Artificial Neural Network Application for Reservoir Characterization

Chukwuemeka Okoli (https://www.linkedin.com/in/chukwuemeka-okoli-38686923/)

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Introduction

As a Data Scientist in the Oil and Gas Industry, we are often faced with numerous challenges. For instance, with available Gas production data for an unconventional reservoir, can we establish a relationship between several independent variable in the production process and the dependent variable "Gas production per months" using Artificial Neural Network. In this R Markdown (https://rmarkdown.rstudio.com) project, an illustration of how we can apply Artificial Neural Network to Gas production is discussed. The dataset for this project is the "Q4_data.csv" file.

The Goal

Find a relationship between the dependent variable "Gas production per months" and several independent variables. Essentially, we want to determine the Gas Production per month for the given well data based on different factors.

Preliminaries

First, the required packages are installed using the install.packages() function.

```
# install knitr
install.packages("knitr")

# install the rmarkdown package
install.packages("rmarkdown")

# install ggplot for plotting
install.packages("ggplot2")

# install dplyr for data manipulation
install.packages("dplyr")

# install neuralnet for artificial neural network
install.packages("neuralnet")
install.packages("NeuralNetTools")
```

The installed package is then loaded using the library() function.

```
# load libraries
library(knitr)
library(rmarkdown)
library(ggplot2)
library(dplyr)
library(neuralnet)
library(NeuralNetTools)
library(xfun)
```

Data Import

We can call in the dataset into R using the code below. The data is stored in a .csv file.

```
set.seed(440)
data <- read.csv("Q4_data.csv", header=TRUE)</pre>
```

Summaries

The data object field_data is complex. It contains various information about properties of the well such as the perforation interval, fracture volume, proppant, casing depth, etc. Use summary() to get a quick summary of the data.

Summary(data)

Perf_Int epth	Frac_vol	Proppant	N.frac	Tubing.depth	Casing.
Min. : 12.0 7325	Min. : 7478	Min. : 43500	Min. : 1.000	Min. : 4952	Min. :
1st Qu.: 381.8 8176	1st Qu.: 29524	1st Qu.: 164625	1st Qu.: 1.000	1st Qu.: 7652	1st Qu.:
	Median : 36200	Median : 296946	Median : 2.000	Median : 8002	Median :
Mean : 708.1	Mean : 56848	Mean : 662449	Mean : 2.309	Mean : 8020	Mean :
-	3rd Qu.: 56876	3rd Qu.:1135250	3rd Qu.: 3.000	3rd Qu.: 8407	3rd Qu.:
	Max. :3637776	Max. :4437897	Max. :11.000	Max. :12100	Max. :
	Choke.Size	SITHP	SG gas \	Well Type	Latitude
Min. : 0.0	Min. : 6.00	Min. : 300 Mi	n. :0.5810 Mi		
1st Qu.: 450.0	1st Qu.: 14.00	1st Qu.:1615 1s	t Qu.:0.7000 1st	t Qu.:2.000 1s	t Qu.:
Median : 814.1 33	Median : 20.00	Median :2278 Me	dian :0.7000 Med	dian :3.000 Med	dian :
Mean : 905.6	Mean : 28.18	Mean :2250 Me	an :0.6922 Mea	an :2.504 Mea	an : 7
3rd Qu.:1265.0	3rd Qu.: 32.00	3rd Qu.:2913 3r	d Qu.:0.7000 3rd	d Qu.:3.000 3rd	d Qu.:
Max. :2964.0	Max. :128.00	Max. :3715 Ma	x. :1.0320 Max	x. :3.000 Max	x. :3436
Long.	Gas_prod	Acid			
Min. :-97.40	Min. : 95	Min. :1.000			
1st Qu.:-97.35	1st Qu.: 3840	1st Qu.:1.000			
Median :-97.31	Median : 6392	Median :1.000			
	Mean : 13281	Mean :1.193			
Mean •-97 00		11Can			
Mean :-97.00 3rd Qu.:-97.26		3rd Qu.:1.000			

The set.seed() function is useful when running simulations to ensure all results, figures, etc. are reproducible. We can then check that no data point is missing. If we have a missing data point, we need to fix the dataset.

