Human vs. Computer: Can we teach the computer to read residual plots?

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Master

by

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This thesis was written using R markdown with relevant code and data accessible with the text.

https://github.com/shuofan18/ETF5550

Software used to conduct this research is R (R Core Team, 2013), Keras (Chollet et al., 2015), ggplot2 (Wickham, 2009)

Declaration

I hereby declare that this thesis contains no material which has been accepted for the award of any other degree or diploma in any university or equivalent institution, and that, to the best of my knowledge and belief, this thesis contains no material previously published or written by another person, except where due reference is made in the text of the thesis.

Shuofan Zhang

Abstract

Residuals plots are a primary means to diagnose statistical models. It requires human evaluation to determine if structure in the plot is consistent with random variation or not. If not, then the diagnosis is that the model has not adequately captured the relationships between response and explanatory variable in the data. This thesis develops a computer vision model to read residual plots. It compares results with a large database of human evaluations. The evaluations were conducted using a protocol called the "lineup" which places residual plots in a formal framework for statistical hypothesis testing. The comparison between computer and human is made on a very restricted and controlled set of residual plot structures. A new small human subject study is also conducted to compare human vs. computer in reading heteroscedasticity.

Chapter 1

Introduction and literature review

"The multiple regression model for cross-sectional data is still the most widely used vehicle for empirical analysis in economics and other social sciences" (Wooldridge, 2015). Detecting possible violations of the Gauss-Markov assumptions is crucial to interpret the data properly, especially in the early stage of analysis. There are several distribution tests that are commonly used, for instance, the Pearson correlation test for detecting linear relationship; the Breusch-Pagan test and White test for investigating heteroskedasticity. But primarily residual plots are the main diagnostic tool and these rely on human evaluation. Because data plots show a lot more information than a single statistic. A good example here would be Anscome's Quartet. "It is a set of four distinct datasets each consisting of 11 (x,y) pairs where each dataset produces the same summary statistics (mean, standard deviation, and correlation) while producing vastly different plots" (Anscombe, 1973). Matejka and Fitzmaurice also did an interesting study on this issue, they used 'datasaurus' data from Cairo (2016) and generated a series of data with same statistics but very different plots (Matejka and Fitzmaurice, 2017).

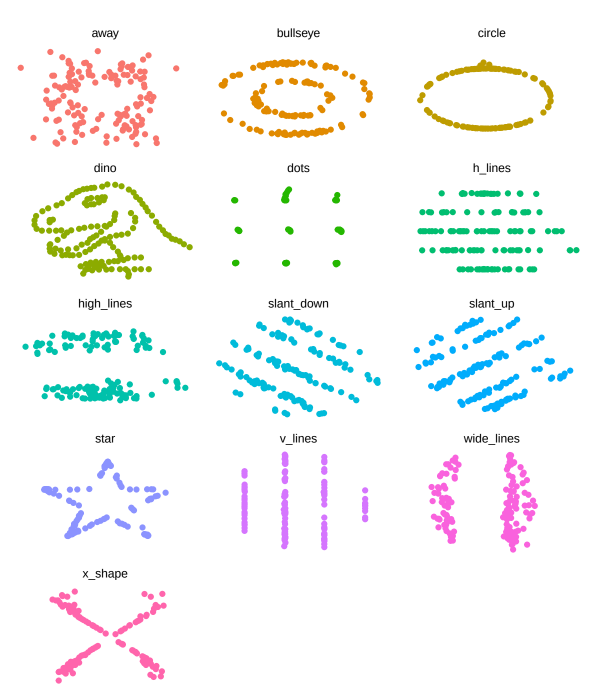


Figure 1.1: Each dataset has the same summary statistics to two decimal places: (E(x)=54.26, E(y)=47.83, Pearson's r=, sd(x)=16.76, sd(y)=26.93)

1.1 Lineup protocol

Former studies have shown that human eyes are sensitive to the systematic patterns in data plots. With proper manipulation, visualized plots can be used as test statistics and perform valid hypothesis test. One example of these protocols that provides inferential validity is lineup which is introduced by Wickham et al. (2010). "The protocol consists of generating 19 null plots (could be other number), inserting the plot of the real data in a random location among the null plots and asking the human viewer to single out one of the 20 plots as most different from the others" (Wickham et al., 2010). If the real plot is chosen, it means the real data is different from the null hypothesis, so we reject the null hypothesis with 5% chance to be wrong (Type I error). Figure 1.2 is an example of lineup. Which plot do you think is the most different? If you choose one, we are 95% confident to reject the no-relationship assumption between the two variables, hp and disp (Simchoni, 2018). This protocol has proved valid and powerful theoretically as well as practically through human experiments, especially when the assumptions for doing conventional tests are violated (Majumder, Hofmann, and Cook, 2013).

