The strong contour line relationship between two-sided p values of simple RCT two sample t-tests and BIC based Bayes factor  $P_{BIC}(D|H0)$  outputs for fixed sample sizes

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#### **Executive Summary**

For simple Randomised Control Trial (RCT) analysis there is a strong non-linear relationship between the two-sided p values of two sample t-tests and the corresponding BIC based Bayes factor output  $P_{BIC}(D|H0)$ . In particular, the spearman rank correlation coefficient  $cor_{spearman,RCT}(p_{value},P_{BIC}(D|H0)) \approx 1$  and a strong contour line relationship dependent on sample size exists between the two statistics that allows pseudo ROC curve interpretation and cutoff estimation. In practice, as P(H0) varies  $(0 \leq P(H0) \leq 1)$  the position of the distribution of observed data on a given sample size contour line varies according to the statistical power (ie. effect size) and P(H0) probability. Proceeding to posterior odds calculations and derivation of  $P_{BIC}(H0|D)$ ,  $P_{BIC}(H1|D)$  probabilities, the above BIC based results have significant relevance under the common prior odds assumption  $P(H0)/P(H1) \sim 1$ .

#### Introduction

There is continuing disagreement about the correlation between

- p<sub>value</sub> the two sample t-test probability of the observed data given the Null Hypothesis and
- P(H0|D) the Bayesian posterior probability of the Null Hypothesis given the observed data.

In this paper, the correlation under repeated sampling for fixed sample sizes is re-examined empirically between Bayes Factor estimates and p-values using (i) BIC based estimates for Bayes Factor outputs  $P_{BIC}(D|H0)$  in relative posterior probability calculations, and explicitly (ii) the Spearman Rank correlation calculation (to allow for a non-linear relationship).

Firstly, the evidence for [1] and against [2], a weak (0.397) correlation coefficient between  $p_{value}$  and P(H0|D) is briefly described. Then an existing BIC based method for Bayes Factor calculation is described and used to produce evidence for a strong non-linear relationship between  $P_{BIC}(D|H0)$  and two sample t-test p-values under repeated sampling for a fixed sample size. Finally, estimates of pseudo ROC curve cutoff points are attempted for the  $P_{BIC}(D|H0)$  (and normalised  $P_{BIC}(D|H0)$ ) contour line for fixed sample sizes to provoke discussion about optimality or otherwise of  $\alpha = 0.05$   $p_{values}$ .

Under the commonly used prior odds assumption  $\frac{P(H0)}{P(H1)} \approx 1$ , the above finding has significant relevance to the posterior probability P(H0|D), p-value correlation for fixed sample sizes.

Previously, Trafimow and Rice [1] raised the issue that frequentist based Null Hypothesis Significance Test (NHST) p-values are not strongly correlated (ie. 0.397) with Bayesian estimates of P(H0|D) based on Bayes formula updating. The simulation used randomly drawn values of  $P(D|H0) \rightarrow p_{value}$ , P(H0), and P(D|H1) as input to Bayes Formula calculations.

As a critique of the relevance of the evidence presented in [1], Lakens [2]

1. Replicated (with more accuracy) the results from [1], which Lakens identified as employing Pearson correlation calculations (ie. 0.37) of the relationship between p-values and posterior P(H0|D) estimates using Bayes Rule updating under randomly drawn values of P(D|H0), P(H0), and P(D|H1).

- 2. Then argued that such an approach did not replicate the situation in Randomised Control Trial (RCT) analysis where the statistical power P(D|H1) for a known H1 value is determined by the sample size and coefficient of variation of the data distribution. This pre-determined power along with the existing P(H0) then strengthens the p-value (P(D|H0)) and P(H0|D) relationship under repeated sampling rather than the simulation studied in [1].
- 3. Used simulations to empirically demonstrate the strong non-linear relationship under repeated sampling for fixed P(H0|D) and P(D|H1) for several values of these parameters. Further, Lakens then argued that the results [1] may be inadequate because it assumes a linear regression relationship (ie. Pearson correlation) for the correlation analysis.

