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# Distributions of *p*-values smaller than .05 in Psychology: What is going on?

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Previous studies provided mixed findings on pecularities in *p*-value distributions in psychology. This paper examined 258,050 test results across 30,710 articles from eight high impact journals to investigate the existence of a peculiar prevalence of *p*-values just below .05 in the psychological literature, and a potential increase thereof over time. We indeed found evidence for a bump just below .05 in the distribution of exactly reported *p*-values in the journals Developmental Psychology, Journal of Applied Psychology, and Journal of Personality and Social Psychology, but the bump did not increase over the years and disappeared when using recalculated *p*-values. We found clear and direct evidence for the QRP "incorrect rounding of *p*-value" (John et al., 2012) in all psychology journals. Finally, we also investigated monotonic excess of *p*-values, an effect of certain QRPs that has been neglected in previous research, and developed two measures to detect this by modeling the distributions of statistically significant *p*-values. Using simulations and applying the two measures to the retrieved test results, we argue that, although one of the measures suggests the use of QRPs in psychology, it is difficult to draw general conclusions concerning QRPs based on modeling of *p*-value distributions.

# Distributions of *p*-values smaller than .05 in Psychology: What is going on?

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# 7 ABSTRACT

Previous studies provided mixed findings on pecularities in *p*-value distributions in psychology. This paper examined 258,050 test results across 30,710 articles from eight high impact journals to investigate the existence of a peculiar prevalence of *p*-values just below .05 in the psychological literature, and a potential increase thereof over time. We indeed found evidence for a bump just below .05 in the distribution of exactly reported *p*-values in the journals Developmental Psychology, Journal of Applied Psychology, and Journal of Personality and Social Psychology, but the bump did not increase over

- the years and disappeared when using recalculated *p*-values. We found clear and direct evidence for the QRP "incorrect rounding of *p*-value" (John et al., 2012) in all psychology journals. Finally, we also investigated monotonic excess of *p*-values, an effect of certain QRPs that has been neglected in previous research, and developed two measures to detect this by modeling the distributions of statistically significant *p*-values. Using simulations and applying the two measures to the retrieved test results, we argue that, although one of the measures suggests the use of QRPs in psychology, it is difficult to draw general conclusions concerning QRPs based on modeling of *p*-value distributions.
- <sup>9</sup> Keywords: p-values, NHST, QRP, data peeking, Caliper test

# 10 INTRODUCTION

A set of *p*-values can be informative of the underlying effects that are investigated, but can also be 11 indicative of potential research biases or questionable research practices (QRPs). Masicampo and 12 Lalande (2012) found a bump of p-values just below .05 in three main psychology journals (i.e., Journal 13 of Personality and Social Psychology, JPSP; Journal of Experimental Psychology: General, JEPG; 14 *Psychological Science*, PS), which could be explained by research biases. A bump has occurred when 15 *p*-values just below .05 are more prevalent than smaller *p*-values. The observation of a bump was one 16 of several signals of a crisis of confidence in research findings in psychological science (Pashler and 17 Wagenmakers, 2012; Ferguson, 2015). Leggett et al. (2013) later corroborated this bump of p-values for 18 JPSP and JEPG, and observed that it was larger in 2005 than in 1965. Considering that research biases 19 can lead to overemphasis on statistical significance, this result suggested that the state of psychology may 20 have even deteriorated over the years. Additional corroboration in samples of published articles from 21 various fields was provided by Head et al. (2015), who documented the bump of p-values below .05 in 22 1,048,575 articles across 16 disciplines including psychology. Ginsel et al. (2015) found similar biased 23 reporting of *p*-values in medical abstracts, but noted the variety of potential causes (e.g., publication bias, 24 25 fraud, selective reporting). At the same time, other studies failed to find a bump of *p*-values below .05 (Jager and Leek, 2014; 26 Krawczyk, 2015; Vermeulen et al., 2015). Reanalysis of data from Masicampo and Lalande (2012) and 27 Head et al. (2015) indicated that the original results may have been confounded by publication bias 28 and tendencies to round p-values (Lakens, 2015b; Hartgerink, 2015). Publication bias refers to the fact 29 that the probability of getting published is higher for statistically significant results than for statistically 30 nonsignificant results (Gerber et al., 2010; Franco et al., 2014). Publication bias only changes the p-value 31 distribution above .05 and cannot cause a bump. Krawczyk (2015) analyzed a sample of around 5,000 32 psychology articles and found no bump in *p*-values that were *recalculated* on the basis of reported test 33

statistics and degrees of freedom (cf. Bakker and Wicherts, 2011). However, he did observe a bump

for *reported p*-values. As such, this highlights an important difference between reported *p*-values and recalculated *p*-values, and stresses the need to distinguish both types of results when studying signs of

<sup>37</sup> questionable research practices.