The question that arises today is whether we can train a computer to read scatter plots, particularly with a computer vision approach such as deep learning.

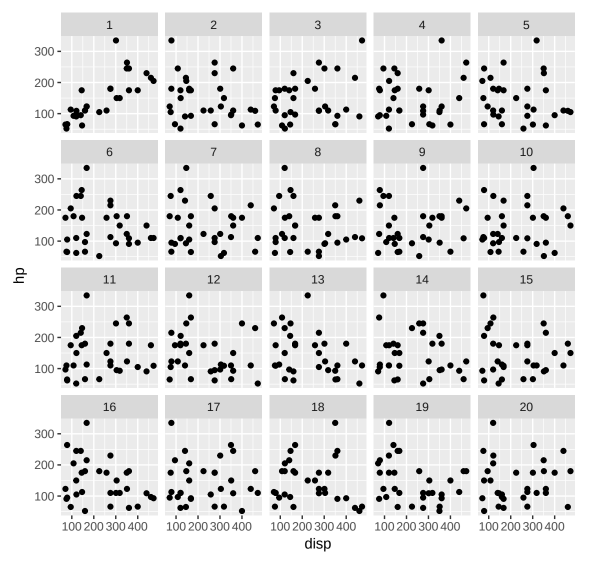


Figure 1.2: Scatterplot lineup example: one plot is the data, the rest are generated from a null model assuming no relationship between the two variables. In this lineup it is easy to see that plot 1, which is the data plot, is different from the rest.

1.2 Computer vision

Motivation for the task is provided in a blog post by Giora Simchoni (Simchoni, 2018). He has designed a deep learning model to test the significance of linear relationship between two variables for samples of size 50. The model reached over 93% accuracy on unseen test data. He also mentioned that the computer fails to pick up a strong non-linear relationship even though the Pearson'r is as high as -0.84 (Simchoni, 2018). So the short conclusion is the computer vision is not perfect, in that it is not as flexible as human vision. As Simchoni explained in his article, the model can only distinguish linear relationship

from no-relationship as trained. However, we think this fact is just another example reflecting the importance of visualization as we discussed above. Strong correlation does not necessarily mean linear relationship. We should always refer to the plot before making any statement. What's more, if we want the model to be more flexible, we could simply adjust our design of training accordingly. Therefore, in this article, we are trying to further Simchoni's study. More specifically, we will build a computer model to perform two hypothesis tests as following.

 H_0

: There are no relationships between the two variables.

 H_1

: There is linear relationship between the two variables where all Gauss-Markov assumptions are met.

 H_0

: There is linear relationship between the two variables where all Gauss-Markov assumptions are met.

 H_1

: There is linear relationship between the two variables where the variance of the error term is not a constant while all other Gauss-Markov assumptions are met.

For ease of exposition, only regression model with one explanatory variable will be considered in this paper, but many of the results can be generalized to other cases including multiple regression model. Because the "statistics" we will use is scatter plot, in terms of teaching the computer of reading the plot, one variable is enough to generate different patterns in that plot for convnets to learn. And this makes the design process much simpler.

The model we will use is the convolutional neural networks, also known as convnets, a type of deep-learning model "almost universally used in computer vision applications" (Chollet and Allaire, 2018). Unlike the classical programming where human input rules, in deep learning paradigm, we provide data and the answers associated with the data. Deep learning algorithm will output the rules, and these rules can then be used on new data to make predictions. We can also think of the deep learning neural network as a complex nonlinear model which could estimate millions of parameters (w) with big enough dataset. As usual regression problem, to get the estimates of unknown parameters (w), we need to provide the model with dependent variable (y_i) and independent variables (\mathbf{x}_i) . In this case, the independent variable will be the images of data plots (in forms of matrices) simulated from the null distribution and the alternative distribution, and dependent variable will be the labels of that plot indicating the true relationship of the original data. Once we have the estimated parameters $(\hat{\mathbf{w}})$, we then can use them to classify unseen data plots, eg. to perform hypothesis tests. The estimation method for deep-learning model is called Backpropagation algorithm which "is a way to train chains of parametric operations using gradient-descent optimization". (Chollet and Allaire, 2018) The gradient-descent optimizer is meant to find the set of parameters such that the cost function reaches its minimum. The form of the cost functions or loss function, should be determined according to each question. In both of the two experiments conducted in this paper, the deep learning model is expected to complete binary classification task, eg. tell "linearly correlated" variables from "independent" variables for the first experiment, tell "heteroskedasticity errors" from "normal errors" for the second experiment. "Crossentropy is usually the best choice (as the loss function) when you're dealing with models that output probabilities" as introduced by Chollet and Allaire (2018). Originated from Information Theory, Crossentropy is a quantity measuring the distance between probability distributions. In deep learning world, it measures the distance between the true distribution and the predictions. Therefore, in this paper, the binary crossentropy loss function will be used. The associated cost function is of the form,

