Quantitative Risk Management Project 2

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Problem 1

Remember from last week we discussed that skewness and kurtosis functions in statistical packages are

often biased. Is your function biased? Prove or disprove your hypothesis.

Answer

The python has both unbiased and biased estimator of skewness and kurtosis. The difference between the

unbiased kurtosis estimator and the biased kurtosis estimator is significant. The biased kurtosis estimator

is biased when the sample size is small, but as the sample size increases, the biased kurtosis estima-

tor becomes unbiased. Even though the result of the kurtosis estimator is reasonable, the difference

between the unbiased skewness estimator and the biased skewness estimator is very small. Both are

unbiased in the experiment. There are some possible reasons for this, which I will discuss in the following

analysis.

1. Background

It is up to the user to decide whether the estimation of skewness and kurtosis in scipy, a Python package,

is biased or not. There is a parameter called bias in the estimation function. It can be used to choose whether

to use an unbiased estimator or not. If you set the bias to False, then the calculations will be corrected for the

bias and the value that is calculated will be the adjusted Fisher-Pearson standardised coefficient of moments:

 $G_1 = \frac{k_3}{k_2^{\frac{3}{2}}} = \frac{\sqrt{N(N-1)}}{N-2} \frac{m_3}{m_2^{\frac{3}{2}}}$

2. Experiment

Objection: Test if the kurtosis and skewness calculated by scipy package is biased.

1

Step:

- 1. Sample 5,10,100,100000 standardized random normal values separately.
- 2. Calculate the kurtosis and skewness by setting bias to False and True separately.
- 3. Sample the kurtosis and skewness by repeating <steps 1 and 2> 100 times.
- 4. Calculate the first 2 moments of kurtosis and skewness, which are mean kurtosis $\overline{\mu_k}$, mean skewness $\overline{\mu}_{sk}$, standard deviation $\overline{\sigma}_k$, standard deviation $\overline{\sigma}_{sk}$.
- 5. Set the significance level $\alpha=0.005$. Given the null hypotesis as $h\mu_k=0$ and $\mu_{sk}=0$, calculate the T statistic of skewness and kurtosis separately. (T-test)
- 6. Given the condition of two-side test, calculate the p-value of skewness and kurtosis.
- 7. If the p-value is lower than significance level, then we reject the null hypotesis that skewness or kurtosis is unbiased.

3. Results:

1. Sample size = 5.

Table 1: Sample moments of estimator When sample size = 5

Characteristic	Description	Unbiased skewness	Unbiased kurtosis	Biased skewness	Biased kurtosis
Moments	Mean	-0.14544599	0.23005863	-0.09756814	-0.94248534
	Variance	0.92341645	4.43911036	0.4155374	0.2774444
P-value	_	0.13332058	0.27751873	0.13332058	0.

2. Sample size = 10.

Table 2: Sample moments of estimator When sample size = 10

Characteristic	Description	Unbiased skewness	Unbiased kurtosis	Biased skewness	Biased kurtosis
Moments	Mean	-0.04352551	-0.13300726	-0.03670394	-0.62069098
	Variance	0.43281135	1.57664148	0.30777696	0.5044738
P-value	_	0.50976544	0.29205202	0.50976544	6.1284311e-14

Reasoning

Let's first look at the unbiased skewness and the biased skewness. We can see that although their means are different, their p-values are the same and both are above the significant level, which means that we should accept the null hypothesis. They are statistically unbiased. However, this contradicts our past experience and doesn't make sense.

Firstly, I'm inclined to attribute such a confusing result to rounding error, because the sample means

of unbiased and biased skewness are too close for the computer to differentiate their p-values.

Secondly, we get a sample from a standard normal distribution, which means that the sample is a random variable. Its randomness leads to the randomness of the p-value. Therefore, such a result could be just a coincidence. Therefore, I did some additional experiments and I find that the majority of p-value is greater than the significant level.

Third, the distribution of the unbiased skewness estimator and the biased skewness estimator is not normal. T-test requires its random variable to be normal or approximately normal. If its random variable does not fulfil such condition, then T-test requires large sample to make the statistic close to t-distribution. In fact, under these circumstances, the distribution is asymptotically normal. However, our sample size is only 5 and 10, which would reduce the efficiency of the t-test. In addition, the result is not very precise and not very stable.