<sup>38</sup> In this paper we differentiate between two forms of peculiar prevalence of *p*-values just below .05; a

- <sup>39</sup> bump and monotonic excess. Monotic excess signifies a higher than expected frequency of p-values just <sup>40</sup> below .05, but in the absence of a bump, as in Figure 1b below.
- In light of the aforementioned conflicting findings and interpretations, the present paper attempts to answer two questions: (1) Does a bump or monotonic excess of *p*-values below .05 exist in psychology? and (2) Did evidence for a bump increase over time in psychology? We chose to focus on psychology
- and (2) Did evidence for a bump increase over time in psychology? We chose to focus on psychology
   because of the availability of an extensive database on statistical results in psychology (used in Nuijten
- et al., 2015) and because discussions on research practices are particularly salient in this discipline (e.g.,
- Pashler and Wagenmakers, 2012; John et al., 2012; Simmons et al., 2011; Wagenmakers et al., 2012;
- 47 Asendorpf et al., 2013).

First we clarify how the two research questions relate to questionable research practices (QRPs). QRPs 48 are defined as practices that are detrimental to the research process (Panel on Scientific Responsibility 49 and the Conduct of Research, 1992), with a recent focus on those which "increase the likelihood of 50 finding support for a false hypothesis" (p.524 John et al., 2012). Several QRPs related to significance 51 testing are known to affect *p*-values of statistical tests and consequently the decisions based on these 52 tests. Specifically, particular QRPs may yield results that are just significant and can create a bump 53 54 of *p*-values, such as ad hoc exclusion of outliers (Bakker and Wicherts, 2014), repeatedly sampling new participants and checking the results (i.e., data peeking, Armitage et al., 1969), including various 55 combinations of covariates until a significant result is reached, or operationalizing a measure in different 56 ways until significance is reached (Simmons et al., 2011). 57

These QRPs have been used by many researchers at least once in their career. For instance, data peeking and the ad hoc exclusion of outliers were admitted by 63% and 38% of psychological researchers, respectively (John et al., 2012). On the other hand, other QRPs mainly yield very small and (clearly) significant *p*-values, such as analyzing multiple conditions or correlated variables and selecting only the smallest *p*-value out of this set of analyses (van Aert et al., 2015; Ulrich and Miller, 2015) and do not lead to a bump. To summarize, different QRPs may differently affect the distribution of statistically significant *p*-values.

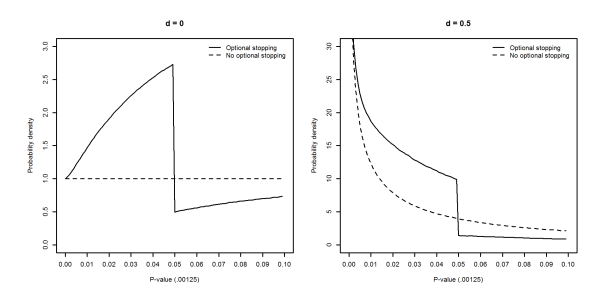
In the absence of QRPs, the distribution of significant *p*-values can be expected to have a certain shape. 65 Under the null-hypothesis all *p*-values are equally probable (i.e., follow a uniform distribution). If there 66 is truly an effect, smaller *p*-values are more likely than larger *p*-values (i.e., the distribution decreases 67 monotonically in the *p*-value). Consequently, because some hypotheses are false and some are true, the 68 distribution of observed *p*-values arises from a mixture of uniform and right-skewed distributions and 69 should also decrease monotonically.<sup>1</sup> Deviation from a monotonically decreasing distribution (i.e., a 70 bump) could indicate evidence of QRPs that aim to obtain just significant results. Hence answering our 71 research questions of whether a bump exists and whether this bump changed over time may also inform 72 us on the prevalence of these particular QRPs and its development over time. 73