$$J(\mathbf{w}) = -\frac{1}{N} \sum_{i=1}^{N} (y_i \log \hat{y}_i + (1 - y_i) \log (1 - \hat{y}_i))$$

where $\hat{y}_i = g(\mathbf{w} \times \mathbf{x}_i) = \frac{1}{1 + e^{-\mathbf{w} \times \mathbf{x}_i}}$ and g(z) is the logistic function.

1.3 Comparing human vs. computer

Chapter 2 will compare computer performanace against database of human evaluation in reading linear relationship. Steps of constructing computer experiment will be discussed, Turk's study will be explained, the comparison results will be given. Chapter 3 will compare computer performance against the results from the new human subject study. Details of this new human subject study will be provided. The results will also be presented.

Chapter 2

Comparing computer performance against database of human evaluation

A database of human evaluations of scatterplots of residuals against fitted is available from prior studies. This is used to compare the performance of the computer model. The computer model is trained on a broader parameter simulation framework, and tested on the same data as the human evaluations.

2.1 Turk study explanation

A large database of results from a human subjects was collected examine the performance of the lineup protocol relative to a classical tests. The work is published in Majumder, Hofmann, and Cook (2013). This database forms the basis of the test set used to examine the computer model performance.

In Majumder, Hofmann, and Cook (2013), "three experiments were conducted to evaluate the effectiveness of the lineup protocol relative to the equivalent test statistic used in the regression setting." In each experiment, they simulated data from a controlled setting and then generated associated lineup for human to evaluate.

Sample size (n)		Error SD(sigma)	Experiment 1 beta2	Experiment 2 beta1
	100	5	0,1,3,5,8	0.25, 0.75, 1.25, 1.75, 2.75
	100	12	1,3,8,10,16	0.5, 1.5, 3.5, 4.5, 6
	300	5	0,1,2,3,5	0.1, 0.4, 0.7, 1, 1.5
	300	12	1,3,5,7,10	0, 0.8, 1.75, 2.3, 3.5

Table 2.1: Combination of parameter values used for simulation in Turk's study.

The controlled model in their first experiment is

$$Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \epsilon_i$$

where $\beta_0 = 5$, $\beta_1 = 15$, $X_1 \sim Poisson(\lambda = 30)$, $\epsilon_i \sim N(0, \sigma^2)$, i = 1, 2, ..., n. While in the null model $\beta_2 = 0$, and the null data was generated by simulating from $N(0, \hat{\sigma}^2)$. This experiment was aimed to test the ability of human on detecting the effect of X_2 .

Their second experiment is very similar to the first one, but there is only one continuous variable X_1 on the right hand side. The actual data model is

$$Y_i = \beta_0 + \beta_1 X_{i1} + \epsilon_i$$

where $\beta_0 = 6$, $X_1 \sim N(0,1)$, and the null data was generated from $N(X\hat{\beta}, \hat{\sigma}^2)$.

The third experiment in their paper contains contaminated data where the actual data were in fact generated from two different specifications.

$$Y_{i} = \begin{cases} \alpha + \beta X_{i} + \epsilon_{i} & X_{i} \sim N(0,1) \ i = 1, ..., n \\ \lambda + \eta_{i} & X_{i} \sim N(\mu, 1/3) \ i = 1, ..., n_{c} \end{cases}$$

where $\epsilon_i \sim N(0, \sigma)$, $\eta_i \sim N(0, \sigma/3)$, $\mu = -1.75$, $\beta \in (0.1, 0.4, 0.75, 1.25, 1.5, 2.25)$. And n = 100, $n_c = 15$, alpha = 0, $\lambda = 10$, $\sigma = 3.5$. The null plots were generated from $N(0, \hat{\sigma}^2)$.