Then we focus on the unbiased kurtosis and the biased kurtosis. We could quickly find that the mean of those two are more different and their p-value obviously suggest that unbiased kurtosis estimator are unbiased, biased kurtosis estimator are biased.

3. Sample size = 100.

Table 3: Sample moments of estimator When sample size = 100

Characteristic	Description	Unbiased skewness	Unbiased kurtosis	Biased skewness	Biased kurtosis
Moments	Mean	-0.01493056	0.01284804	-0.01470566	-0.04719138
	Variance	0.05716795	0.20712172	0.05545869	0.18720098
P-value	-	0.53376663	0.77829632	0.53376663	0.27804886

4. Sample size = 100000.

Table 4: Sample moments of estimator When sample size = 100000

Characteristic	Description	Unbiased skewness	Unbiased kurtosis	Biased skewness	Biased kurtosis
Moments	Mean	-0.00150312	0.00083446	-0.0015031	-0.00077442
	Variance	6.09705474e-05	0.00022669	6.09705474e-05	0.00022667
P-value	-	0.05709829	0.58067351	0.05709829	0.60813631

Reasoning

As we can see, the biased estimator tends to converge to the unbiased estimator as the sample size increases. The variance of the biased and unbiased estimators decreases a lot and their mean is very close. Moreover, their p-values are all above the significant level, so we could consider both as unbiased.

4. Conclusion

When the sample size is small, The kurtosis estimator behaves as expected, but the skewness estimator does not. An unbiased kurtosis estimator is statistically significant in unbias and a biased kurtosis estimator is biased. For skewness, both are unbiased, but I will attribute this to rounding error and small sample size. When the sample size is large, the biased estimator converge to unbiased estimator and biasedness disappears.

Problem 2

- 1. Fit the data in problem2.csv using OLS and calculate the error vector. Look at its distribution. How well does it fit the assumption of normally distributed errors?
- 2. Fit the data using MLE given the assumption of normality. Then fit the MLE using the assumption of a T distribution of the errors. Which is the best fit?
- 3. What are the fitted parameters of each and how do they compare? What does this tell us about the breaking of the normality assumption in regards to expected values in this case?

Answer

Problem 2.1

The residuals produced by OLS is not normally distributed.

reasoning

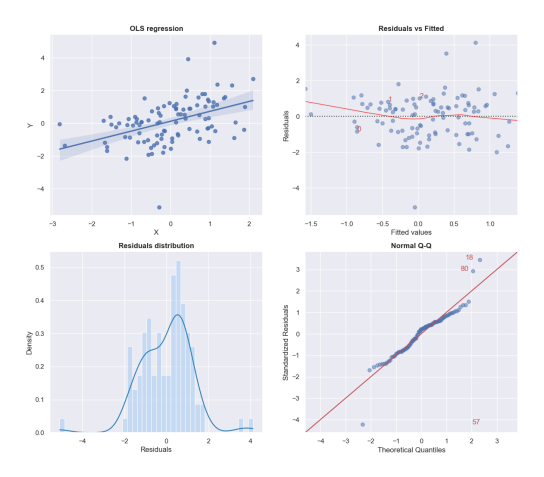
Given the dataset, we use OLS to fit the data and here is our reslut:

Now let's focus on the distribution of the residuals. We might find that it is more likely to be a left skewed distribution than a normal distribution. With such a guess, we could see that the Q-Q plot also shows that the residuals want to leave the diagonal line when they are away from 0.

With a visual check, we could now perform a normal test on the residuals. We use a method based on D'Agostino and Pearson's test, which combines skewness and kurtosis to produce an omnibus test of normality. And we find that the p-value = 0.0008475313945909529, which is less than our expected significance level of 0.05. We therefore reject the null hypothesis. The residuals are not normally distributed.

Problem 2.2

The MLE assuming T-distribution of error terms is better than that assuming normality.



reasoning

First, we use MLE assuming the normality of errors to fit the data. Then, assuming that the error term is T-distributed, we use MLE to run the regression. We could get the corresponding parameters for each MLE and we use them to caculate the residuals. Here are some visuals for residuals distribution and linear moded:

From the images above, we could see that the difference between these two MLE is not very large and their residuals have a large amount of overlap. As a result, it is not easy to tell from the image which of the MLE is the better one. One interesting thing is that when we use MLE given error is T distributed, its residuals are still not normal distribution. It seems more left biased distributed.