# Deriving linked $P_{BIC}(D|H0)$ values and p-values under repeated sampling for fixed sample sizes

Masson [3] demonstrated how to calculate estimates of relative posterior probabilities using

$$\frac{P(H0|D)}{P(H1|D)} = \frac{P(D|H0)}{P(D|H1)} \cdot \frac{P(H0)}{P(H1)}$$
(1)

$$= BF \cdot \frac{P(H0)}{P(H1)} \tag{2}$$

where BF is the Bayes Factor of the ratio of the likelihoods of the data given H0 and H1 respectively, by using Bayesian Information Criteria (BIC) model fit calculations

$$BIC = -2ln(L) + kln(n) \tag{3}$$

where L is the maximum likelihood of the fitted model, k is the number of free model parameters and n is the sample size.

Explicitly, the BIC based estimate [3] for the Bayes Factor component in (2), is given by

$$\frac{P(H0|D)}{P(H1|D)} \approx \frac{P_{BIC}(D|H0)}{P_{BIC}(D|H1)} \cdot \frac{P(H0)}{P(H1)} \tag{4}$$

$$\approx e^{\frac{(\Delta BIC)}{2}} \cdot \frac{P(H0)}{P(H1)} \tag{5}$$

$$\rightarrow e^{\frac{(\Delta BIC)}{2}}$$
 as  $\frac{P(H0)}{P(H1)} \rightarrow 1$  (6)

where

$$\Delta BIC = BIC_{H1} - BIC_{H0} \tag{7}$$

In comparison, the  $p_{value}$  of the frequentist based NHST method is calculated via the probability of the observed t statistic under a two-sided null hypothesis

$$p_{value} = P(|t||H_{null}, D)$$
(8)

where  $H_{null}: \mu_T = \mu_C ; H_{alt}: \mu_T \neq \mu_C$ 

In practice, in the calculations presented in the paper, the two model populations used were (i) an intercept only model and (ii) an intercept with additive treatment effect model

$$H0: \mu_T = \mu_C \tag{9}$$

$$H1: \mu_T = \mu_C + \delta \tag{10}$$

To obtain  $P_{BIC}(D|H0)$ , as explained in [3] the relationship

$$P_{BIC}(D|H0) = \frac{BF}{(BF+1)} \tag{11}$$

is used and the same (r package) lm r-object output was explicitly employed for  $p_{value}$  and BF (BIC based Bayes Factor) calculations using summary(lm) and BIC(lm) commands respectively.

Providing theoretical support for an information equivalence between  $p_{value}$  and BIC based Bayes Factor statistics Francis [4] describes the mapping between various summary statistics for two sample t-tests. In particular, for independent samples the t statistic estimate is related to the difference in maximum likelihoods of the two model fits [4] by the expression

$$t = \sqrt{(n-2)\left[\exp\left(\frac{2(\ln(L_{H1}) - \ln(L_{H0}))}{n}\right) - 1\right]}$$
 (12)

where the  $p_{value}$  can then be obtained from the t-statistic via equation (8) and  $\Delta BIC$  obtained from  $(ln(L_{H1}) - ln(L_{H0}))$  via equations (3) and (7).

Empirical behaviour of  $P_{BIC}(D|H0)$  vs  $p_{value}$  under repeated sampling for different fixed sample sizes

Under A/A conditions ie. P(H1) = 0

## P<sub>BIC</sub>( D | H0 ) vs p\_value for different total N in RCT analysis

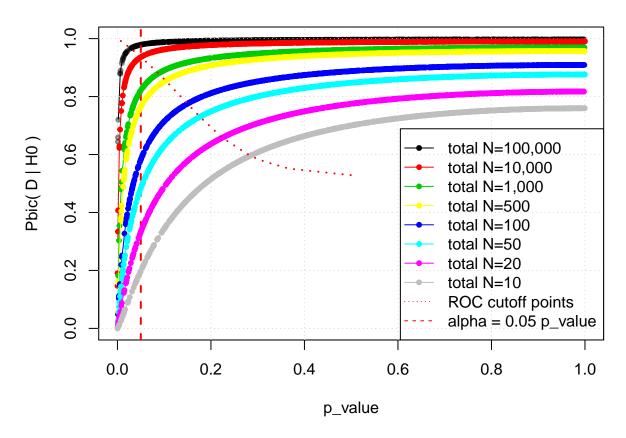


Figure 1.  $P_{BIC}(D|H0)$  as a function of p value for different sample sizes under simple RCT analysis when P(H0) = 1.

