### BIOLOGY LETTERS

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#### Review



**Cite this article:** Halsey LG. 2019 The reign of the *p*-value is over: what alternative analyses could we employ to fill the power vacuum? *Biol. Lett.* **15**: 20190174. http://dx.doi.org/10.1098/rsbl.2019.0174

Received: 4 March 2019 Accepted: 1 May 2019

#### Subject Areas:

bioinformatics

#### **Keywords:**

AIC, Bayesian, confidence intervals, effect size, statistical analysis

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Electronic supplementary material is available online at https://dx.doi.org/10.6084/m9. figshare.c.4498919.



#### Animal behaviour

# The reign of the *p*-value is over: what alternative analyses could we employ to fill the power vacuum?

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The *p*-value has long been the figurehead of statistical analysis in biology, but its position is under threat. p is now widely recognized as providing quite limited information about our data, and as being easily misinterpreted. Many biologists are aware of p's frailties, but less clear about how they might change the way they analyse their data in response. This article highlights and summarizes four broad statistical approaches that augment or replace the *p*-value, and that are relatively straightforward to apply. First, you can augment your *p*-value with information about how confident you are in it, how likely it is that you will get a similar *p*-value in a replicate study, or the probability that a statistically significant finding is in fact a false positive. Second, you can enhance the information provided by frequentist statistics with a focus on effect sizes and a quantified confidence that those effect sizes are accurate. Third, you can augment or substitute *p*-values with the Bayes factor to inform on the relative levels of evidence for the null and alternative hypotheses; this approach is particularly appropriate for studies where you wish to keep collecting data until clear evidence for or against your hypothesis has accrued. Finally, specifically where you are using multiple variables to predict an outcome through model building, Akaike information criteria can take the place of the *p*-value, providing quantified information on what model is best. Hopefully, this quick-and-easy guide to some simple yet powerful statistical options will support biologists in adopting new approaches where they feel that the *p*-value alone is not doing their data justice.

#### 1. Introduction

The reified position of the *p*-value in statistical analyses was unchallenged for decades despite criticism from statisticians and other scientists (e.g. [1-4]). In recent years, however, this unrest has intensified, with a plethora of new papers either driving home previous arguments against *p* or raising additional critiques (e.g. [5-11]). Catalysed by the part that the *p*-value has played in science's reproducibility crisis, this criticism has brought us to the brink of an uprising against *p*'s reign.

Consequently, an analysis power vacuum is forming, with a range of alternative approaches vying to fill the space. Commentaries that criticize the *p*-value often suggest alternative paradigms of statistical analysis, and now a number of options have taken seed in the field of biology. New statistical methods typically involve concepts that are counterintuitive to our *p*-based training; they represent radically different ways of interrogating data that involve disparate approaches to generating evidence, different software packages and a host of new assumptions to understand and justify. The steep curves for learning new methods could stifle further expansion of their use in lieu of *p*-centred statistical analyses in the biological sciences.

To provide clarity and confidence for biologists seeking to expand and diversify their analytical approaches, this article summarizes some tractable alternatives to *p*-value centricity. But first, here is a brief overview about the limits of the *p*-value and why, on its own, it is rarely sufficient to interpret our hard-earned data. Along with many other august statisticians, Jacob Cohen and John Tukey have written cogently about their concerns with the fundamental concept of null hypothesis significance testing. Because the *p*-value is predicated on the null hypothesis being true, it does not give us any information about the alternative hypothesis-the hypothesis we are usually most interested in. Compounding this problem, if our *p*-value is high and so does not reject the null hypothesis this cannot be interpreted as the null being true; rather, we are left with an 'open verdict' [2]. Moreover, with a big enough sample size, inevitably the null hypothesis will be rejected; perversely, a p-value based statistical result is as informative about our sample as it is about our hypothesis [12,13].