Other parameters in the "actual data sets" of experiment one and two are shown in the table below.

Their experiment 2 examined the performance of humans in recognising linear association between two variables, in direct comparison to conducting a t-test of H_0 : $\beta_k = 0$ vs

 $H_a: \beta_k \neq 0$ assessing the importance of including variable k in the linear model. An example linear is shown in Figure 2.6. For this linear, 63 of the 65 people who examined it selected the data plot (position 20) from the null plots. There is clear evidence that the data displayed in plot 20 is not from $H_o: \beta_k = 0$.

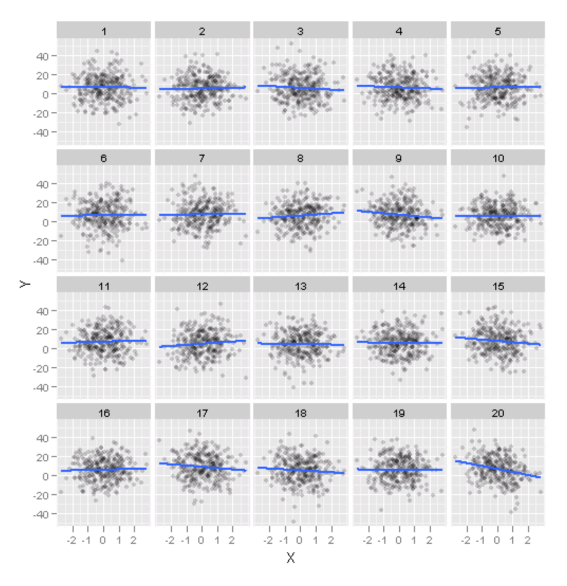


Figure 2.1: One of 70 lineups used in experiment 2 Majumder et al (2012). Of the 65 people who examined the lineup, 63 selected the data plot, which is in position 20.

This experiment 2 utilised 70 lineups of size 20 plot, with varying degrees of departure from the H_0 : $\beta_k = 0$. There were 351 evaluations by human subjects. These results will be used for comparison with the deep learning model.

2.2 Linear relationship simulation

The design for this model is similar to what Simchoni (2018) did in his blog but is tailored to compare the computer performance with the Turk study results.

The model is designed as:

$$Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i, i = 1, \dots, n$$

which is the same with the data generating model of Turk's experiment 2. And all the parameters in our model will be designed to cover the range used in the second experiment in Majumder, Hofmann, and Cook (2013). The relevant parameters are generated using the following specification.

- $X \sim N[0, 1]$
 - Distributions of X has impact on the shape of the scatters. For instance, if X is generated from a uniform distribution, then the plots will look like a square especially when the sample size is large; while more like a circle if X follows normal distribution.
- $\beta_0 = 0$ Intercept is set to be zero, because it will not change the patterns in the data plots.
- $\beta_1 \sim U[-10, -0.1] \cup [0.1, 10]$ β_1 is designed to be uniformly generated from -10 to 10 (excluding -0.1 to 0.1).
- $\varepsilon \sim N(0, \sigma^2)$ where $\sigma \sim U[1, 12]$ ε is designed to be uniformly generated from 1 to 12.
- n = U[50,500]
 The sample sizes of each data set vary from 50 to 500.

Figure 2.2 shows four example plots generated using the specifications above. Under this controlled structure, a total number of 240,000 data sets are simulated. The histograms of the simulated n, β , σ , the estimated p value and the scatter plots of β against n, σ against n in figure 2.3 show good coverage over all the values.

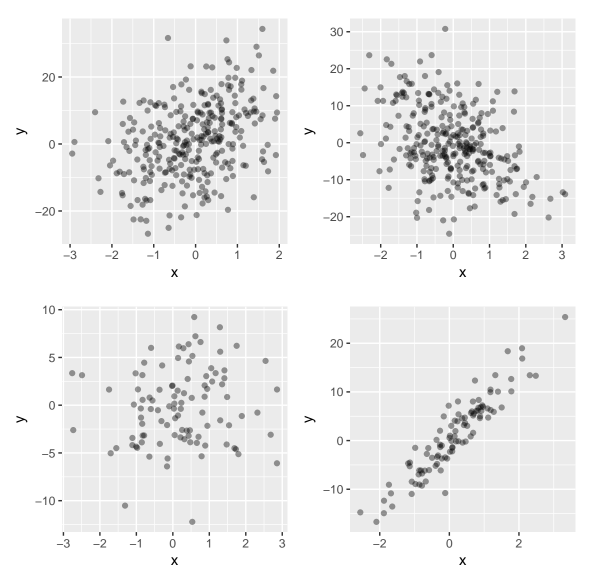


Figure 2.2: Four examples of data plots generated from the classic linear model.

