02 -0.024669 0.010955 0.14381 1.125
## 8 -0.047471 1.85e-01 3.13e-02 -0.036008 0.090584 -0.30196 0.896
## 9 -0.000547 2.67e-03 9.55e-04 0.006028 -0.014997 0.02107 1.238
## 10 -0.008460 8.05e-03 -2.53e-02 0.016126 -0.003117 -0.04691 1.241
## 11 -0.033009 -5.69e-01 -2.76e-01 0.664444 -1.244026 1.42824 0.741
## 12 -0.025510 -5.09e-02 7.21e-05 0.049597 -0.006575 0.11025 1.134
## 13 0.007812 -1.81e-02 -9.37e-03 -0.000558 -0.014652 -0.02615 1.250
## 14 -0.000389 -1.61e-03 7.15e-05 0.000713 -0.001883 -0.00488 1.148
## 15 0.012296 5.30e-03 -3.71e-02 0.002013 -0.001564 0.05819 1.182
```

```

## 16 -0.017022 -4.31e-02 -6.33e-02 0.084668 -0.057912 0.20256 1.054
## 17 0.235268 -5.35e-02 5.72e-02 -0.303436 0.095100 0.40778 1.036
## 18 -0.057605 7.39e-02 -5.88e-02 0.036101 0.101646 -0.20714 1.219
## 19 -0.221919 4.84e-02 2.91e-01 0.174087 -0.166830 -0.57681 0.709
## 20 0.002387 7.62e-03 -1.99e-02 -0.003218 -0.004869 -0.06309 1.122
## 21 -0.052828 -8.12e-02 1.63e-01 -0.023468 0.081194 -0.23178 1.098
## 22 -0.146492 1.63e-01 2.01e-01 0.041939 0.203775 0.34590 1.069
## 23 0.061153 7.86e-02 -1.55e-01 -0.036661 0.075346 0.27039 1.073
## 24 -0.152293 1.93e-01 -1.58e-01 0.159679 0.083500 -0.44227 1.011
## 25 0.020092 6.51e-03 5.33e-02 -0.046052 0.055511 0.14957 1.071
## 26 0.071108 9.14e-02 -2.42e-02 -0.080966 -0.098656 -0.27162 1.000
## 27 0.016596 -2.02e-02 -7.07e-02 0.020076 -0.005414 0.14244 1.132
## 28 0.402992 1.43e-01 -4.24e-01 -0.288096 -0.283276 -0.51725 1.460
## 29 -0.009464 2.56e-02 4.10e-02 0.008083 -0.085716 -0.18242 1.127
## 30 -0.088716 -1.38e-01 1.24e-01 0.071170 -0.017018 -0.21995 1.065
## 31 -0.049072 -5.53e-02 7.62e-02 0.039222 0.028848 0.10637 1.171
## 32 0.011393 -6.01e-02 -4.89e-03 -0.002487 -0.012640 -0.06681 1.442
## 33 -0.019970 -1.75e-03 -3.80e-03 0.025276 -0.005219 -0.03437 1.227
## 34 0.291080 9.70e-03 -2.82e-01 -0.297496 0.209852 0.52382 1.047
## 35 0.002238 -9.51e-02 1.05e-02 0.013510 -0.044242 -0.11347 1.198
## 36 0.035812 -1.99e-02 -2.89e-02 -0.011934 -0.037269 0.07001 1.142
## 37 -0.023790 3.95e-02 7.55e-02 -0.054191 0.135471 -0.16331 1.262
## 38 -0.139407 -4.61e-01 1.88e-01 0.193251 -0.169119 -0.53682 1.019
## 39 0.062823 -5.58e-03 -5.47e-02 -0.051111 -0.004418 0.07465 1.262
## 40 -0.317583 1.58e-01 -9.62e-02 0.357681 0.020151 -0.55733 0.925
## 41 0.015278 -1.23e-03 -1.79e-02 -0.013387 0.009648 0.03141 1.231
## 42 0.120516 -4.05e-02 4.94e-02 -0.136805 -0.009497 0.24190 1.086
## 43 0.018620 2.71e-01 6.79e-02 -0.025291 -0.096144 0.45875 1.013
## 44 -0.222055 -1.08e-01 4.13e-02 0.298455 -0.017639 0.38846 1.037
## 45 -0.040916 -6.45e-02 4.00e-02 0.022311 0.117523 0.20328 1.099
## 46 -0.005268 2.62e-05 -3.35e-03 0.005718 0.000217 -0.01658 1.153
## 47 0.006164 9.92e-02 2.62e-01 -0.320275 0.559123 -0.67816 1.056
## 48 -0.374426 7.95e-02 1.66e-01 0.354831 0.010543 -0.43422 0.976
## 49 0.069886 6.29e-02 -1.10e-01 -0.057052 0.044388 0.17475 1.136
## 50 0.186254 9.61e-02 -1.55e-01 -0.154629 -0.146677 -0.28910 1.147
##      cook.d      hat inf
## 1 2.38e-02 0.1406
## 2 5.05e-02 0.2473 *
## 3 5.68e-02 0.1443
## 4 4.71e-02 0.0862
## 5 2.88e-03 0.3848 *
## 6 3.82e-02 0.0896
## 7 4.19e-03 0.0494
## 8 1.77e-02 0.0374
## 9 9.08e-05 0.0965
## 10 4.50e-04 0.1003
## 11 3.64e-01 0.2398 *
## 12 2.47e-03 0.0428
## 13 1.40e-04 0.1054
## 14 4.88e-06 0.0257
## 15 6.92e-04 0.0593
## 16 8.22e-03 0.0426
## 17 3.28e-02 0.0951
## 18 8.71e-03 0.1157

```



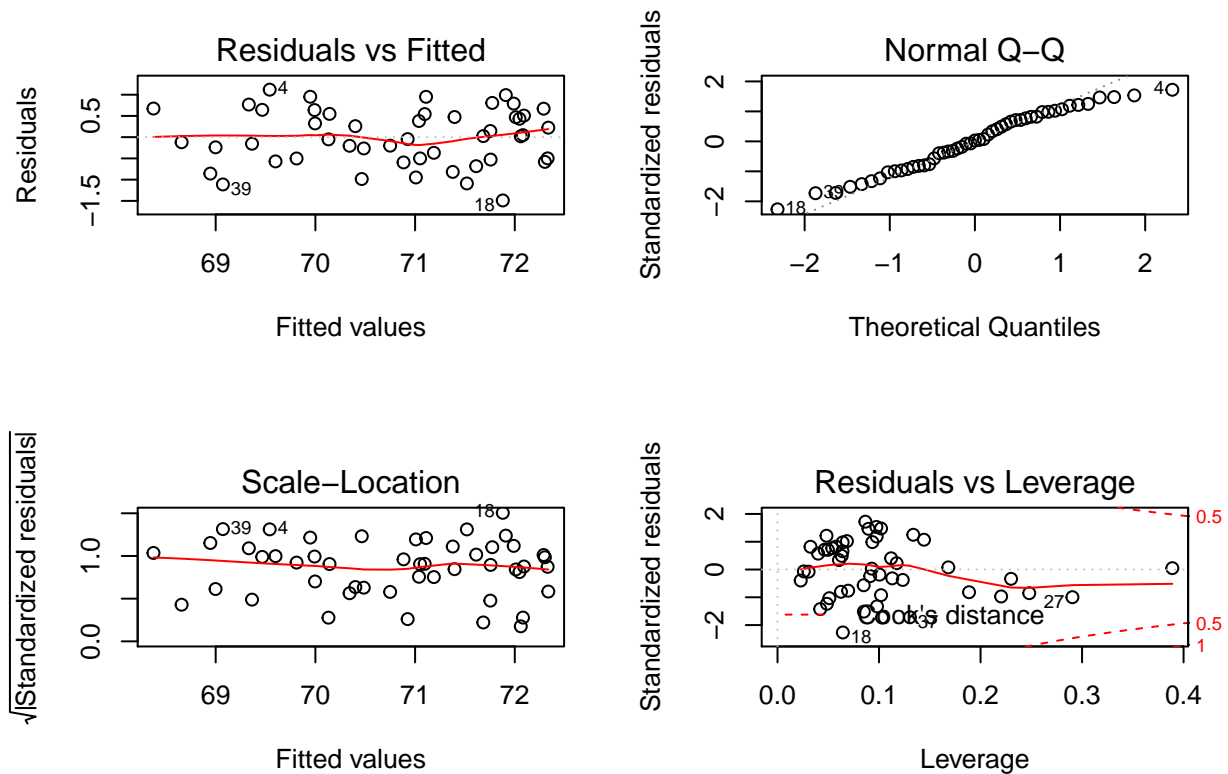
```
## 19 6.13e-02 0.0642
## 20 8.11e-04 0.0225
## 21 1.08e-02 0.0654
## 22 2.38e-02 0.0884
## 23 1.46e-02 0.0682
## 24 3.84e-02 0.0969
## 25 4.51e-03 0.0321
## 26 1.46e-02 0.0485
## 27 4.12e-03 0.0519
## 28 5.39e-02 0.2886      *
## 29 6.73e-03 0.0622
## 30 9.70e-03 0.0510
## 31 2.31e-03 0.0629
## 32 9.13e-04 0.2252      *
## 33 2.42e-04 0.0893
## 34 5.39e-02 0.1295
## 35 2.62e-03 0.0814
## 36 1.00e-03 0.0353
## 37 5.43e-03 0.1313
## 38 5.63e-02 0.1240
## 39 1.14e-03 0.1174
## 40 5.99e-02 0.1029
## 41 2.02e-04 0.0921
## 42 1.17e-02 0.0642
## 43 4.13e-02 0.1017
## 44 2.98e-02 0.0901
## 45 8.32e-03 0.0572
## 46 5.62e-05 0.0305
## 47 8.96e-02 0.1717
## 48 3.69e-02 0.0850
## 49 6.18e-03 0.0636
## 50 1.68e-02 0.1020
```

a) Identify any leverage and/or influential points and take appropriate measures.

Based on Cook's distance, Observation 11 is a potentially influential observation. Remove this point and fit the model again.