Check that no data is missing apply(data,2,function(x) sum(is.na(x)))

Perf_Int ke.Size	Frac_vol	Proppant	N.frac Tub	oing.depth Ca	asing.depth	FTP	Cho
0	0	0	0	0	0	0	
SITHP 0	SG_gas 0	Well_Type 0	Latitude 0	Long. 0	Gas_prod 0	Acid 0	

We can see that there is no missing data. In the case where missing data exist, data cleaning will have to take place.

Data Analysis

To analyze the data, we first proceed by randomly splitting the data into a *train* and a *test* dataset. A linear regression model is fit to the train dataset, and tested on the test dataset.

```
# Train-test random splitting for linear model
index <- sample(1:nrow(data), round(0.75*nrow(data)))
train <- data[index,]
test <- data[-index,]

# Fitting linear model to the train dataset
lm.fit <- lm(Gas_prod~., data=train)
summary(lm.fit)</pre>
```

```
Call:
lm(formula = Gas_prod ~ ., data = train)
Residuals:
   Min
            1Q Median
                            3Q
                                  Max
                 -1262
 -47265
       -10594
                          6029 1130490
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)
             4.242e+06 6.496e+06
                                   0.653 0.51423
Perf Int
             7.347e+00 9.449e+00
                                   0.778 0.43739
Frac vol
            -2.151e-03 1.828e-02 -0.118 0.90643
Proppant
            -2.850e-03 8.633e-03 -0.330 0.74148
N.frac
             3.468e+03 3.381e+03
                                  1.026 0.30573
Tubing.depth 2.724e+00 7.987e+00
                                   0.341 0.73332
Casing.depth -4.627e+00 7.114e+00 -0.650 0.51589
FTP
                                   1.766 0.07839
             1.334e+01 7.556e+00
Choke.Size -1.336e+02 1.828e+02 -0.731 0.46519
SITHP
            -1.010e+01 5.776e+00 -1.749 0.08126 .
                                  0.244 0.80744
SG gas
            2.025e+04 8.302e+04
            -1.712e+04 5.909e+03 -2.897 0.00402 **
Well_Type
Latitude
            -1.619e+00 2.527e+00 -0.641 0.52226
Long.
            4.283e+04 6.657e+04
                                   0.643 0.52049
Acid
            -7.482e+03 9.953e+03 -0.752 0.45275
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 64930 on 319 degrees of freedom
Multiple R-squared: 0.05826,
                             Adjusted R-squared:
F-statistic: 1.41 on 14 and 319 DF, p-value: 0.1465
```

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```
# Predicted data from linear model fit
pr.lm <- predict(lm.fit, test)

# Test MSE
MSE.lm <- sum((pr.lm - test$Gas_prod)^2)/nrow(test)</pre>
```

Before fitting a neural network, some preparation needs to be done. Neural networks are not that easy to train and tune. As a first step, we need to address data preprocessing. It is good practice to normalize your data before training a neural network. This step is important because, depending on your dataset, avoiding normalization may lead to useless results or to a very difficult training process (most of the time, the algorithm will not converge before the number of maximum iterations allowed). You can choose different methods to scale the data (z-normalization, min-max scale, etc.). The data were scaled using the min-max method and scaled in the interval [0,1]. We therefore scale and split the data before moving on.