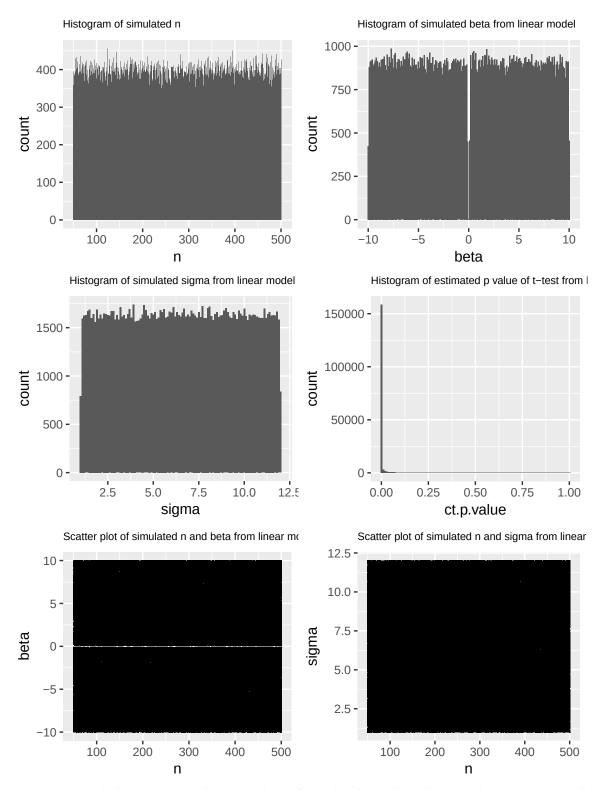


Figure 2.3: The histograms and scatter plots of simulated variables show good coverage over all the values.

2.3 Null plot simulation

This is the null scenario in our first experiment, eg. the two variables under tested are independent of each other. If the data arises from this situation, then the data plots will not show any systematic patterns theoretically.

The model is designed the same as the linear model:

$$Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i, i = 1, \dots, n$$

with elements of the model generated using the same specification as the linear model, except

•
$$\beta_1 = 0$$

The coefficient of X_i is always zero. So X_i and Y_i are uncorrelated of each other.

Figure 2.4 are four example plots generated using the specifications above. Same as the linear model simulation, a total number of 240,000 data sets are simulated under this structure. The histograms of the simulated n, β , σ , the estimated p value and the scatter plots of β against p0 against p1 in figure 2.5 show good coverage over all the values.