```
state_df_no_11 =
  state_df %>%
  mutate(index = 1:nrow(state_df)) %>%
  filter(!index %in% c(11))

final_fit_no_11 = lm(life_exp ~ population + murder + hs_grad + frost, data = state_df_no_11)
par(mfrow=c(2,2))
plot(final_fit_no_11)
```



b) Check the model assumptions.

The residuals are spread equally along the fitted values. We can assume that the residuals have mean 0 and equal variance, and are independent of each other. The QQ plot shows a minor deviate from normal distribution at the tail, which may be caused by outliers.

Model Validation

Using the 'final' model chosen in part 4, focus on MSE to test the model predictive ability.

a) Use a 10-fold cross-validation (10 repeats).

```
cv_df =
  crossv_kfold(state_df, 10) %>%
  mutate(model = map(train, ~lm(life_exp ~ population + murder + hs_grad + frost, data = .x))) %>%
  mutate(mse = map2_dbl(model, test, ~mse(model = .x, data = .y)))
```

Below shows the 10 mse's of the 10-fold cross-validation.

```
cv_df$mse

##          1          2          3          4          5          6          7
## 0.4741850 0.5431650 0.7532672 0.5256390 0.4621004 0.5299299 0.3237438
##          8          9         10
## 0.6292451 1.1462354 0.2424295

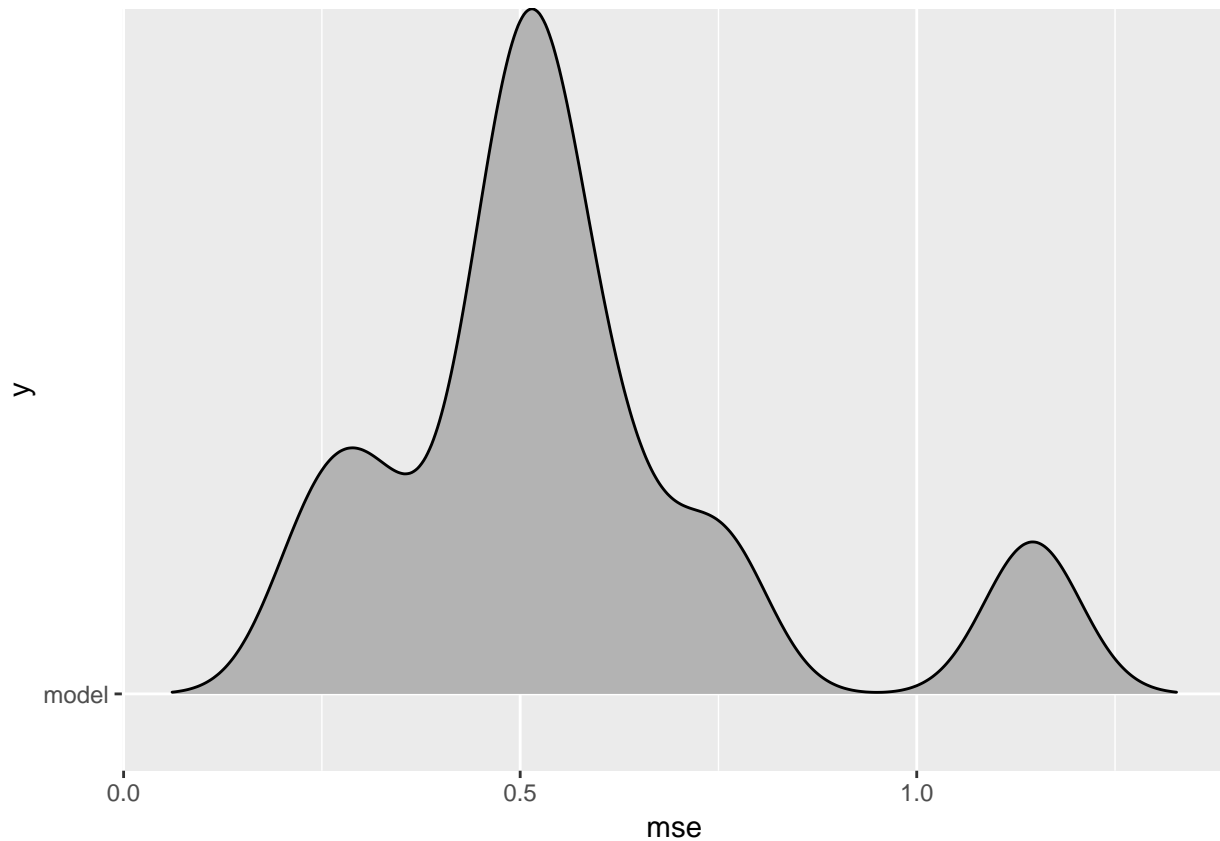
summary_continuous(cv_df$mse)