```
# Neural network fitting

# Scaling data for the Neural Network
maxs <- apply(data, 2, max)
mins <- apply(data, 2, min)
scaled <- as.data.frame(scale(data, center = mins, scale = maxs - mins))

# Train-test split
train_ <- scaled[index,]
test_ <- scaled[-index,]</pre>
```

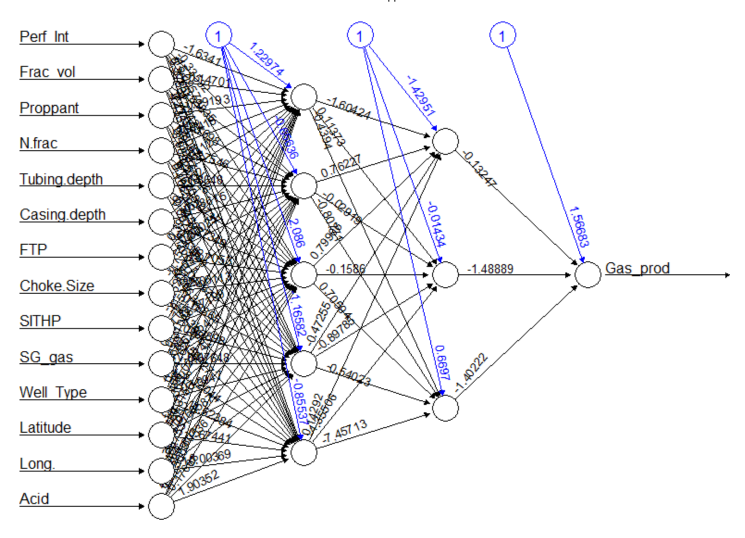
Since there is no fixed rule as to how many layers and neurons to use, we are going to use two (2) hidden layers with 5 and 3 neurons. To fit the network, we use the following code:

```
# Training the neural network
n <- names(train_)
f <- as.formula(paste("Gas_prod ~", paste(n[!n %in% "Gas_prod"], collapse = " + ")))
neural_net <- neuralnet(f, data = train_, hidden = c(5,3), linear.output = T)</pre>
```

Note that the hidden argument accepts a vector with the number of neurons for each hidden layer, while the argument linear.output is used to specify whether we want to do regression **linear.output = TRUE** or **classification linear.output = FALSE**

The neuralnet package provides a nice tool to plot the model. Use the following code to plot the neural network in R:

```
# Visual plot of the model plot(neural_net)
```



Now we can try to predict the values for the test set and calculate the mean squared error (MSE). Remember that the net will output a normalized prediction, so we need to scale it back in order to make a meaningful comparison (or just a simple prediction). The mean squared error is one metric used to measure prediction accuracy. The MSE is calculated as:

$$MSE = rac{1}{n}\sum_{i=1}^n (Y_i - \hat{Y_i}^2)$$

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```
# Prediction
pr.nn <- compute(neural_net, test_[,1:14])

# Results from Neural Networks are normalized (scaled)
# Descaling for comparison
pr.nn_ <- pr.nn$net.result*(max(data$Gas_prod)-min(data$Gas_prod))+min(data$Gas_prod)
test.r <- (test_$Gas_prod)*(max(data$Gas_prod)-min(data$Gas_prod))+min(data$Gas_prod)
# Calculating MSE
MSE.nn <- sum((test.r - pr.nn_)^2)/nrow(test_)</pre>
```

We then compare the two mean squares error obtained from the linear regression and the neural network fitting process using the code:

```
# Compare the two Mean Squared Errors (MSEs)
print(paste(MSE.lm, MSE.nn))
```

```
[1] "183730606.293678 132133926.405935"
```

Plot predictions
par(mfrow=c(1,2))

This shows that the network is doing a better job at predicting Gas_prod than the linear model. The prediction from the neural network is **129413554.366771** which is better than the **229551018.949411** obtained from the linear model. We can perform a fast cross visualization in order to be more confident of the result. A visual approach to the performance of the network and the linear model is plotted below.