#### 0.8 Pbic(D|H0) 9.0 total N=100,000 total N=10,000 total N=1,000 0.4 total N=500 total N=100 total N=50 0.2 total N=20 total N=10 **ROC** cutoff points 0.0 alpha = 0.05 p\_value 0.0 0.2 0.4 0.6 8.0 1.0 p\_value

## P<sub>BIC</sub>( D | H0 ) vs p\_value for different total N in RCT analysis

Figure 2.  $P_{BIC}(D|H0)$  as a function of p value for different sample sizes under simple RCT analysis when P(H0)=1. The contour line barely changes when P(H0)<1 as  $cor_{spearman,RCT}(p_{value},P_{BIC}(D|H0))\approx 1$  remains but the distribution of observed p values and  $P_{BIC}(D|H0)$  values moves to the left. As evidence, of fixed contour line behaviour the fitted lines on the graph are from P(H1)=0 data and these lines fit the P(H1)=1 data points very well.

Looking at Figure 1, with P(H1) = 0 under repeated sampling for fixed sample size there is a strong non-linear contour line behaviour between  $P_{BIC}(D|H0)$  and  $p_{value}$ . The curvature of the contour line depends on the fixed sample size. As the the sample size increases, the density of the observed data pairs of  $(P_{BIC}(D|H0), p_{value})$  with  $P_{BIC}(D|H0) < 0.8$  in the region  $0 \le p_{value} \le 0.05$  decreases.

Using the Spearman Rank correlation calculation  $cor_{spearman,RCT}(p_{value}, P_{BIC}(D|H0)) \approx 1$  which is consistent with a strong non-linear relationship between  $(P_{BIC}(D|H0), p_{value})$  observed data in simple RCT analysis when P(H0) = 1.

Looking at Figure 2, with P(H1) = 1 and using the fitted contour lines from P(H1) = 0 data from figure 1, the data using P(H1) = 1 lies closely on the same contour line for a fixed sample size but the distribution of the observed data shifts (left and down) to lower  $P_{BIC}(D|H0)$  and  $p_{values}$  consistent with the measured treatment effect results being determined as statistically significant at a higher rate.

Again using the Spearman Rank correlation calculation  $cor_{spearman,RCT}(p_{value},P_{BIC}(D|H0)) \approx 1$  which is consistent with a strong non-linear relationship between  $(P_{BIC}(D|H0),p_{value})$  observed data in simple RCT

analysis when P(H0) < 1. As the treatment effect increases further, more observed data points will shift more dramatically (left and down) to lower  $P_{BIC}(D|H0)$  and  $p_{values}$  consistent with increasing statistical power.

# ROC curve interpretation and analysis of $P_{BIC}(D|H0)$ vs $p_{value}$ behaviour for different fixed sample sizes

It can also be observed from figures 1 & 2 that the  $P_{BIC}(D|H0)$  vs  $p_{value}$  behaviour for fixed sample sizes has the form of a Receiver Operating Characteristic (ROC) curve which appears often for binary classification problems. In this case the binary classification objective is assigning statistical significance labels to particular observed  $P_{BIC}(D|H0)$  and/or  $p_{value}$  estimates.

Using the approximation equation (6), such data analysis would also be used to estimate statistical significance for  $\hat{P}(H0|D)$ .

Using the ROC curve interpretation, the optimal threshold for  $P_{BIC}(D|H0)$  and by extrapolation  $\hat{P}(H0|D)$  using the BIC based relative posterior probability equations (5) - (6) could be calculated by the elbow point of the ROC curve behaviour. Two common methods are the Youden Index and the maximum height from the diagonal (between minimum and maximum values). In this paper, both methods gives very similar results for the elbow point of  $P_{BIC}(D|H0)$  vs  $p_{value}$  behaviour.

The elbow point cutoff  $p_{value}$  behaviour as a function of sample size model analysis is shown in figure 3 as a log-log plot. The cutoff points are shown in figures 1 & 2 at the intersection of the contour lines and the red dotted line. The alternate cutoff points using the standard  $\alpha = .05$  NHST method is shown in figures 1 & 2 at the intersection of the contour lines and the vertical red dashed line. For total sample size of 10,000, the optimal ROC cutoff point is  $p_{value} \sim 0.05$  agreeing closely with the standard NHST method.