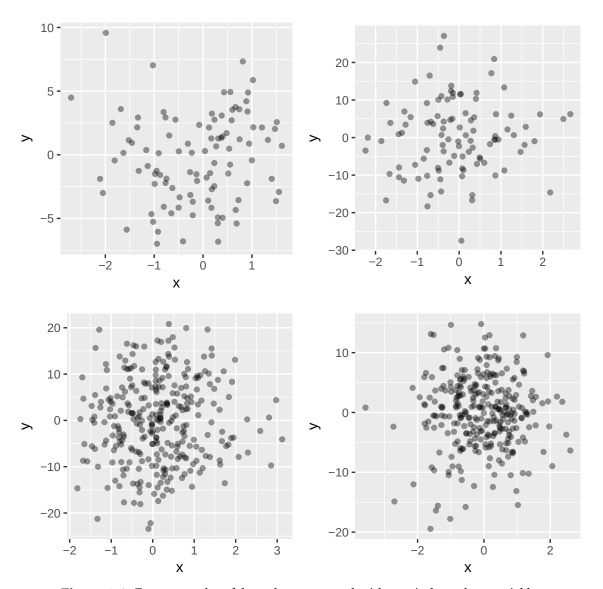


Figure 2.4: Four examples of data plots generated with two independent variables

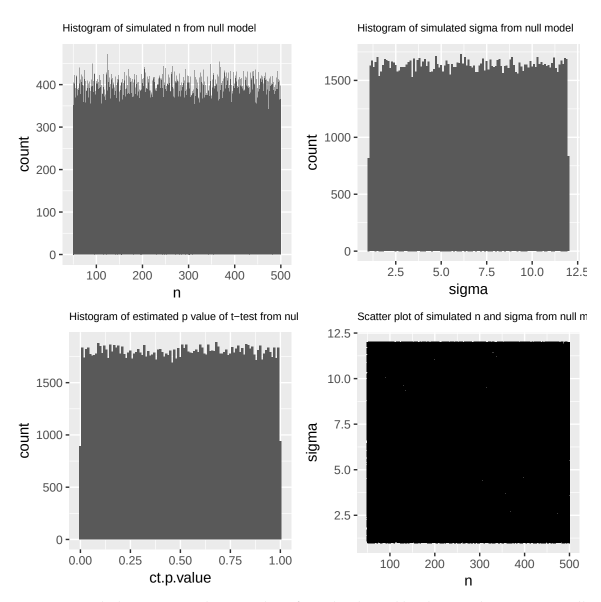


Figure 2.5: The histograms and scatter plots of simulated variables show good coverage over all the values.

2.4 Computer model

A convolutional neural network will be used to build the computer model because it proves powerful in all kinds of computer vision applications. The plots used for training and testing in this section is the scatter plot between the dependent variable Y and the independent variable X. The R package Wickham (2009) is used to generate the plots. All plots are saved as png and will be resized to have width and height both equal to 150 pixels. This size is similar to the plot size used in the lineup for human evaluation. As for the labels given to each image, we use the true population as the samples' identification directly. It is true that there will be some undesired patterns formed out of randomness, especially when the sample size is small. Unlike what Simchoni did in his post, no conventional tests will be used to sort out the "significant" observations. Because the answer to the question that if the deep learning model can distinguish from patterns formed by chance and by nature is also interested.

As mentioned above, 240,000 data sets are generated for each of the two groups in the first experiment. 100,000 of them are set apart for training. Another 40,000 of the data sets are set apart as validation set in order to monitor during training the accuracy of the model on data it has never seen before. And the leftover (100,000 data sets) become the unseen test set. We make the test set so large that we can compare the performance of the convnet with the conventional t-test properly.

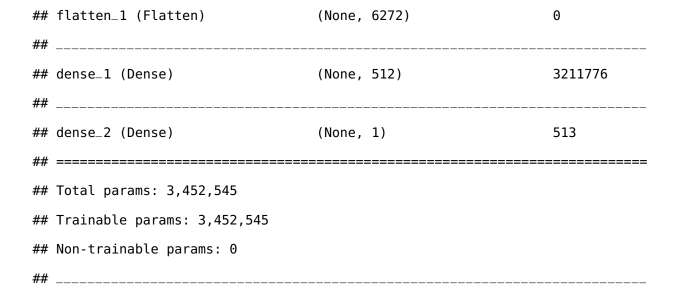
"A convnet takes as input tensors of shape (image height, image width, image channels)." (Chollet and Allaire, 2018) In this case, the input tensors are of shape $150 \times 150 \times 1$ because they are grayscale images. Therefore the convnet will be configured to process inputs of size (150, 150, 1). We'll do this by passing the argument input_shape = c(28, 28, 1) to the first layer.

```
layer_conv_2d(filters = 64, kernel_size = c(3, 3), activation = "relu") %>%
layer_max_pooling_2d(pool_size = c(2, 2)) %>%
layer_conv_2d(filters = 128, kernel_size = c(3, 3), activation = "relu") %>%
layer_max_pooling_2d(pool_size = c(2, 2)) %>%
layer_conv_2d(filters = 128, kernel_size = c(3, 3), activation = "relu") %>%
layer_max_pooling_2d(pool_size = c(2, 2)) %>%
layer_flatten() %>%
layer_dense(units = 512, activation = "relu") %>%
layer_dense(units = 1, activation = "sigmoid")
model
```

##	Model		
##			
##	Layer (type)	Output Shape	Param #
##			
##	conv2d_1 (Conv2D)	(None, 148, 148, 32)	320
##			
##	<pre>max_pooling2d_1 (MaxPooling2D)</pre>	(None, 74, 74, 32)	0
##			
##	conv2d_2 (Conv2D)	(None, 72, 72, 64)	18496
##			
##	<pre>max_pooling2d_2 (MaxPooling2D)</pre>	(None, 36, 36, 64)	0
##			
##	conv2d_3 (Conv2D)	(None, 34, 34, 128)	73856
##			
##	<pre>max_pooling2d_3 (MaxPooling2D)</pre>	(None, 17, 17, 128)	0
##			
##	conv2d_4 (Conv2D)	(None, 15, 15, 128)	147584
##			
##	max_pooling2d_4 (MaxPooling2D)	(None, 7, 7, 128)	Θ
##			

Table 2.2: Performance of three checkpoints of the deep learning model on the unseen test set, where the power means the accuracy on the linear ones, (1-alpha) is the accuracy on the no-relationship ones. All numbers have been rounded to four decimal places.