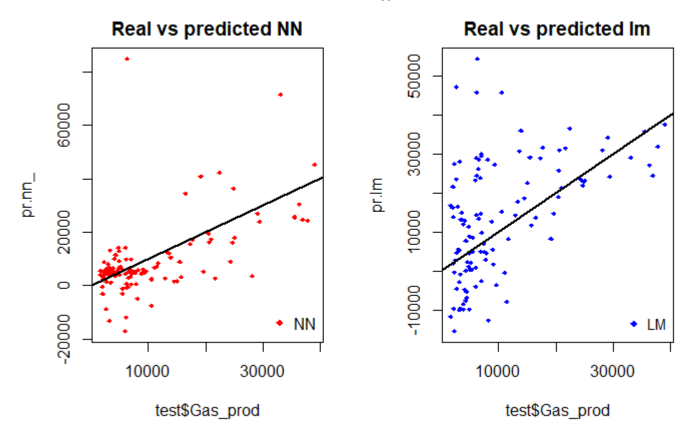
plot(test\$Gas_prod, pr.nn_, col='red', main='Real vs predicted NN', pch=18, cex=0.7)
abline(0,1,lwd=2)

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```
legend('bottomright', legend='NN', pch=18, col='red', bty='n')
plot(test$Gas_prod, pr.lm, col='blue', main='Real vs predicted lm', pch=18, cex=0.7)
```

```
abline(0,1,lwd=2)
legend('bottomright', legend='LM', pch=18, col='blue', bty='n', cex=.95)
```



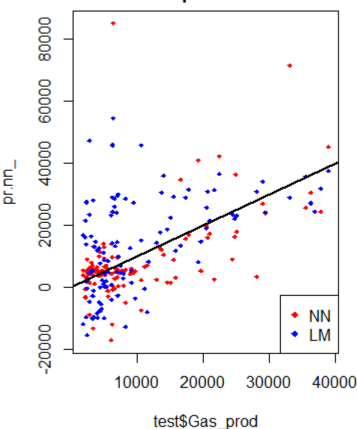
By visually inspecting the plot, we can see that the predictions made by the neural network are more concentrated around the line than those made by the linear model. We can obtain a more useful visual comparison using the code below

```
# Compare predictions on the same plot
plot(test$Gas_prod, pr.nn_, col='red', main='Real vs predicted NN', pch=18, cex=0.7)
points(test$Gas_prod, pr.lm, col='blue', pch=18, cex=0.7)

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abline(0,1,lwd=2)
legend('bottomright', legend=c('NN','LM'), pch=18, col=c('red','blue'))
```

Real vs predicted NN



The comparison of prediction on the same plot is shown above. In analyzing the model above, we allowed the network select the training, validation and testing dataset itself by randomly splitting the data into a train and a test set, then a linear regression model is fit and then tested on the test set.

If we choose to select the training, validation and testing datasets ourselves, we normalize our data and split into training and test data as before.

```
# Neural net fitting
set.seed(440)
mydata <- read.csv("Q4_data.csv", header=TRUE)

#Normalizing the dataset
normalize <- function(x) {
  return ((x - min(x)) / (max(x) - min(x)))
}