However, there are at least two problems with using *p*-value distributions to examine the prevalence 74 75 of QRPs. First, as we previously argued, not all QRPs lead to a bump of p-values just below .05. Hence, examining the distribution of p-values just below .05 will not inform us on the prevalence of QRPs that do 76 not aim to obtain just significant results but yield mainly small and clearly significant p-values (van Aert 77 et al., 2015; Ulrich and Miller, 2015). Second, the QRPs yielding just significant results do not necessarily 78 result in a non-monotonic *p*-value distribution, that is, a distribution with a *bump*. For instance, consider 79 Figure 1 that shows the result of simulations done for data peeking, which is known to result in mainly 80 just significant p-values (Armitage et al., 1969; Lakens, 2015b; Wagenmakers, 2007). The dashed lines in 81 both panels correspond to 20 million simulated p-values under the null-hypothesis and a medium effect 82 size (d = .5), respectively, in a two-sample t-test with 24 participants per group. The solid lines show 83 the distributions of 20 million simulated *p*-values under these same effect sizes and designs, but after a 84 maximum of three rounds of data peeking with each round adding 1/3 of the original sample size. Figure 1 85 illustrates that data peeking may result in non-monotonic excess (i.e., bump; left panel), but can also cause

<sup>&</sup>lt;sup>1</sup>One exception to this rule is when the alternative hypothesis is wrongly specified, that is, if the true effect size is negative whereas the alternative hypothesis states that the true effect is positive. In this case the distribution of the p- value is left-skewed and monotonically increasing.

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monotonic excess (right panel), even if all researchers use data peeking. Specifically, if all underlying 87 effects are genuinely and substantially different from zero (right panel), data peeking will generally not 88 lead to a bump below .05. In the present paper, we therefore examine the peculiar prevalence of *p*-values 89 just below .05 by both investigating the presence of a bump and monotonic excess in distributions of 90 statistically significant results.



**Figure 1.** Distributions of 20 million *p*-values each, when d = 0 (bump; left) and d = .5 (monotonic excess; right), given data peeking (solid) or no data peeking (dashed). Simulations were run for two-sample t-tests with  $n_k = 24$ . For data peeking, a maximum of three rounds of additional sampling occurred if the result was nonsignificant, with each round adding 1/3 of the original sample size.

In answering our research questions, whether a bump or monotonic excess exist and whether the bump 92 93 changed over time, we improve previous studies on four dimensions. First, we eliminate the distortive effects of publication bias on the *p*-value distribution by inspecting only statistically significant results. 94 Second, we use a large dataset on *p*-values from entire articles instead of only *p*-values from abstracts 95 (as in Jager and Leek, 2014; de Winter and Dodou, 2015). Third, we distinguish between reported and 96 recalculated *p*-value distributions for the same set of test results and show that this distinction affects 97 answers to the two questions because of common mismatches (Bakker and Wicherts, 2011). Fourth, we 98 fit analytic models to p-value distributions to investigate for monotonic excess, where previous research 99 only investigated whether there was non-monotone excess (i.e., a bump). 100

Publication bias distorts the *p*-value distribution, but distortions caused by this bias should not be 101 confounded with distortions caused by QRPs. Publication bias refers to the selective publication of 102 disproportionate amounts of statistically significant outcomes (Gerber et al., 2010; Franco et al., 2014). 103 Publication bias contributes to a higher frequency of *p*-values just below .05 relative to the frequency 104 of *p*-values just above .05, but only does so by decreasing the frequency of *p*-values *larger* than .05. 105 Masicampo and Lalande (2012) and de Winter and Dodou (2015) indeed found this relatively higher 106 frequency, which is more readily explained by publication bias, which affects the distribution of p-values 107 larger than .05. QRPs that lead to a bump affect only the distribution of p-values smaller than .05 (Lakens, 108 2015b). We focus only on the distribution of significant p-values, because this distribution is affected by 109 QRPs that cause a bump or monotonic excess and not, or to a much lesser extent, by publication bias. 110 The second extension is the use of more extensive data for psychology than previously used to inspect 111

QRPs that cause a bump or monotonic excess, improving our ability to examine the prevalence of QRPs. 112 Masicampo and Lalande (2012) and Leggett et al. (2013) manually collected *p*-values from a relatively 113 small set of full research articles (i.e., 3,627 and 3,701), whereas Jager and Leek (2014) and de Winter 114 and Dodou (2015) used automated extraction of *p*-values from only the abstracts of research papers. 115 However, *p*-values from abstracts are not representative for the population of *p*-values from the entire 116 paper (Benjamini and Hechtlinger, 2014; Ioannidis, 2014), even though some have argued against this 117

(Pautasso, 2010). Our large scale inspection of full-text articles is similar to papers by Head et al. (2015)
 and Krawczyk (2015), but with the addition of being able to recalculate all *p*-values from the test statistics
 extracted from the text.