One immediate comment about the ROC curve cutoff point for N > 10,000 becoming a  $p_{value} < .05$  is that [5,6] Bayesian arguments have already been made that the NHST approach ( $\alpha = 0.05$ ) seems too high for large samples based on increasing Bayesian probabilities of false positives and "it must be the case that a large enough sample will produce a significant result" [5]. A possible reply to that argument based on figure 1 data, is that the contour lines for N > 10,000 arises from P(H0) = 1.

Two resolutions to the above may be: Firstly, the proper application of the observed data for posterior probability calculations rather than just using the Bayes Factor approach would therefore be that the prior odds  $\frac{P(H0)}{P(H1)}$  be included rather than the common usage of  $\frac{P(H0)}{P(H1)} \approx 1$ . However, using P(H1) = 0 is problematic for prior odds estimation. Secondly, the BIC based Bayes Factor approach [3] is an approximation and may not be sensitive enough for the  $P(H0) \to 1$  case.

A second comment is that the standard  $\alpha=0.05$  approach can be interpreted as essentially assigning a fixed cutoff threshold to the ROC curve. In comparison to the elbow cutoff point behaviour, using  $\alpha=0.05$  is aggressive for sample sizes < 10,000. In the last section, the data distribution for different sample sizes under both  $\alpha=0.05$  and 95% statistical power constraints is illustrated.

In the appendix, the relationship between the BIC based Bayes Factor itself  $BF_{BIC} = \frac{P_{BIC}(D|H0)}{P_{BIC}(D|H1)}$  and  $p_{value}$  is illustrated and for  $p_{value} = 0.05$  the approximate asymptotic limit for the BIC based Bayes Factor for simple RCT analysis is  $\frac{BF_{BIC}}{\sqrt{N}} \sim 0.15 = (1/6.666)$  as N  $\rightarrow \infty$ .

#### 1/N vs $p_{value}$ for $P_{BIC}(D \mid H0)$ vs $p_{value}$ ROC curve cutoff points

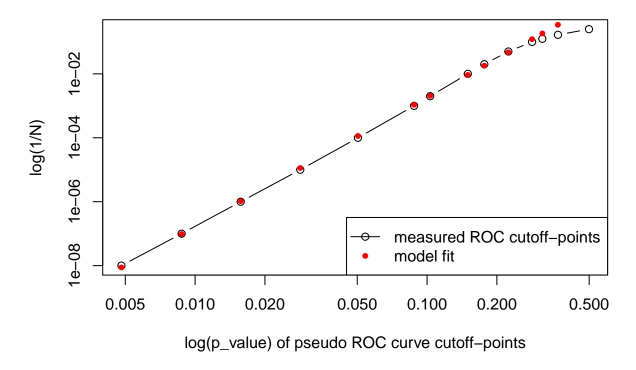


Figure 3. Linear regression fitted log-log scatterplot of 1/(N) vs  $p_{value}$  for ROC curve cutoff point analysis.

#### Normalised $P_{BIC}(D|H0)$ values

It can be seen in figure 1 that  $P_{BIC}(D|H0) < 1$  when P(H0) = 1. For small sample the evidence for the dominance of H0 or H1 is weaker (consistent with low statistical power) and  $P_{BIC}(D|H0) \ll 1$ . However, as discussed for Figure 2 the contour line for fixed sample size doesn't move with treatment effect, only the observed data shifts (left and down) along the contour line as statistical power increases.

To improve interpretation of  $P_{BIC}(D|H0)$  with respect to P(H0), a simple normalisation factor by inspection can be added to equation (11)

$$norm P_{BIC}(D|H0) = \frac{BF}{(BF+1)} \cdot \frac{1}{(1+\frac{1}{\sqrt{N}})}$$
 (13)

such that  $normP_{BIC}(D|H0)$  has the interval (0,1].

The behaviour of  $normP_{BIC}(D|H0)$  is shown in Figure 4 and the same ROC cutoff  $p_{value}$  apply as for  $P_{BIC}(D|H0)$ . To calculate the ROC cutoff point an additional method now becomes available  $\frac{dnormP_{BIC}(D|H0)}{dp} = 1$  and is in close agreement with the Youden Index and maximum height from the diagonal (which now has fixed minimum=0 and maximum=1 for  $normP_{BIC}(D|H0)$  irrespective of sample size). As  $N \to \infty$  the slope of the ROC cutoff line approaches -1.