maxmindf <- as.data.frame(lapply(mydata, normalize))</pre>
```

```
# Display head of data head(mydata)
```

	Perf_Int <int></int>	Frac_vol <dbl></dbl>	Proppant <int></int>	N.frac <int></int>	Tubing.depth <dbl></dbl>	Casing.depth <dbl></dbl>	FTP <dbl></dbl>	Choke.Size <dbl></dbl>	•
1	280	19454.76	408550	2	7754	8067	185.0	42	2815
2	757	58477.00	407000	1	8843	9081	490.0	40	1915
3	532	27262.00	129400	1	7963	8515	565.0	22	3015
4	345	31622.74	321300	1	7832	8039	615.0	20	2350
5	856	18596.76	390532	2	8602	9065	715.0	24	3250
6	470	34785.71	190000	1	7470	7810	1235.2	10	2250
6 r	ows 1-10	of 15 colum	nns						

Display tail of data
tail(mydata)

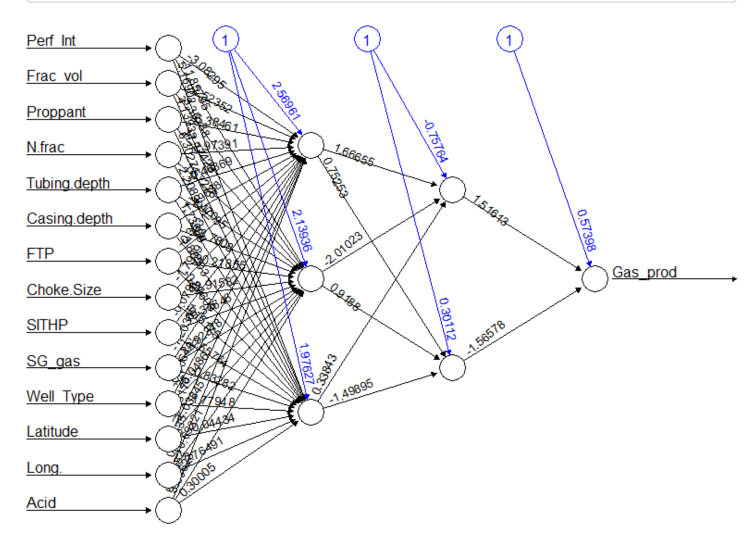
	Perf_Int <int></int>	Frac_vol <dbl></dbl>	Proppant <int></int>	N.frac <int></int>	Tubing.depth <dbl></dbl>	Casing.depth <dbl></dbl>			SI <dbl></dbl>
441	718	52929.00	701000	1	8851	8869	700	32	2200
442	575	28945.00	465800	2	8115	8532	680	24	2450
443	786	52379.00	352572	3	8426	9151	1365	20	2910
444	740	23752.00	203500	2	8149	9200	439	18	2714
445	702	24676.00	294765	2	8484	9245	1365	14	3070
446	372	28928.57	115250	1	7370	7644	1500	9	2865

Since we know the size of the data, we then proceed to test and train the network. We have chosen to split the training and testing dataset into a 70/30 ratio .i.e. 70% (for training) and 30% (for testing) instead of allowing the network do the splitting for us.

```
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```

```
# Training and Test Data
trainset <- maxmindf[1:312, ]
testset <- maxmindf[313:446, ]</pre>
```

We fit the network and arbitrarily decide on the number of hidden neurons. Deciding on the number of hidden layers in a neural network is not an exact science. In fact, there are instances where accuracy will likely be higher without any hidden layers. Therefore, trial and error plays a significant role in this process. One possibility is to compare how the accuracy of the predictions change as we modify the number of hidden layers. If we use 3 and 2 hidden layers, we obtain:



The output from the neural network training is shown above. Our neural network has been created using the training data. We then compare this to the test data to gauge the accuracy of the neural network forecast.

	Gas_prod <dbl></dbl>	Perf_Int <dbl></dbl>	Frac_vol <dbl></dbl>	Proppant <dbl></dbl>	N.frac <dbl></dbl>	Tubing.depth <dbl></dbl>	Casing.depth <dbl></dbl>	
313	0.0115296075	0.14902047	0.006530593	0.08508562	0.0	0.4555120	0.2342327	C
314	0.0045662511	0.05414924	0.005322704	0.04983619	0.2	0.4542529	0.2495274	C
315	0.0041483181	0.14880035	0.005769774	0.05718191	0.0	0.3940963	0.2488400	C
316	0.0029264959	0.08056350	0.004923563	0.01567906	0.0	0.3567431	0.1031105	C
17	0.0009303225	0.11182038	0.007402423	0.07036233	0.3	0.4847510	0.3101908	C
18	0.0042217903	0.15100154	0.003430199	0.03379303	0.1	0.4956631	0.2758206	C
row	s 1-9 of 15 colu	mns						_
								•

nn.results <- compute(nn, temp_test)

The predicted results are compared to the actual results. The code to do this and a snippet of the result is shown below:

#Comparison of Predicted to Actual
results <- data.frame(actual = testset\$Gas_prod, prediction = nn.results\$net.result)
results</pre>

	actual <dbl></dbl>	prediction <dbl></dbl>
313	0.0115296075	0.0019106450
314	0.0045662511	0.0024498630
315	0.0041483181	0.0053135253
316	0.0029264959	0.0061688705
317	0.0009303225	0.0071988651
318	0.0042217903	0.0023350951
319	0.0052641792	0.0032005803
320	0.0065977434	0.0061051029
321	0.0033872906	0.0021572123
322	0.0051492189	-0.0008556349
1-10 of 134 rows	Previous 1 2 3	3 4 5 6 14 Next

We then test the accuracy of the model

Hide

```
#Testing The Accuracy Of The Model
Gas_prod <- trainset$Gas_prod
predicted = results$prediction * abs(diff(range(Gas_prod))) + min(Gas_prod)
actual = results$actual * abs(diff(range(Gas_prod))) + min(Gas_prod)
comparison = data.frame(predicted, actual)

deviation = ((actual-predicted)/actual)
comparison = data.frame(predicted, actual, deviation)</pre>
```

In the above code, we are converting the data back to its original format. Note that we are also converting our data back into standard values given that they were previously scaled using the max-min normalization technique.

We compute the accuracy of network with (3,2) hidden layer. An accuracy of 61.8% on a mean absolute deviation basis (i.e. the average deviation between estimated and actual Gas production per month) was obtained.

```
accuracy = 1-abs(mean(deviation))
accuracy
[1] 0.6187621
```

You can see that we obtain a high accuracy of 61.8 % accuracy using a (3,2) hidden configuration. This is quite good, especially considering that our dependent variable is in the interval format. However, let's see if we can get it higher!

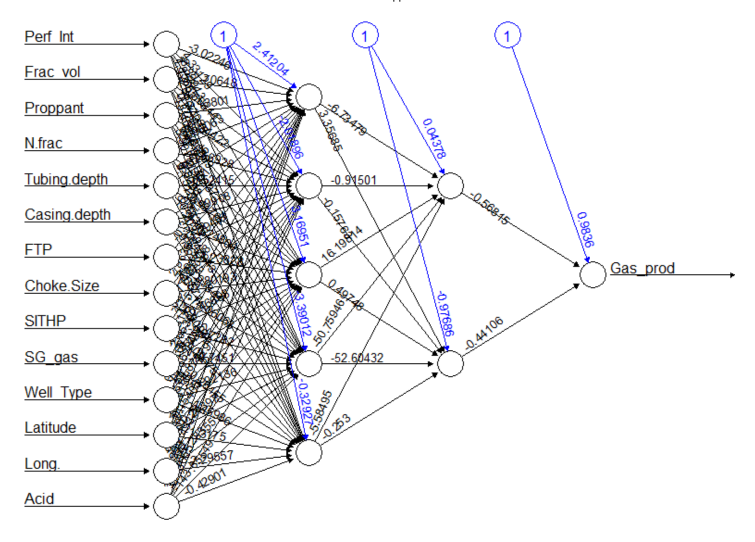
What happens if we now use a (5,2) hidden configuration in our neural network? Here is the generated output:

```
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```

	[,1]
error	3.653272e-03
reached.threshold	8.863541e-03
steps	1.902000e+03
Intercept.to.1layhid1	2.412039e+00
Perf_Int.to.1layhid1	-3.022457e+00
Frac_vol.to.1layhid1	-2.706479e+00
Proppant.to.1layhid1	2.138009e+00
N.frac.to.1layhid1	6.900633e+00
Tubing.depth.to.1layhid1	3.020498e+00
Casing.depth.to.1layhid1	-1.606061e-01
FTP.to.1layhid1	-3.427725e-01
Choke.Size.to.1layhid1	2.963539e+00
SITHP.to.1layhid1	-7.742261e-02
SG_gas.to.1layhid1	-3.814676e-01
Well_Type.to.1layhid1	6.916603e-01
Latitude.to.1layhid1	-8.057662e-01
Longto.1layhid1	-1.101212e+00
Acid.to.1layhid1	1.165234e+01
Intercept.to.1layhid2	-2.018957e+00
Perf_Int.to.1layhid2	4.331287e+00
Frac_vol.to.1layhid2	5.084424e-01
Proppant.to.1layhid2	-7.344221e+00
_ · · ·	-3.569281e+00
N.frac.to.1layhid2	
Tubing.depth.to.1layhid2	-5.241477e-01
Casing.depth.to.1layhid2	-1.099082e+00
FTP.to.1layhid2	-1.129401e+01
Choke.Size.to.1layhid2	3.903377e+00
SITHP.to.1layhid2	4.751302e+00
SG_gas.to.1layhid2	2.653925e+00
Well_Type.to.1layhid2	7.466284e+01
Latitude.to.1layhid2	-1.541364e+00
Longto.1layhid2	9.906799e+00
Acid.to.1layhid2	1.626440e+00
Intercept.to.1layhid3	3.169506e+00
Perf_Int.to.1layhid3	-8.899430e+00
Frac_vol.to.1layhid3	-1.330377e+00
Proppant.to.1layhid3	8.086839e+01
N.frac.to.1layhid3	-6.402368e+00
Tubing.depth.to.1layhid3	-4.970220e-01
Casing.depth.to.1layhid3	-3.024804e+01
FTP.to.1layhid3	-1.423828e+01
Choke.Size.to.1layhid3	4.380193e+01
SITHP.to.1layhid3	1.046698e+01
SG_gas.to.1layhid3	9.484196e+00
Well_Type.to.1layhid3	5.987929e+01
Latitude.to.1layhid3	1.540507e-01
Longto.1layhid3	-7.249328e+01
Acid.to.1layhid3	-8.130646e+00
Intercept.to.1layhid4	-3.390121e+00
Perf_Int.to.1layhid4	7.855518e+00
Frac_vol.to.1layhid4	7.786203e-01
Proppant.to.1layhid4	-2.010102e+01
oppone. co. i i a yniu-	2.0101026101