Third, we examine the prevalence of QRPs that cause a bump or monotonic excess by investigating 121 both reported and the accompanying recalculated *p*-values. Previous studies do not fully compare results 122 from reported *p*-values and recalculated *p*-values. This distinction is relevant, because reported *p*-values 123 are subject to reporting bias such as rounding errors, particularly relevant around the .05 threshold. Such 124 reporting biases result in inaccurate *p*-value distributions. For example, there is evidence that reporting 125 errors that affect statistical significance occur in approximately 10-15% of papers in psychology (i.e., 126 gross inconsistencies Bakker and Wicherts, 2011; García-Berthou and Alcaraz, 2004; Nuijten et al., 2015; 127 Veldkamp et al., 2014). The advantage of analyzing recalculated p-values is that they contain more 128 decimals than typically reported and correct reporting errors. Some previous studies analyzed reported 129 p-values (de Winter and Dodou, 2015; Jager and Leek, 2014; Head et al., 2015), whereas others looked 130 at recalculated *p*-values (Masicampo and Lalande, 2012) or a mix of reported and recalculated (Leggett 131 et al., 2013). Only Krawczyk (2015) used both reported and recalculated *p*-values for a subset of the 132 data, and they found that the peculiar prevalence below .05 disappeared when the recalculated data were 133 used. Hence, this distinction between reported and recalculated *p*-values allows us to distinguish between 134 peculiarities due to reporting errors and peculiarities due to QRPs such as data peeking. 135

Fourth, we examine the prevalence of *p*-values just below .05 by taking into account various models 136 to test and explain characteristics of p-value distributions. We applied tests and fitted models to p-values 137 138 below .05, in two ways. We first applied the non-parametric Caliper test (Gerber et al., 2010) comparing frequencies of *p*-values in an interval just below .05 to the frequency in the adjacent lower interval; a 139 higher frequency in the interval closest to .05 is evidence for QRPs that seek to obtain just significant 140 results. The Caliper test has also been applied to examine publication bias, by comparing just significant 141 to just nonsignificant p-values (Kühberger et al., 2014), and to detect QRPs (Head et al., 2015). However, 142 the Caliper test can only detect a a bump but not monotonic excess, as illustrated by the distributions of 143 *p*-values in Figure 1. Therefore, we also attempted to model the distribution of significant *p*-values in 144 order to investigate for all forms of excess (i.e., both a bump and monotonic excess), and illustrate the 145 results and difficulties of this approach. 146

In short, this paper studies the distribution of significant *p*-values in four ways. First, we verified whether a bump is present in *reported p*-values just below .05 with the Caliper test. Second, to examine how reporting errors might influence *p*-value distributions around .05, we analyzed only the recalculated *p*-values corresponding to those reported as .05. Third, we used the Caliper test to examine if a bump effect is present in *recalculated p*-values and whether evidence for a bump changed over time. Finally, we modeled the distribution of significant recalculated *p*-values in an attempt to also detect a monotonic excess of *p*-values below .05.

## 154 DATA AND METHODS

#### 155 Data

We investigated the *p*-value distribution of research papers in eight high impact psychology journals (also used in Nuijten et al., 2015). These eight journals were selected due to their high-impact across different subfields in psychology and their availability within the Tilburg University subscriptions. This selection also encompasses the journals covered by Masicampo and Lalande (2012) and Leggett et al. (2013). A summary of the downloaded articles is included in Table 1.