Recently, further concerns have been documented about p, linking the p-value to problems with experimental replication [5]. Cumming [7] and Halsey et al. [6] demonstrated that *p* is 'fickle' in that it can vary greatly between replicates even when statistical power is high, and argued that this makes interpretation of the *p*-value untenable unless *p* is extremely small. Colquhoun [8,14] has argued that significant p-values at just below 0.05 are extremely weak evidence against the null hypothesis because there is a 1 in 3 chance that the significant result is a false positive (aka type 1 error). Interpreting *p* dichotomously as 'significant' or 'not significant' is particularly egregious for many reasons, but most pertinent here is that this approach encourages failed experiment replication. Studies are often designed to have 80% statistical power, meaning that there is an 80% chance that an effect in the data will be detected. As Wasserstein & Lazar [9] explain, the probability of two identical studies statistically powered to 80% both returning  $p \le 0.05$  is at best 80%  $\times$ 80% = 64%, while the probability of one of these studies returning  $p \le 0.05$  and the other not is  $2 \times 80\% \times 20\% =$ 32%. Together, these papers and calculations demonstrate that the *p*-value is typically highly imprecise about the amount of evidence against the null hypothesis, and thus pshould be considered as providing only loose, first pass evidence about the phenomenon being studied [6,15,16].

With the broadening realization among biologists that p-values provide only tentative evidence about our dataand, indeed, that exactly what this evidence tells us is easy to misinterpret-it is important that we equip ourselves with a broad understanding of what statistical options are available that can clarify, or even supplant, p. While it will be hard to extricate ourselves from our indoctrinated approach to interpreting every statistical analysis through the prism of significance or non-significance, we can be motivated by the knowledge that there really are other ways, and indeed more intuitive ways, to investigate our data. Below, I provide a quick-and-easy guide to some simple yet powerful statistical options currently available to biologists conducting standard study designs. Each distinct statistical approach interrogates the data through a different lens, i.e. by asking a fundamentally different scientific question; this is reflected in the subsection headings that follow. We shall start with the option least disruptive to the *p*-value paradigm—augmenting *p* with information about its variability.

## 2. *p*-Value: how much evidence is there against the null hypothesis?

*p* provides unintuitive information about your data. However, perhaps it can best be interpreted as characterizing the evidence in the data against the null hypothesis [10,17]. And despite its limitations, the *p*-value has attractive qualities. It is a single number from which an objective interpretation about data can be made. Moreover, arguably that interpretation is context independent; p-values can be compared across different types of studies and statistical tests [18] (though see [10]). Huber [19] argues that focusing on the *p*-value is a suitable first step for screening of multiple hypotheses, as occurs in 'high throughput biology' such as gene expression analysis and genome-wide association studies.

However, p is let down by the considerable variability it exhibits between study samples-variability disguised by the reporting of p as a single value to several decimal places. Arguably, then, if you want to continue calculating *p* as part of your analyses of single tests, you ought to provide some additional information about this variability, to inform the reader about the uncertainty of this statistic. One way to achieve this is to provide a value that is somewhat akin to the confidence interval around an effect size, which characterizes the uncertainty of your study *p*-value and is termed the *p*-value prediction interval [7]. Another option is to calculate the prediction interval that characterizes the uncertainty of the p-value of a future replicate study. Lazzeroni et al. [18] provide a simple online calculator for both [18]. Based on this calculator, if the p-value from your experiment is, for example, 0.01, it will have a 95% prediction interval of  $5.7^{-6}$  to 0.54. Clearly, this would provide us with little confidence that *p* is replicable under this experimental scenario. A *p*-value of 0.0001 has a 95% prediction interval of 0–0.05. In this second scenario, the 95% prediction interval of a future replicate study is 0-0.26. Vsevolozhskaya et al. [20] argue that the prediction interval around p calculated by this method returns underestimates of both the lower and upper bounds. Nonetheless, the width of the prediction interval, however calculated, will be surprisingly large to those of us accustomed to seeing the *p*-value as a naked single value reported to great precision.

If you have calculated the planned power of your study and are prepared to quantify the level of belief you had before conducting the experiment that the null hypothesis is true, you can augment p with the estimated likelihood that if you get a significant p-value it is falsely rejecting the null hypothesis. This is termed the estimated false positive (discovery) risk, and can be easily estimated from a simple Bayesian framework (see later) ([9] and the comment by Altman annexed to [9]):

Estimated false positive risk =  $\frac{p \cdot \pi_0}{p \cdot \pi_0 + (1 - \beta) (1 - \pi_0)}$ 

where *p* is the *p*-value of your study,  $\pi_0$  is the probability that the null hypothesis is true based on prior evidence and  $(1 - \beta)$  is study power.