## # A tibble: 1 x 6
##   mean    sd median maximum minimum  IQR
##   <dbl> <dbl> <dbl>   <dbl>   <dbl> <dbl>
```

```
## 1 0.563 0.250 0.528 1.15 0.242 0.143
```

```
cv_df %>%
  ggplot(aes(x = mse, y = "model")) +
  geom_density_ridges()
```

```
## Picking joint bandwidth of 0.0604
```



This is the summary of the 10-fold cross-validation. `summary_continuous` function was written in Part 1.

b) Bootstrapping

```
fit_final = lm(life_exp ~ population + murder + hs_grad + frost, data = state_df)

bootstrap_df =
  state_df %>%
  add_predictions(fit_final) %>%
  mutate(resid = life_exp - pred)

bootstrap = function(x){
  bootstrap_df_new =
    bootstrap_df %>%
    mutate(y_new = pred + sample(bootstrap_df$resid, replace = TRUE))
  model = lm(y_new ~ population + murder + hs_grad + frost, data = bootstrap_df_new)
  mse(model, bootstrap_df_new)
}

result_df_10 =
  map(1:10, bootstrap) %>%
  c() %>% as.numeric %>% as.tibble()
```

```

result_df_1000 =
  map(1:1000, bootstrap) %>%
  c() %>% as.numeric %>% as.tibble()

summary_continuous(result_df_10$value)

## # A tibble: 1 x 6
##   mean      sd median maximum minimum   IQR
##   <dbl> <dbl> <dbl>   <dbl>   <dbl> <dbl>
## 1 0.448 0.0669  0.453    0.558    0.320 0.0785

summary_continuous(result_df_1000$value)

```

```

## # A tibble: 1 x 6
##   mean      sd median maximum minimum   IQR
##   <dbl> <dbl> <dbl>   <dbl>   <dbl> <dbl>
## 1 0.419 0.0690  0.417    0.631    0.219 0.0908

```

Here shows the discriptive statistics of the 10X and 1000X bootstrap.