Journal	Acronym	Timespan	Articles downloaded	Articles with extracted results (%)	APA results extracted
Developmental Psychology	DP	1985-2013	3,381	2,607 (77%)	37,658
Frontiers in Psychology	FP	2010-2013	2,126	702 (33%)	10,149
Journal of Applied Psychology	JAP	1985-2013	2,782	1,638 (59%)	15,134
Journal of Consulting and Clinical Psychology	JCCP	1985-2013	3,519	2,413 (69%)	27,429
Journal of Experimental Psychology General	JEPG	1985-2013	1,184	821 (69%)	18,921
Journal of Personality and Social Psychology	JPSP	1985-2013	5,108	4,346 (85%)	101,621
Public Library of Science	PLOS	2000-2013	10,303	2,487 (24%)	31,539
Psychological Science	PS	2003-2013	2,307	1,681 (73%)	15,654
		Total	30.710	16.695 (54%)	258,105

**Table 1.** Articles downloaded, articles with extracted APA results, and number of extracted APA test results per journal.

For these journals, our sample included articles published from 1985 through 2013 that were available in HTML format. For the PLOS journals, HTML versions of articles were downloaded automatically with the rplos package (v0.3.8; Chamberlain et al., 2015). This package allows an R user to search the PLOS database as one would search for an article on the website.<sup>2</sup> We used this package to retrieve search results that include the subject 'psychology' for (part of) an article. For all other journals, HTML versions of articles were downloaded manually by the first author.

APA test results were extracted from the downloaded articles with the R package statcheck (v1.0.1; 167 Epskamp and Nuijten, 2015). The only requirement for this package to operate is a supply of HTML (or 168 PDF) files of the articles that are to be scanned and statcheck extracts all test results reported according 169 170 to the standards of the American Psychological Association (APA; American Psychological Association, 2010). This format is defined as test results reported in the following order: the test statistic and degrees 171 of freedom (encapsulated in parentheses) followed by the p-value (e.g., t(85) = 2.86, p = .005). This 172 style has been prescribed by the APA since at least 1983 (American Psychological Association, 1983, 173 2001), with the only relevant revision being the precision of the reported *p*-value, changing from two 174 decimal places to three decimal places in the sixth edition from 2010. statcheck extracts t, F,  $\chi^2$ , Z 175 and r results reported in APA style. Additional details on the validity of the statcheck package can be 176 found in Nuijten et al. (2015). 177 From the 30,710 downloaded papers, statcheck extracted 258,105 test results. We removed 55 178

From the 50,710 downloaded papers, statcheck extracted 258,105 test results. We removed 55 results, because these were impossible test results (i.e., F(0,55) = ... or r > 1). The final dataset thus included 258,050 test results. The extracted test results can have four different formats, where test results or *p*-values are reported either exactly (e.g., p = .042) or inexactly (e.g., p < .05). Table 2 shows the composition of the dataset, when split across these (in)exactly reported *p*-values and (in)exactly reported test results.

	Exact test statistic	Inexact test statistic	
Exact <i>p</i> -value	68,776	274	69,050 (27%)
Inexact <i>p</i> -value	187,617	1,383	189,000 (73%)
	256,393 (99.36%)	1,657 (0.64%)	258,050 (100%)

**Table 2.** Composition of extracted APA test results with respect to exact and inexact reporting of *p*-values or test statistics.

From this dataset, we selected six subsets throughout our analyses to investigate our research questions 184 regarding a bump below .05. We analyzed (i) all reported p-values (N = 258,050) for a bump in their 185 distribution just below .05. Subsequently we analyzed (ii) only exactly reported p-values (N = 69,050). 186 It is possible that reporting or rounding errors have occurred among the reported *p*-values. To investigate 187 the degree to which this happens at p = .05, we analyzed (iii) exactly reported test statistics that are 188 accompanied by an exactly reported p-value of .05 (i.e., p = .05). This subset contains 2,470 results. To 189 debilitate rounding errors and other factors influencing the reporting of *p*-values (e.g., Ridley et al., 2007), 190 we also investigated the recalculated *p*-value distribution with (iv) *p*-values that were accompanied by 191 exactly reported test statistics (N = 256, 393). To investigate whether evidence for a bump differs for 192 inexactly and exactly reported p-values, (v) 68,776 exactly reported test statistics with exactly reported 193 *p*-values were analyzed. Finally, we used (vi) all recalculated *p*-values in 0-.05 for *t*, *r*, and  $F(df_1 = 1)$ 194 values to model the effect size distribution underlying these *p*-values to investigate evidence of both a 195 bump and monotonic excess. 196

#### 197 Methods

We used the Caliper test and two new measures to examine if the observed *p*-value distribution shows evidence for a bump or monotonic excess below .05. We applied the two measures to the observed *p*-value distribution and we examined their performance to detect a bump or monotonic excess using a simulation study on data peeking. Data peeking was chosen because it is one of the most frequently used and well-known QRPs. Below, we explain the Caliper test, how the *p*-value distributions are modeled with the two new measures, and describe the design of the simulation study in more detail.