For example, if you have powered your study to 80% and before you conduct your study you think there is a 30% possibility that your perturbation will have an effect (thus  $\pi_0 =$  0.7), and then having conducted the study your analysis returns p = 0.05, the estimated false positive risk is 13%. That is, many replicates of this experiment would indicate a

statistically significant effect of the perturbation and be wrong in doing so about 13% of the time. Bear in mind, however, that given the aforementioned fickleness of p, this estimate of false positive risk could be equally capricious. This concern can be circumvented for high throughput studies, replacing p in the equation above for  $\alpha$  (the significance threshold of the statistical test), and estimating  $\pi_0$  from observed p-values [9,21].

For those not conducting high throughput studies and who do not like the idea of subjectively quantifying their *a priori* expectations about the veracity of their experimental perturbation, the calculations can be flipped such that your *p*-value is accompanied by a calculation of the prior expectation that would be needed to produce a specified risk (e.g. 5%) of a significant *p*-value being a false positive ([8]; and the author provides an easy-to-use web calculator for this purpose: http://fpr-calc.ucl.ac.uk/). This provides an alternative way of assessing the likelihood that a significant *p*-value is a true positive. If, for example, your *p*-value is 0.03 for a study powered to about 70%, to limit the risk of a false positive to 5% your prior expectation that the perturbation will have an effect would need to be 77% (based on the '*p*-equals' case; [8]).

## 3. Effect size and confidence interval: how much and how accurate?

A statistically significant result tells us relatively little about the phenomenon we are studying—only that the null hypothesis of no 'effect' in our data (which we already knew wasn't true to some level of precision; [13]) has been rejected [22]. Instead of the *p*-value scientific question 'is there or isn't there an effect?', considerably more information is garnered by asking 'how strong is the effect in our sample?' coupled with the question 'how accurate is that value as an estimate of how strong the population effect is?'.

The most straightforward way to analyse your data in order to answer these two questions is to calculate the effect size in the sample along with the 95% confidence intervals around that estimate [6,7,23–26]. Fortunately, the effect size is often easy to calculate or extract from statistical outputs, since it is typically the mean difference between two groups or the strength of the correlation between two variables. And while the definition of a confidence interval is complex, Cumming & Calin-Jageman [27] compellingly argue that it is reasonable to interpret a confidence interval as an indication of the accuracy of the effect size estimate; it is the likely error estimation.

The calculations of confidence intervals and *p*-values share the same mathematical framework [28,29], but this does not detract from the fact that focusing interpretation of data on effect sizes and their confidence intervals is a fundamentally different approach from that of focusing interpretation on whether or not to reject the null hypothesis [11]. These two procedures ask very different questions about the data and elicit distinct answers [30]. For example, a study on the effects of two different ambient temperatures on paramecium diameter returning an effect size of 20 µm and a p-value of 0.1, if centred on p-value interpretation would conclude 'no effect' of temperature, despite the best supported effect size being 20, not 0. An interpretation based on effect size and confidence intervals could, for example, state: 'Our results suggest that paramecium kept at the lower temperature will be on average 20 µm larger in size,

however a difference in size ranging between -4 and 50 µm is also reasonably likely'. As Amrhein *et al.* [11] point out, the latter approach acknowledges the uncertainty in the estimated effect size while also ensuring that you do not make a false claim either of no effect if p > 0.05, or an overly confident claim. And if all the values within the confidence interval are biologically unimportant, then a statement that your results indicate no important effect can also be made [11]. (This is an example of where focusing on effect size and uncertainty also allows clear yes/no interpretations if desired; see also [31].)