```
N.frac.to.1layhid4
                           6.809802e+00
Tubing.depth.to.1layhid4
                          4.636598e-01
Casing.depth.to.1layhid4
                          1.147943e+01
FTP.to.1layhid4
                          1.591844e+01
Choke.Size.to.1layhid4
                          -7.406063e+01
SITHP.to.1layhid4
                          -9.272816e+00
SG gas.to.1layhid4
                          -1.074510e+01
Well Type.to.1layhid4
                          -1.392136e+01
Latitude.to.1layhid4
                          2.439652e-01
Long..to.1layhid4
                          1.728355e+01
Acid.to.1layhid4
                          -1.436555e+02
Intercept.to.1layhid5
                          -3.292692e-01
Perf Int.to.1layhid5
                          1.132030e+00
Frac vol.to.1layhid5
                          -1.340956e+01
Proppant.to.1layhid5
                          3.907111e+00
N.frac.to.1layhid5
                           3.751044e+00
Tubing.depth.to.1layhid5 -2.037910e-01
Casing.depth.to.1layhid5
                          9.653307e-01
FTP.to.1layhid5
                           2.657104e+00
Choke.Size.to.1layhid5
                          -9.004901e-01
SITHP.to.1layhid5
                          -1.096745e+00
SG gas.to.1layhid5
                          -2.615459e-01
Well_Type.to.1layhid5
                          -1.459861e+00
Latitude.to.1layhid5
                          1.317501e+00
Long..to.1layhid5
                          -2.295573e+00
Acid.to.1layhid5
                          -4.290123e-01
Intercept.to.2layhid1
                          4.378251e-02
1layhid1.to.2layhid1
                          -6.734787e+00
1layhid2.to.2layhid1
                          -9.150114e-01
1layhid3.to.2layhid1
                          1.619814e+01
1layhid4.to.2layhid1
                          -5.075946e+01
1layhid5.to.2layhid1
                          -5.584953e+00
Intercept.to.2layhid2
                          -9.768579e-01
1layhid1.to.2layhid2
                          3.356849e+00
1layhid2.to.2layhid2
                          -1.576312e-01
1layhid3.to.2layhid2
                          4.974827e-01
1layhid4.to.2layhid2
                          -5.260432e+01
1layhid5.to.2layhid2
                          -2.530036e-01
Intercept.to.Gas prod
                          9.835952e-01
2layhid1.to.Gas prod
                          -5.681484e-01
2layhid2.to.Gas prod
                          -4.410568e-01
```

plot(nn)



	Gas_prod <dbl></dbl>	Perf_Int <dbl></dbl>	Frac_vol <dbl></dbl>	Proppant <dbl></dbl>	N.frac <dbl></dbl>	Tubing.depth <dbl></dbl>	Casing.depth <dbl></dbl>	
313	0.0115296075	0.14902047	0.006530593	0.08508562	0.0	0.4555120	0.2342327	C
314	0.0045662511	0.05414924	0.005322704	0.04983619	0.2	0.4542529	0.2495274	C
315	0.0041483181	0.14880035	0.005769774	0.05718191	0.0	0.3940963	0.2488400	C
316	0.0029264959	0.08056350	0.004923563	0.01567906	0.0	0.3567431	0.1031105	C
317	0.0009303225	0.11182038	0.007402423	0.07036233	0.3	0.4847510	0.3101908	C
318	0.0042217903	0.15100154	0.003430199	0.03379303	0.1	0.4956631	0.2758206	C