<sup>&</sup>lt;sup>2</sup>We note there are minor differences in the number of search results from the PLOS webpage and the rplos package for equal searches. This is due to differences in the default search database for the webpage and the package. For technical details on this issue, see https://github.com/ropensci/rplos/issues/75

#### 204 Caliper test

In order to test for a bump of *p*-values just below .05, we applied the Caliper test (e.g., Gerber et al., 2010; Kühberger et al., 2014). This proportion test compares the frequencies of *p*-values in two intervals, such as the intervals .04-.045 and .045-.05. Let *Pr* denote the proportion of *p*-values of the interval .045-.05. Then, independent of the population effect sizes underlying the *p*-values, *Pr* should not be higher than .5 in any situation because the *p*-value distribution should be monotone decreasing. Hence Pr > .5 signifies a bump of *p*-values just below .05.

We carried out one-tailed binomial proportion tests, with  $H_0: Pr \le .5$  and  $H_1: Pr > .5$ . For example, if 40 and 60 *p*-values are observed in the intervals .04-.045 and .045-.05, respectively, then Pr = .6 and the binomial test results in *p*-value = .0284, suggesting evidence for a bump below .05. We applied the Caliper test to the reported *p*-values (subsets one through three as described in the previous section) and recalculated *p*-values (subsets four and five), both for the entire dataset and each of the eight psychology journals.

The Caliper test requires specifying the width of the intervals that are to be compared. For reported 217 p-values, which are frequently rounded to two-decimal values, we selected the intervals (.03875-.04] and 218 [.04875-.05] because there is a strong preference to report *p*-values to the second decimal in research 219 papers (see also Hartgerink, 2015). For recalculated *p*-values we used the same interval width as used by 220 Masicampo and Lalande (2012) and Leggett et al. (2013), which is .00125, corresponding to a comparison 221 of intervals (.0475-.04875] and [.04875-.05). Note that rounding is not a problem for recalculated *p*-values. 222 Considering that some journals might show small frequencies of *p*-values in these intervals, we also 223 carried out Caliper tests with interval widths of .0025, .005, and .01. Note that, on the one hand, increasing 224 interval width increases the statistical power of the Caliper test because more p-values are included in 225 the test, but on the other hand also decreases power because Pr is negatively related to interval width 226 whenever *p*-values correspond to tests of non-zero population effects. In other words, a bump just below 227 .05 will tend more and more towards a monotonically decreasing distribution as the binwidth increases. 228 To verify if evidence for a bump of *p*-values increased over time, we fitted a linear trend to proportion 229 Pr of the Caliper test with binwidths .00125, .0025, .005, and .01. We computed these proportions for 230 each year separately, for both the total dataset and per journal. Time was centered at the start of data 231

collection, which was 1985 except for PLOS (2000), PS (2006; due to 0 *p*-values in the considered interval for preceding years), and FP (2010). The value .5 was subtracted from all Pr values, such that the intercept of the trend corresponds to the bump of *p*-values at the start of data collection, where 0 means no bump. A positive linear trend signifies an increase in the bump of *p*-values below .05 over time.

#### <sup>236</sup> Measures based on *p*-value distributions

Figure 1 demonstrates that data peeking has a different effect on the *p*-value distribution depending on the true effect size. The distribution after data peeking does not monotonically increase for d = 0 (left panel), whereas it does increase monotonically for d = 0.5 (right panel). Consequently, the Caliper test will signal a bump of *p*-values for d = 0 (i.e., it will detect a bump), but not for d = 0.5.