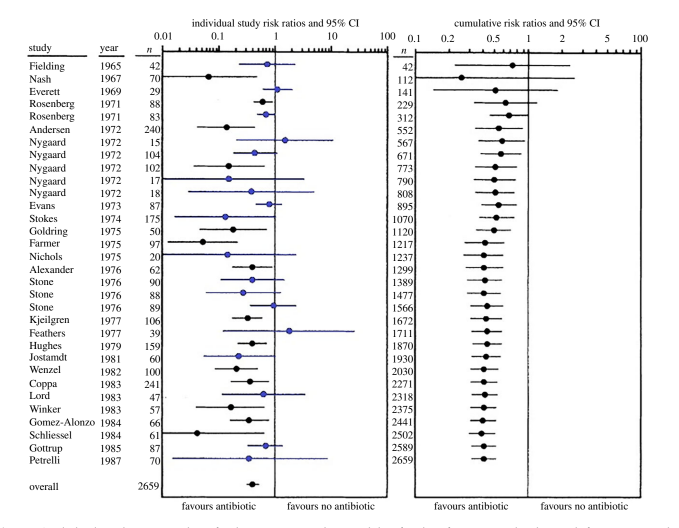
The approach of focusing on effect size estimation is usually accompanied by an emphasis on visualization of the data to support their evaluation. A strong graphical format that achieves this involves a main panel showing the raw data and side panels helping to illustrate the estimated effect size [32]. Refer to the electronic supplementary materials for an example plot (figure S1). Such plots, while intuitive, are not typically available in statistical packages and not easy to code in programming languages. However, Ho and colleagues [32] have recently developed 'Data Analysis with Bootstrap-coupled ESTimation' (DABEST), available in versions for Matlab, Python and R, and also as a webpage https://www.estimationstats.com/#/. All versions have user-friendly, rote instructions to produce graphs that allow full exploration of your data.

Scientific research seeks to home in on 'answers', and estimated effect sizes and their confidence intervals are central to this goal. In biology at least, homing in on an answer almost inevitably requires multiple studies, which then need to be analysed together, through meta-analysis. Effect sizes and confidence intervals are the vital information for this process (e.g. [33]), providing another good argument for their thorough reporting in papers. Typically, the confidence intervals around an effect size calculated from a meta-analysis are much smaller than those of the individual studies [34], thus giving a much clearer picture about the true, populationlevel effect size (figure 1). However, meta-analyses can be deeply compromised by the 'file drawer phenomenon', where non-significant results are not published [36], either because researchers do not submit them, or journals will not accept them [37]. Fortunately, attitudes of science funders, publishers and researchers are starting to change about the value and importance of reporting non-significant results; this momentum needs to continue.

## 4. Bayes factor: what is the evidence for one hypothesis compared to another?

In contrast to the *p*-value providing only information about the likelihood that the null hypothesis is true, the Bayes factor directly addresses both the null and the alternative hypotheses. The Bayes factor quantifies the relative evidence in the data you have collected about whether those data are better predicted by the null hypothesis or the alternative hypothesis (an effect of stated magnitude). For example, a Bayes factor of 5 indicates that the strength of evidence is five times greater for the alternative hypothesis than the null hypothesis; a Bayes factor of 1/5 indicates the reverse.

The Bayes factor is a simple and intuitive way of undertaking the Bayesian version of null hypothesis significance testing. Only recently have Bayes factors been made tractable for the practising biologist, and these are now easily calculable



**Figure 1.** Standard and cumulative meta-analyses of studies investigating antibiotic prophylaxis for colon infection compared to the control of no treatment. In the left panel, the effect size and 95% confidence interval are shown for each study, which are displayed chronologically. Risk ratios (effect size) less than 1 favour a prophylactic; greater than 1 favours no treatment. *n* represents study sample size. The pooled result from all studies is shown at the bottom. Note that the studies where the confidence interval intersects 1 (coloured blue) would be interpreted as statistically non-significant (no efficacy of the prophylaxis); otherwise (black) as statistically significant (the prophylaxis is worth administering). Interpretation of all these studies based on the *p*-value alone would not provide any clarification about the value of an antibiotic prophylaxis with treatment of colon infection, with around half the studies reporting statistical significance. The right panel represents a cumulative meta-analysis of the same studies (*n* represents cumulative sample size). This shows that some degree of efficacy of antibiotic prophylaxis for treatment in colon infection could have been identified as early as 1972, and the efficacy effect size was fairly clear well before the final study. Figure (adapted) and some caption text taken from loannidis & Lau [35].