```

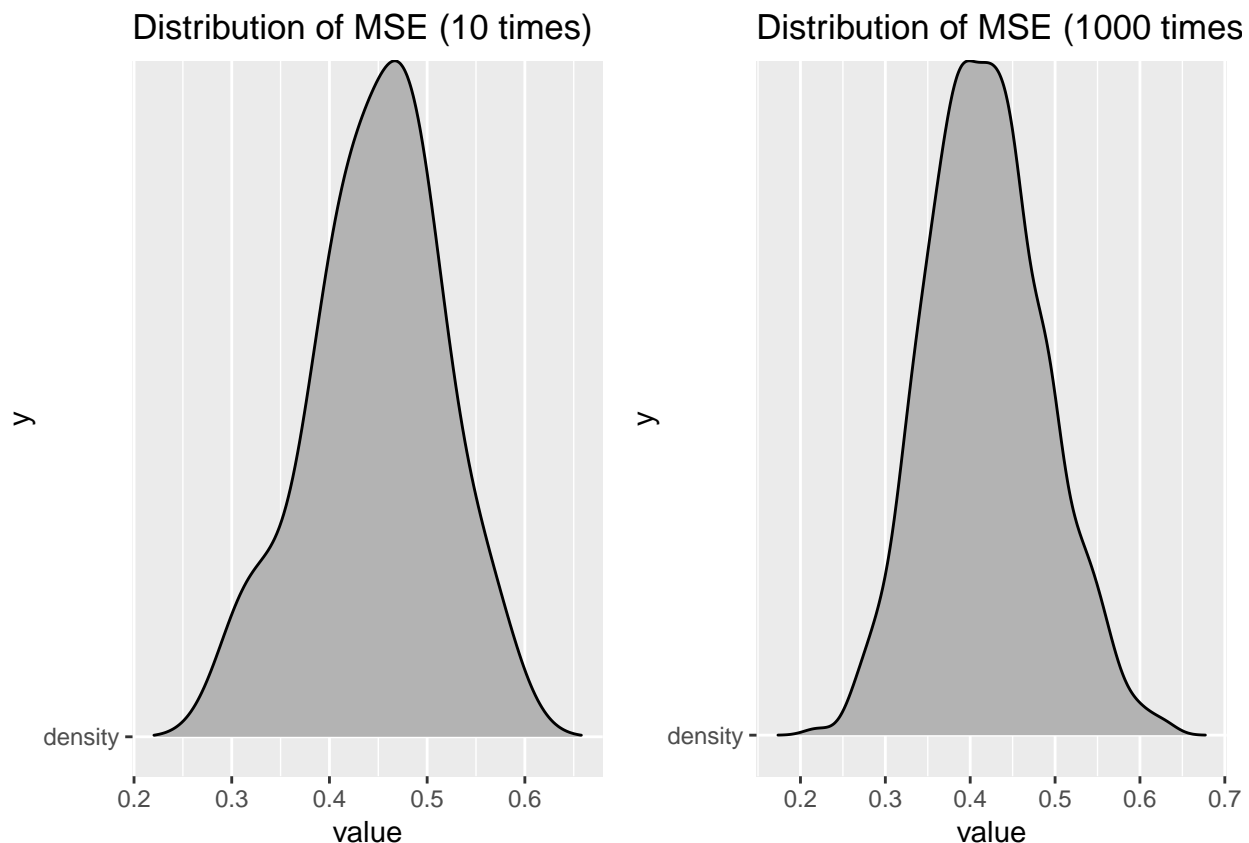
plot_10x =
result_df_10 %>%
  ggplot(aes(y = "density", x = value)) +
    geom_density_ridges() +
    labs(title = "Distribution of MSE (10 times)")
plot_1000x =
result_df_1000 %>%
  ggplot(aes(y="density", x = value)) +
    geom_density_ridges() +
    labs(title = "Distribution of MSE (1000 times)")

plot_10x + plot_1000x

```

```
## Picking joint bandwidth of 0.0333
```

```
## Picking joint bandwidth of 0.0153
```



The plots show the distribution of MSEs from the 10-time and 1000-time bootstrapping.

c) Summary

```
bind_rows(
  summary_continuous(cv_df$mse),
  summary_continuous(result_df_10$value),
  summary_continuous(result_df_1000$value)
) %>%
mutate(method = c('CV-10fold', 'bootstrap-10X', 'bootstrap-1000X')) %>%
select(method, everything()) %>%
knitr::kable(digits = 2)
```

method	mean	sd	median	maximum	minimum	IQR
CV-10fold	0.56	0.25	0.53	1.15	0.24	0.14
bootstrap-10X	0.45	0.07	0.45	0.56	0.32	0.08
bootstrap-1000X	0.42	0.07	0.42	0.63	0.22	0.09

Although the medians of the two methods are similar (0.41 vs 0.42), their means and standard deviations are quite different. For bootstrapping, The means and SD's of MSE are similar in the 10 trials and 1000 trials, showing reliability of this method even with small amount of trials. On the other hand, the standard deviation for the Cross Validation method is significantly higher, which indicates a low stability of this method, especially when k (# of folds) is small. Therefore, I would recommend *bootstrapping* for assessing model performance.