Now, let's look at some useful numbers. Even though direct images could not help us figure out which is better, those numbers may give us some reasonable explanation that T is more suitable for regression this time.

Assuming that the error term is T-distributed, by MLE we could get the following results:

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The R^2 and $adjusted R^2$ of those two MLE are very same, but the AIC and BIC of T-MLE is smaller that Normal-MLE. Therefore, we could conclude that T-MLE is better for regression this time.

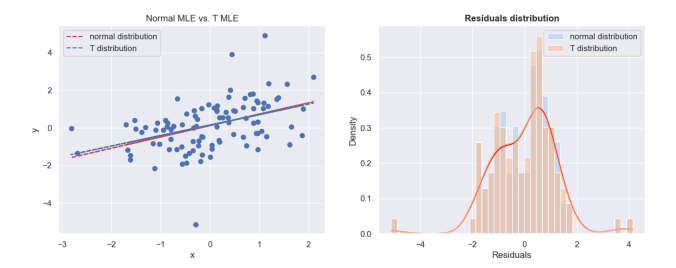


Table 5: MLE under normal error assumption

Characteristic	Description	Value
Parameter	eta_1	0.6052
	eta_0	0.1198
	Variance	1.1984
Goodness of fit	\mathbb{R}^2	0.1946
	$adjusted \ R^2$	0.1864
Information Criteria	AIC	323.9841
	BIC	329.1945

Problem 2.3

Not all error terms follow a normal distribution in our normal life. We could use different assumptions about error terms and compare their AIC, BIC or other statistics to see which one is more reasonable.

reasoning

If error follows a normal distribution:

$$y = 0.1198 + 0.6052x + \epsilon, \qquad \epsilon \sim N(0, 1.1984)$$

If error follows a T distribution:

$$y = 0.1426 + 0.5576x + \epsilon, \qquad \epsilon \sim T(6.2766, 0, 0.9713)$$

From **Problem 2.2**, we could see T distribution is better.

Table 6: MLE under T error assumption

Characteristic	Description	Value
Parameter	eta_1	0.5576
	eta_0	0.1426
	Degree of freedom	6.2766
	Variance	0.9713
Goodness of fit	R^2	0.1931
	$adjusted \ R^2$	0.1849
Information Criteria	AIC	314.9459
	BIC	320.1563

Problem 3

Simulate AR(1) through AR(3) and MA(1) through MA(3) processes. Compare their ACF and PACF graphs. How do the graphs help us to identify the type and order of each process?

Answer

1. AR Process

For AR model, we choose to simulate the following process:

$$x_t = 0.75 + 0.4x_t - 1 + \epsilon, \qquad \epsilon \sim N(0, 0.8)$$

$$x_t = 0.75 + 0.4x_t - 1 + 0.3x_{t-2} + \epsilon, \qquad \epsilon \sim N(0, 0.8)$$

$$x_t = 0.75 + 0.2x_t - 1 + 0.3x_{t-2} + 0.2x_{t-3} + \epsilon, \qquad \epsilon \sim N(0, 0.8)$$

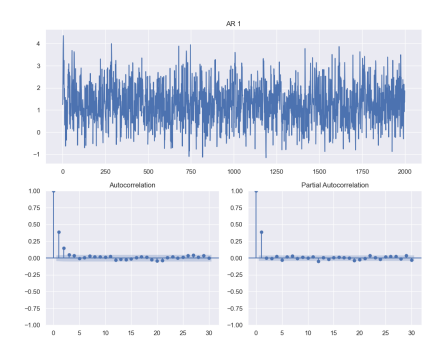
Meanwhile, we calculate and plot the autocorrelation function (ACF) and the partial autocorrelation function (PACF). We can easily see that for the AR(1) model, the PACF becomes insignificant after the first lag; for the AR(2) model, the PACF becomes insignificant after the second lag; for the AR(3) model, the PACF becomes insignificant after the third lag. Their ACF also falls to zero or becomes insignificant more slowly. Therefore, we can safely conclude that the PACF of the AR(p) model would become insignificant after the p^{th} lag and its ACF would slowly converge to zero.