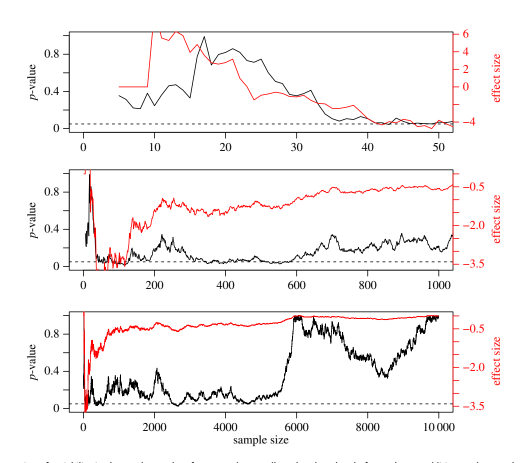
for a range of standard study designs. The Bayes factors for many designs can be run on web-based calculators (e.g. http://pcl.missouri.edu/bayesfactor) and are also available as a new package for R called BayesFactor() [38].

A controversy of the Bayesian approach is the need for you to specify your strength of belief in the effect being studied before the experiment takes place (the prior distribution of the alternative hypothesis) [39]. Thus, your somewhat subjective choice of 'prior' influences the outcome of the analysis. Schönbrodt *et al.* [40] argue that this criticism of Bayesian statistics is often exaggerated because the influence of the prior is limited when a reasonable prior distribution is used. You can assess the influence of the prior with a simple sensitivity analysis whereby the analysis is run using a bounded range of realistic prior probabilities [41]. There is also a default prior that you can use in the common situation when you have little pre-study evidence for the expected effect size.

Nonetheless, undertaking Bayesian analyses is more involved than null hypothesis significance testing, and specifying the prior undoubtedly adds some degree of subjectivity. Fortunately, there is a single, simple formula that you can apply to convert a *p*-value to a form of the Bayes factor without any other information. This simplified Bayes factor, termed the upper bound, states the most likely it is that the alternative hypothesis is true rather than the null hypothesis over any reasonable prior distribution (comment by Benjamin and Berger annexed to [9] and Goodman [42]):

Bayes factor upper bound  $\leq \frac{-1}{e.p.\ln(p)}$ .

For example, if your data generate a *p*-value of 0.07 (sometimes termed a 'trend'), the Bayes factor upper bound is 1.98 and you can conclude that the alternative hypothesis is at most twice as likely as the null hypothesis. A *p*-value of 0.01 indicates the alternative hypothesis is at most 8 times as likely as the null. Benjamin and Berger argue that this approach is an easily-interpretable alternative to *p*, which should satisfy both practitioners of Bayesian statistics



**Figure 2.** A demonstration of variability in the *p*-value as data from a study are collected and analysed after each new addition to the sample. This can result in a study being stopped under the mistaken belief that as soon as a significant *p*-value is obtained, this reflects a real effect. A computer simulates samples drawn at random from two identical, randomly distributed populations (standard deviation = 10), thus the null hypothesis is true. A Student's *t*-test is conducted after five samples are drawn from the two populations. Subsequently, each time one further sample is taken for each population the *t*-test is re-run. The evolution of the *p*-value as sample size increases is presented in the three panels (black line), the upper panel showing the first 50 samples, the middle the first 1000, and the lower panel showing up to 10 000 samples being drawn. The *p*-value varies considerably; another demonstration of its 'fickleness' [6]. In each panel, the red line represents the effect size (mean difference between the samples). Although the *p*-value should typically be high under these circumstances, reflecting a lack of evidence against the null, when the sample size is small it can easily decrease temporarily to below 0.05 (denoted by the dashed line), suggesting that the populations from which the samples are drawn are different. If the sampling is stopped when this happens, *p* will be unrepresentative of reality and return a false positive. (Note that in this simulation, *p* does not tend towards 0 as the sample size becomes very large because as sample size increases the effect size tends towards 0 and thus statistical power does not systematically increase (observed power is inversely related to *p*; [44])).

and practitioners of null hypothesis significance testing (comment by Benjamin and Berger annexed to [9]).