```
6 rows | 1-9 of 15 columns
```

```
nn.results <- compute(nn, temp_test)
results <- data.frame(actual = testset$Gas_prod, prediction = nn.results$net.result)
results</pre>
```

	actual <dbl></dbl>	prediction <dbl></dbl>
313	0.0115296075	0.008300917
314	0.0045662511	0.003234412
315	0.0041483181	0.004158014
316	0.0029264959	0.005043247
317	0.0009303225	0.005416683
318	0.0042217903	0.004526156
319	0.0052641792	0.006067041
320	0.0065977434	0.004446578
321	0.0033872906	0.003426161
322	0.0051492189	0.006288651
1-10 of 134 rows	Previous 1 2 3	4 5 6 14 Next

The predicted results are compared to the actual results. We then test the accuracy of the model

```
#Testing The Accuracy Of The Model
predicted = results$prediction * abs(diff(range(Gas_prod))) + min(Gas_prod)
actual = results$actual * abs(diff(range(Gas_prod))) + min(Gas_prod)
comparison = data.frame(predicted,actual)
deviation = ((actual-predicted)/actual)
comparison = data.frame(predicted,actual,deviation)
```

And compute the accuracy of network with (5,2) hidden layer.

```
accuracy = 1-abs(mean(deviation))
accuracy
```

```
[1] 0.7784113
```

Hide

You can see that we obtain 77.84 % network accuracy using a (5,2) hidden configuration. We see that our accuracy rate has now increased to nearly 78 %, indicating that modifying the number of hidden nodes has enhanced our model! This shows that we can increase the accuracy of the network to predict to a higher value by increasing the number of hidden layers.

Model Interpretability

From the solution above, we can observe that neural networks resemble black boxes a lot: explaining their outcome is much more difficult than explaining the outcome of simpler model such as a linear model. Therefore, depending on the kind of application you need, you might want to take into account this factor too. Furthermore, as you have seen above, extra care will be needed to fit a neural network and small changes can lead to different results.

In addition, we showed that neural network is better at predicting "Gas_prod" than the linear regression model. A better mean squared error (MSE) value was obtained using neural network than linear model. Finally, it is possible for neural networks to be more accurate at prediction than regression; however, it will take trial and error of many different hidden layer configurations to get this better prediction.

Conclusion

This project was developed to illustrate the relationship between a dependent variable and several independent variables using **Artificial Neural Network**. We have been able to develop a way such that we were able to select the training, validation and testing datasets. We also investigated how varying number of hidden layers and neurons affect the artificial neural network results. The result using Artificial Neural Network was compared to result from using linear regression. We can conclude that the Artificial Neural Network is better at prediction than the linear regression.