## norm\_P<sub>BIC</sub>( D | H0 ) vs p\_value for different total N in RCT analysis

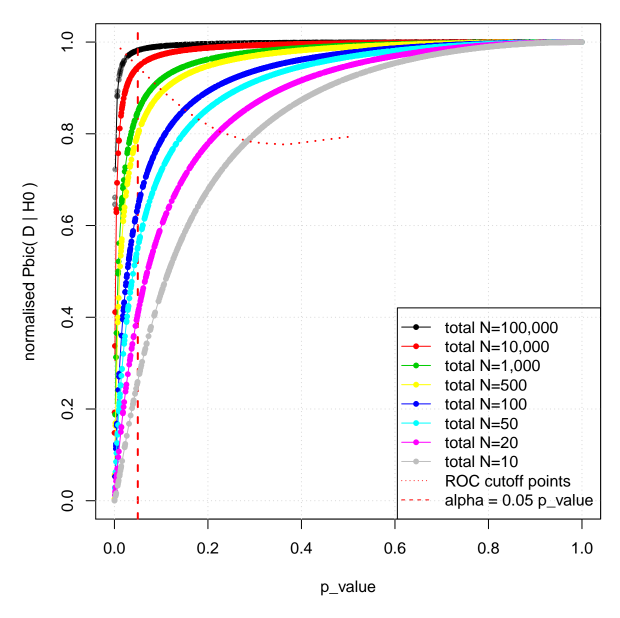


Figure 4.  $normP_{BIC}(D|H0) \equiv \frac{P_{BIC}(D|H0)}{(1+\frac{1}{\sqrt{N}})}$  as a function of p value for different sample sizes under simple RCT analysis when P(H0) = 1. Using the  $normP_{BIC}(D|H0)$  value, the pseudo ROC cutoff point estimate under Youden Index, elbow point and  $\frac{dnormP_{BIC}(D|H0)}{dp} = 1$  methods are in very close agreement.

 $\alpha=0.05$  and 95% statistical power case contour plots for  $normP_{BIC}(D|H0)$  displaying only 100 repeated samples

To appreciate the benefit of having sufficiently powered sample sizes to cope with expected minimum treatment effect sizes, Figure 5 shows the  $p_{value}$ ,  $normP_{BIC}(D|H0)$  relationship for simple RCT designs meeting both

 $\alpha = 0.05$  and 95% statistical power constraint. For clarity, only 100 repeated samples are shown, so on average only ~5 data points should have  $p_{value} > 0.05$ . The contour lines underneath the data points were generated from figure 1 where P(H0) = 1 again showing the constancy of the contour line positions for a given sample size.

## norm\_P<sub>BIC</sub>( D | H0 ) vs p\_value for 95% statistical power cases

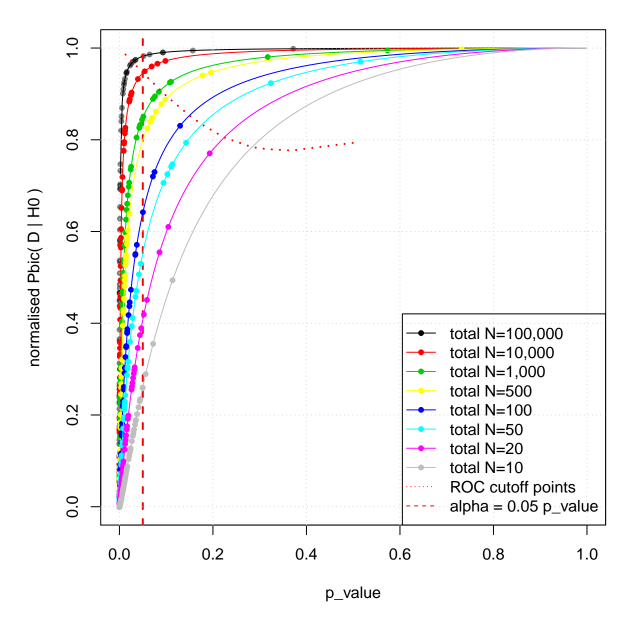


Figure 5.  $P_{BIC}(D|H0)$  as a function of p value for different sample sizes under simple RCT analysis when  $\alpha=0.05$  and the statistical power is ~95%. In this simulation the effect size is different for each fixed sample size to achieve ~95% and for clarity only 100 repeated samples are shown. The underlying contour lines behind the data points were obtained from figure 1 using P(H0)=0.