Tests	Power	1-Alpha	Overall accuracy
4 epoch	0.8790	0.9757	0.9273
6 epoch	0.8910	0.9569	0.9239
10 epoch	0.8826	0.9687	0.9257
T-test	0.9003	0.9487	0.9245



10 epoches (1 epoch = 1 iteration over all samples) will be done for training. The model specification given by each epoch will be saved, the one gives the highest test power is chosen to represent the computers.

"target_size: Tuple of integers (height, width), default: (256, 256). The dimensions to which all images found will be resized." Chollet et al. (2015)

All convolutional neural network related work will be done by Keras (Chollet et al., 2015) package in R.

From the plot of the traning history, we can see the fourth specification gives the overall best performance with lowest loss and highest accuracy on validation set. Its accuracy on the unseen test set is also the best among other models. After the fourth checkpoint, the model starts to overfit.

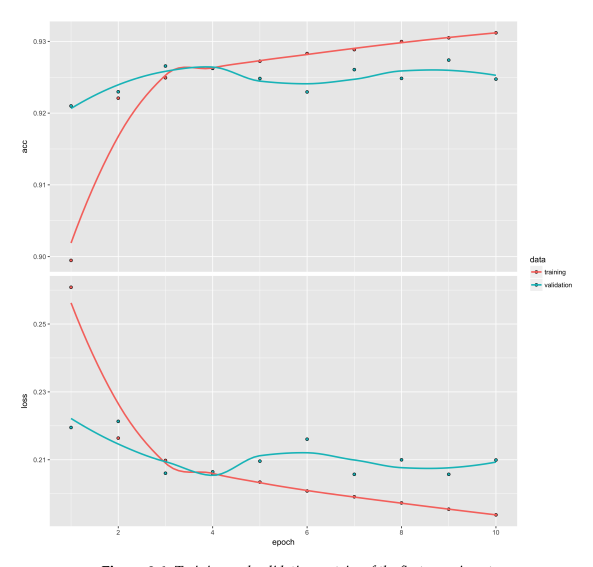


Figure 2.6: *Training and validation metrics of the first experiment*

However, since the majority of the data plots in Turk's experiment have been generated with linear relationship, we will choose the sixth model here because it has the highest test power which provides the deep learning model a better chance. What's more, the α of the sixth model, which is the probability of Type I error, is the closest to 0.05. We can see the sixth model's performance is quite close to the conventional T-test.

2.5 Comparing results

The conclusion of human evaluation is calculated using the p-value calculation introduced in section 2 of Majumder, Hofmann, and Cook (2013), the null hypothesis is rejected if the calculated p-value is smaller than 0.05.

Table 2.3: Accuracy of testing the 70 data plots evaluated by human computer and the conventional *t-test*

Tests	No. of correct	Accuracy
human	47	0.6714
computer	44	0.6286
t-test	43	0.6143

The α of the conventional t-test we used in this comparison is 0.05.

From this table, we can see human evaluation achieves the best accuracy on the Turk's experiment data set. Deep learning model and the conventional t-test perform similarly.

2.6 Aside discussion related to the comparing results

As we can see from the table, t-test and deep learning model behave quite similarly on both the test data set and Turk's experiment data. It is possible that the "Backpropagation algorithm" used in deep learning model is equivalent to maximum likelihood estimation in terms of linear regression problem in this case. Because all the error terms are following normal distribution as designed, they are also equivalent to the ordinary least square estimation. Since the fourth-epoch model behaves the best in all 10 models, and its estimated Type I error (α) is not 0.05. We think the deep learning model may be in fact just finding the "best" α for t-test in this experiment, where "best" means giving the highest expected accuracy of the null and alternative hypothesis. To confirm this idea, a small simulation experiment was conducted to search for the "best" α for t-test. The data was simulated using the same specification as in the section "Linear relationship simulation" and "null plots simulation". The range of the α is from (0.005, 0.010, 0.015, ..., 0.100), with 200,000 iterations, the "best" α is found to be α = 0.02 which is very close to the one suggested by the fourth-epoch model 0.02435.