Schönbrodt et al. [40] make the case that the Bayes factor can be used to inform when a study has secured a sufficient sample size and can be halted. Effective stopping rules in research can be invaluable for controlling time and financial costs while increasing study replicability, and are ethically important for certain animal studies or intrusive human studies; the use of subjects should be minimized while ensuring that the experiments are robust and reproducible (https://www.nc3rs.org.uk/the-3rs; [43]). Arguably, stopping rules should be used a lot more than they currently are, and can be a far more effective method for targeting a suitable sample size than power analysis. A big mistake often made, however, is to implement the *p*-value in the stopping rule; the study is stopped when the data thus far collected return a statistically significant *p*-value. The underlying assumption is that increasing the sample size further would probably decrease p further. A simple model demonstrates this thinking to be spurious and thus that it drives very bad practice (figure 2). For those of us basing our study on the *p*-value, it is far preferable to continue a study until a pre-determined sample size is reached that has been decided by *a priori* power analysis [45]. However, this approach is greatly influenced by the associated *a priori* effect size estimate we have provided and there can be a strong temptation to increase sample size beyond the pre-determined number; researchers longing for a statistically significant result can easily succumb to the temptation of collecting extra data points when their *p*-value is 0.06 or 0.07 [46].

The Bayes factor is much more appropriate here. It provides evidence for the null, and with a large enough sample the Bayes Factor will converge on 0 (the null is true) or infinity (the alternative is true). If the Bayes Factor of your data reaches 10 or 1/10, this almost certainly represents the true situation and your study can stop. Alternatively, if your study must be stopped for logistical reasons then the final Bayes Factor can still be interpreted, for example a Bayes factor of 1/7 would indicate moderate evidence for the null hypothesis. Moreover, you are entitled to continue sampling if you feel the data are not conclusive enough; if the results are unclear, collect more data. All such decisions do not affect interpretation of the Bayes Factor [40]. A final big motivation for employing the Bayes factor over the *p*-value in stopping procedures is that in the long run, the former uses a smaller sample while at the

same time generating fewer interpretation errors. A general consensus has not yet been reached about the most suitable priors for each situation, and tractable Bayes factor procedures have thus far only been produced for some experimental designs. But do not let this put you off. Instead of the Bayes factor, the Bayes factor upper bound, as described above, can be used.

## 5. Akaike information criterion: what is the best understanding of the phenomenon being studied?

If your study involves measuring an outcome variable and multiple potential explanatory variables, then you have many possible models you could build to explain the variance in your data. Stepwise procedures of model building often focus on *p*-values, by holding onto only those explanatory variables associated with a low *p*. Aside from the general concerns about p, specific criticisms of p-value-based model building include the inflated risk of type 1 errors [47,48]. An alternative approach to model assessment is the Akaike information criterion (AIC), which can be easily calculated in statistical software packages, and in R using AIC() [49]. The AIC provides you with an estimate of how close your model is to representing full reality [50], or in other words its predictive accuracy [51]. Couched within the principle of simplicity and parsimony, a fundamental aspect of the AIC is that it trades off a model's goodness of fit against that model's complexity to insure against over-fitting [52].

Let's imagine you have generated three models, returning AICs of 443 (model 1), 445 (model 2) and 448 (model 3). Your preferred model in terms of relative quality will be the one that returns the minimum AIC. But you should not necessarily discard the other models. With the AIC calculated for multiple models, you can easily compute the relative likelihood that each of those models is the best of all presented models given your data, i.e. the relative evidence for each of them. For example, the preferred model will always have a relative evidence of 1, and in the current example the second best model, model 2, has relative evidence 0.37, and model 3 has 0.08. Finally, you can then compute an evidence ratio between any pair of models; following the above example, the evidence for model 1 over model 2 is 1/ 0.37 = 2.7, i.e. the evidence for model 1 is 2.7-times as strong. In this scenario, although model 1 has the absolute lowest AIC, the evidence that model 1 rather than model 2 is the best from those generated is not strong, and with some explanatory variables present in only one of the models, the most suitable response could be to make your inferences based on both models [50]. The AIC approach encourages you to think hard about alternative models and thus hypotheses, in contrast to *p*-value interpretation that encourages rejecting the null when p is small, and supporting the alternative hypothesis by default [53]. More broadly, the AIC paradigm involves dropping hypotheses judged implausible, refining remaining hypotheses and adding new hypotheses-a scientific strategy that Burnham et al. [50] argue promotes fast and deep learning about the phenomenon being studied.

Although the AIC is mathematically related to the *p*-value (they are different transformations of the likelihood ratio;

[29]), the former is far more flexible in the models it can compare. The AIC is a strong option for choosing between multiple models that you have generated to explain your data, i.e. to choose what model represents your best understanding of the phenomenon you have measured, particularly when the observed data are complex and poorly understood and you do not expect your models to have particularly strong predictive power [54].