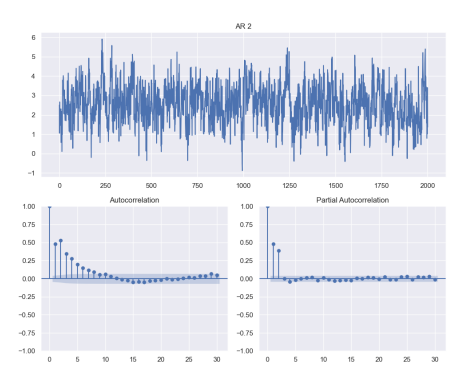
2. MA Process

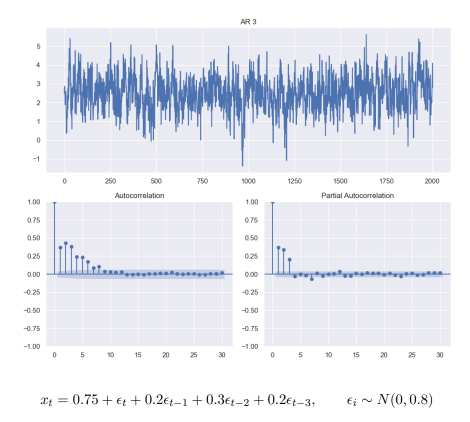
For AR model, we choose to simulate the following process:

$$x_t = 0.75 + \epsilon_t + 0.4\epsilon_{t-1}, \qquad \epsilon_i \sim N(0, 0.8)$$

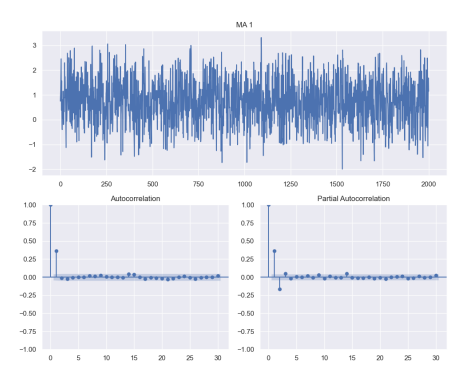
 $x_t = 0.75 + \epsilon_t + 0.4\epsilon_{t-1} + 0.3\epsilon_{t-2}, \qquad \epsilon_i \sim N(0, 0.8)$

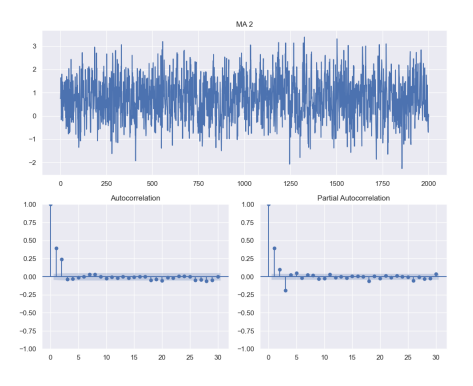


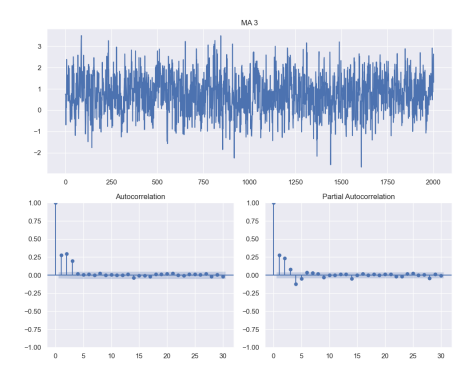




Also, ACF and PACF are calculated and plotted. From the following figures, we can find that The ACFs are not significant after the first lag for MA(1), after the second lag for MA(2) and after the third lag for MA(3). However, there is no obvious pattern about PACF. Therefore, for MA(\mathbf{q}) model, we could guess their oder by looking at ACF. If its q^{th} lag of ACF becomse insignificant and its PACF slowly converge to zero, then it has great possibility to be MA(\mathbf{q})







Feedback

- 1. Full Credit
- 2. Full Credit
- 3. -0.25 I would suggest using python package, some of your graph seems wrong(MA(3) for example)
 - (a) **AR:** abs(ACF) for AR is always non-zero, decaying through time. PACF for AR is only non-zero for the number of lags in the model.
 - (b) **MA:** ACF for MA is only non-zero for the number of lags in the model. abs(PACF) for MA decays through time.

See the formuals for autocovariance (which is scaled correlation) in the notes.(Feixed my code bugs.)