#### Conclusions

Under repeated sampling, for a fixed sample size the Spearman Rank correlation coefficient  $cor_{spearman,RCT}(p_{value},P_{BIC}(D|H0)) \approx 1$  where  $P_{BIC}(D|H0)$  is derived from BIC based Bayes Factor calculations [3] indicating a strong relationship between the NHST  $p_{value}$  and Bayesian  $P_{BIC}(D|H0)$ . This behaviour is consistent with the earlier work [2,4] also showing a strong relationship between NHST  $p_{value}$  and Bayesian P(D|H0) calculations.

With the evidence of the  $(p_{value}, P_{BIC}(D|H0))$  contour line positions being independent of (i) treatment effect size and (ii) P(H1) the above results gives interesting information on the common interpretation of relative posterior probability based estimates of P(H0|D) equation (2) assuming  $\frac{P(H0)}{P(H1)} \approx constant$ .

The corresponding  $BF_{BIC}=\frac{P_{BIC}(D|H0)}{P_{BIC}(D|H1)}$  and  $p_{value}$  relationship is illustrated in the Appendix where is estimated for  $p_{value}=0.05$  that the approximate large sample asymptotic limit for the BIC based Bayes Factor for simple RCT analysis is  $\frac{BF_{BIC}}{\sqrt{N}}\sim 0.15=(1/6.666)$  as N  $\rightarrow \infty$ 

Finally, a normalised version of  $P_{BIC}(D|H0)$  provides easier ROC point cutoff calculation and interpretation.

#### References

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## Appendix A: BIC based Bayes Factor vs $p_{value}$ behaviour for different sample sizes

The following three graphs show the same RCT analysis results (under repeated sampling for different fixed sample sizes), indicating the relationship between the BIC based Bayes Factor itself  $BF_{BIC} = \frac{P_{BIC}(D|H0)}{P_{BIC}(D|H1)}$  and  $p_{value}$ .

Figure 7 shows the raw scatterplot as a log plot where the different sample size contour lines seem to be at least approximately, shifted versions of each other.

Figure 8 shows the log plot relationship using a normalised BIC based Bayes Factor  $\frac{BF_{BIC}}{\sqrt{N}}$ . It can be seen that as  $p_{value} \to 1$  the normalised contour lines are in good agreement. The upper limit  $\frac{BF_{BIC}}{\sqrt{N}} = 1$  and an approximate large N intersection point of  $\frac{BF_{BIC}}{\sqrt{N}} \approx 0.15$  when  $p_{value} = 0.05$  are indicated by dotted horizontal lines.

Finally, Figure 9 shows the  $\log(\frac{BF_{BIC}}{\sqrt{N}})$ - $\log(p_{value})$  relationship which more accurately estimates that  $\frac{BF_{BIC}}{\sqrt{N}} \sim 0.15 = (1/6.666)$  as N  $\to \infty$ . As N becomes small, the intersection point with  $p_{value} = 0.5$  dcreases to lower  $\frac{BF_{BIC}}{\sqrt{N}}$  values.