A key limitation of the AIC is that it provides a relative, not absolute, test of model quality. It is easy to fall into the trap of assuming that the best model is also a good model for your data; this may be the case, or instead the best model may have only half an eye on the variance in your data while all other models are blind to it. Quantifying the absolute quality of your best model(s) requires calculation of the effect size, as discussed earlier (in the case of models, typically  $R^2$  is suitable).

#### 6. Conclusion

Good science generates robust data ripe for interpretation. There are several broad approaches to the statistical analysis of data, each interrogating the collected variables through a distinct line of questioning. Popper [55] argued that science is defined by the falsifying of its theories. Taking this approach to science, p-values might be the rightful centrepiece of your statistical analysis since they provide evidence against the null hypothesis [10,17]. Building on this paradigm, you can easily enhance interpretation of the *p*-value by augmenting p with a prediction interval and/or an estimate of the false positive risk—information about p's reliability. A counter argument, however, is that because the *p*-value does not test the null hypothesis nor the alternative hypothesis you can never use it to actually falsify a theory [56]. Converting the *p*-value into a Bayes factor attends to this concern, providing relative evidence for one hypothesis or the other. But many have argued that hypothesis testing by any approach is superseded by focusing on the effect in the data-specifically both its magnitude and accuracy-because your best estimate of the magnitude of the phenomenon you are studying is ultimately what you want to know. And if you conduct multi-variate analysis, particularly when the phenomenon under study is poorly understood, you can be well served by the AIC, which encourages consideration of multiple hypotheses and their gradual refinement.

It is important to emphasize that these manifold approaches are not all mutually exclusive; for example, many would argue that effect size estimates are an essential component of most analyses. Indeed, Goodman et al. [57] go so far as to recommend the use of a hybrid for decision making that requires a low *p*value coupled with an effect size above an *a priori* determined minimum to be relevant/important in order to reject the null hypothesis. p-values can also be presented alongside Bayes factors for each statistical test conducted ('a B for every p'). Continuing to present *p*-values as part of your statistical output while diluting their interpretive power by including other statistical approaches should ensure your submission is not jeopardized, and indeed this approach is probably the best way to nudge reviewers and editors towards acceptingeven encouraging-the application of alternative inferential paradigms (and see Box 2 in [43]). Whatever your chosen statistical approach, it is important that this has been determined before data collection. Arming oneself with more statistical options could risk the temptation of trying different approaches until an exciting result is achieved; this must be resisted.

Regardless of the statistical paradigm you employ to investigate patterns in your data, many have recommended that the outputs from statistical tests should always be considered as secondary interrogations. Primarily, the argument goes, you should prioritize interpretation of graphical plots of your data, where possible, and treat statistical analyses as supporting or confirmatory information [25,58–60]. A plot that does not appear to support the findings of your statistical analysis should not be automatically explained away as a demonstration that your analysis has uncovered patterns that are deeper than can be visualized.

Finally, while I hope that this review might help readers feel a little more informed, and confident, about some of the additional and alternative statistical options to the *p*-value, it is worth reminding ourselves of Sir Ronald Fisher's pertinent words from his Presidential Address to the First Indian Statistical Congress in 1938 [61]: 'To call in a statistician after the experiment is done may be no more than asking him to perform a post-mortem examination: he may be able to say what the experiment died of.' Without a good dataset, none of the statistical tools mentioned here will be effective. Moreover, even a good dataset represents just a single study, and it must not be forgotten that a single study provides limited information. Ultimately, replication is key to refining, and having confidence in, our understanding of the biological world.

Ethics. Consent was not required for this review.

Data accessibility. All data were generated by R code; the code will be made available on request.

Competing interests. I have no competing interests.

Funding. This study was not supported by funding.

Acknowledgements. The Laboratory Animal Care seminar in 2019 entitled '3R seminar: Study design', organised by the University of Oulu Graduate School, provided the catalyst for writing this article. I appreciate the feedback that I received on drafts of this article from Michael Pedersen, Dr Louise Soanes and Dr Mircea Iliescu, and Professor Stuart Semple.

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