Chapter 3

New experiment comparing human vs. computer on reading heteroskedasticity

3.1 Heteroskedasticity simulation

Linear model with heteroskedasticity is the model implied in the alternative hypothesis of the second experiment in this paper, where the constant variance assumption of the linear model is violated while all other conditions are met. By the definition given in Wooldridge (2015), "The homoskedasticity states that the variance of the unobserved error, u, conditional on the explanatory variables, is constant. Homoskedasticity fails whenver the variance of the unobserved factors changes across different segments of the population, where the segments are determined by the different values of the explanatory variables." There are countless types of heteroskedasticity since the "change of the variance" could be related to "the explanatory variables" in various ways. It is not feasible to list out all kinds of heteroskedasticity by a single function. For simplicity, we will focus on one example of them, a linear correlation between the explanatory variable X and the standard deviation of the error term. The conclusion from this experiment though can be generalized to more complicated cases.

A new human experiment will be rendered to produce the human's performance on detecting heteroskedasticity by reading residual plots. After the deep learning model is trained on the data generated from this section, it will be tested on the same data set that is used for the human experiment. And the accuracy of the model will be compared to the human's performance from this new human experiment. To provide a reference to evaluate how computer perform in detecing heteroskedasticity issues, a special case of the White test will be used. Each image in test set will be evaluated using White test. The associated accuracy of White test will be compared to computers' accuracy on the same set.

The model structure is the same with the classic linear model:

$$Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i, i = 1, \dots, n$$

with elements of the model generated by the following processes

- $X \sim U[-1, 1]$ To better present the heteroskedasticity in the data, a uniform distribution of X is used instead of normal distribution. The range is set to be small (from -1 to 1) in order to produce the data with weak heteroskedasticity more frequently.
- $\beta_0 = 0$ Intercept is set to be zero for the same reason stated above.
- $\beta_1 \sim U[0.5, 1] \beta_1$ has little impact in this case so it is set to be uniformly generated from 0.5 to 1. Because the residual plot but not data plot will be used here, therefore, the information contained in β_1 will be extracted by the linear regression we fit to the data.
- $\varepsilon \sim N(0, (aX + v)^2)$ The variance of the error term is a quadratic function of the explanatory variable which controls the magnitude of heteroskedasticity in the model.
- a ~ U(-5, -0.05) ∪(0.05, 5) The parameter a here, following uniform distribution from -5 to 5 (excluding -0.05 to 0.05), is the correlation coefficient between X and the standard deviation. Larger a gives stronger heteroskedasticity. This range is wide enough for our purpose.

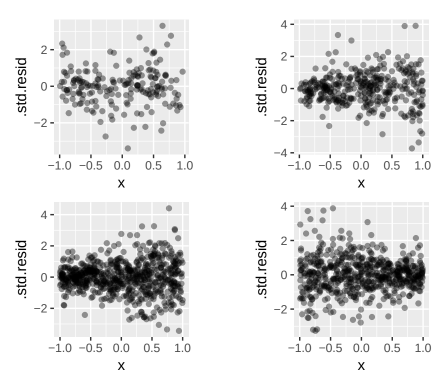


Figure 3.1: (#fig:heter_plot)Four examples of residual plots generated from linear model with heteroskedasticity

- $v \sim N(0, 1)$ This new error term is added to the variance of ε so the relationship between the data can be more flexible.
- $n \sim U[20, 1500]$ The sample sizes will be randomly generated from 20 to 1500. We choose 20 to 1500, because when the sample size is smaller than 20, there is hardly any systematic patterns to see. And 1500 is large enough to give a good description about the true relationship within the data, and is light enough to be processed. Since we make the transparency of the points to be 0.4, more points being added to the plot will just make the plot looks darker.

In general, the choice of the parameters is an empirical work. Primarily, we want the residual plots to show more variation; on the other hand, we need to limit the range of these parameters in order to keep the key features in the data.

3.2 Human subject experiment

3.3 Deep learning model training

For the second experiment, a linear model will be fit to the data firstly. Residuals from the fitted model will be standardized and extracted. The residual plot will be made of standardized residuals against X.

3.4 Results

Chapter 4

Conclusion

Based on the results from the two experiments,

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