## BIC based Bayes Factor vs p\_value for 95% statistical power cases

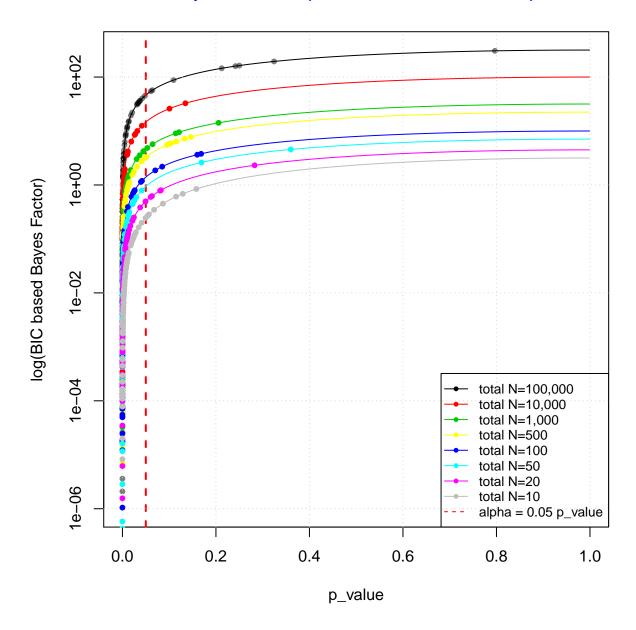


Figure 6.  $log(BIC\ based\ Bayes\ Factor)$  as a function of  $p_{value}$  for different sample sizes under simple RCT analysis when  $\alpha=0.05$  and the statistical power is ~95%, for clarity only 100 repeated samples are shown. The underlying contour lines behind the data points were obtained from figure 1 using P(H0)=0.

## BIC based Bayes Factor vs p\_value for 95% statistical power cases

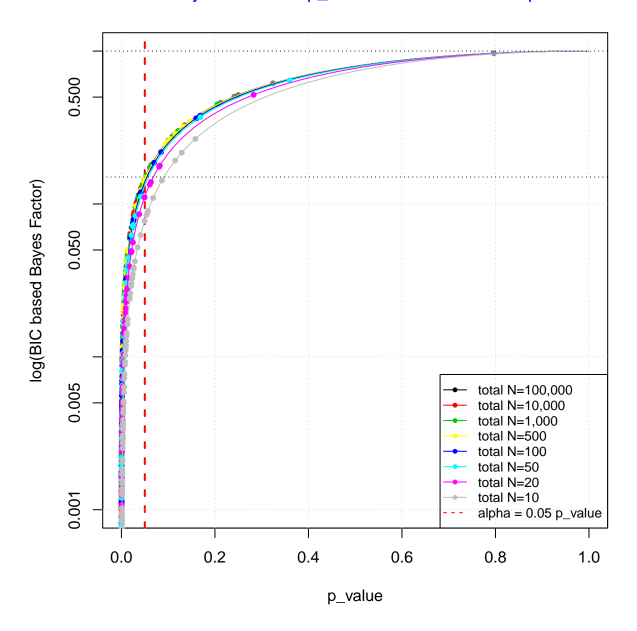
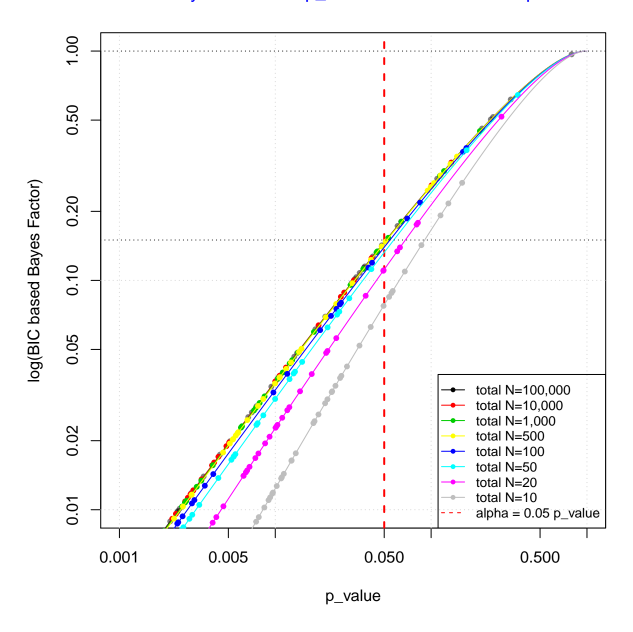


Figure 7.  $log((BIC\ based\ Bayes\ Factor)/sqrt(N))$  as a function of  $p_{value}$  for different sample sizes under simple RCT analysis when  $\alpha=0.05$  and the statistical power is ~95%, for clarity only 100 repeated samples are shown. The underlying contour lines behind the data points were obtained from figure 1 using P(H0)=0.

## BIC based Bayes Factor vs p\_value for 95% statistical power cases



\*\*\*Figure 7.  $\log((\text{BIC based Bayes Factor})/\text{sqrt}(N))$  as a function of  $\log(p_{value})$  for different sample sizes under simple RCT analysis when  $\alpha=0.05$  and the statistical power is ~95%, for clarity only 100 repeated samples are shown. The underlying contour lines behind the data points were obtained from figure 1 using P(H0)=0.