We examined how we may be able to detect both a bump and monotonic excess of *p*-values below 0.05. Figure 1 indicates that, for *p*-values close to zero (e.g.,  $\leq .00125$ ) the *p*-value distributions with data peeking (solid lines) closely match the *p*-value distributions without data peeking (dashed lines). In other words, data-peeking in studies with initially nonsignificant *p*-values rarely results in tiny significant *p*-values, but more often in *p*-values larger than .00125. The basic idea of this analysis is therefore to estimate the 'true' effect size distribution using only these tiny *p*-values (i.e.,  $\leq .00125$ ), assuming that none or a very small proportion of these *p*-values were affected by by data-peeking.

We examined the performance of two measures to detect a bump or monotonic excess of *p*-values 248 below .05. The first method compares the effect sizes estimated on *p*-values smaller than .00125 to effect 249 sizes estimated using all *p*-values smaller than .05. The idea of this first method is that increasing the 250 frequency of just-significant *p*-values *decreases* the effect size estimate. Indeed, the more right-skewed 251 the *p*-value distribution, the higher the effect size estimate when keeping constant studies' sample sizes 252 (Simonsohn et al., 2014; van Assen et al., 2015). According to the first method, there is evidence 253 suggestive of data peeking (or other QRPs leading to a bump of p-values just below .05) if the effect size 254 estimate is considerably lower when based on all p-values than when based on only p-values  $\leq .00125$ . 255

The second method yields a measure of excess of p-values just below .05, for either a bump or monotonic excess, by comparing the observed frequency of p-values in the interval .00125-.05 to the

predicted frequency of p-values in that interval. This prediction is based on the effect size estimated using the p-values smaller than .00125. If the ratio of observed over expected p-values is larger than 1, referred to as statistic D, then this could indicate data peeking. Statistic D is calculated as

$$D = \frac{p_{.00125}^o}{1 - p_{.00125}^o} \times \frac{1 - p_{.00125}^e}{p_{.00125}^e}$$
(1)

with  $p_{.00125}^o$  and  $p_{.00125}^e$  representing the proportion of *p*-values lower than .00125 observed and expected, respectively. Note that *D* is an odds ratio.

#### 258 Modeling *p*-value distributions

For both measures from the previous section the expected *p*-value distribution needs to be derived and compared to the observed *p*-value distribution. The observed *p*-value distribution of the psychology data is based on all exactly reported statistics with test statistics *t*, *r*, and  $F(1,df_2)$ , because these readily provide the same effect measure. We used the Fisher transformed correlation,  $\rho_F$ , as effect size measure. The distribution of the Fisher transformed correlation is approximated well by the normal distribution, with Fisher transformation

$$\rho_F = \frac{1}{2} ln(\frac{1+r}{1-r}) \tag{2}$$

and standard error  $\frac{1}{\sqrt{N-3}}$  or  $\frac{1}{\sqrt{df_2-1}}$ .  $F(1, df_2)$  and t values can be transformed to correlations using

$$r = \frac{\frac{F \times df_1}{df_2}}{\frac{F \times df_1}{df_2} + 1} \tag{3}$$

259 where  $F = t^2$ .

The expected *p*-value distribution was estimated under the assumption that the true effect size, Fisher transformed correlation  $\rho_F$ , is normally distributed with  $\mu_{\rho_F}$  and standard deviation  $\tau_{\rho_F}$ . The two parameters were estimated by minimizing the  $\chi^2$ -statistic

$$\chi_{j-1}^2 = \sum_{j=1}^J \frac{(rf_j^o - rf_j^e)^2}{rf_j^e}$$
(4)

with  $rf^o$  and  $rf^e$  being the relative frequency of observed and expected *p*-values in interval *j*, respectively. Minimization was done with the optim() function in R, where  $\hat{\tau}$  was truncated to be positive. Interval *j* is defined as  $(I_{j-1}, I_j) = ((j-1)x, jx))$ , with width x = .00025 whenever only the significant *p*-values lower than .00125 were modeled (resulting in 5 intervals); .00125 when modeling all significant *p*-values (i.e.,  $p \le .05$ , 40 intervals); .025 when modeling all *p*-values (also 40 intervals). The relative frequencies are conditional probabilities. For instance,  $rf_2^o$  is the proportion of observed *p*-values in interval  $(I_1 = .00025, I_2 = .0005)$  whenever *p*-values lower than .00125 are examined. Expected relative frequency  $rf_i^e$  is computed as

$$rf_{j}^{e} = \frac{\sum_{k=1}^{K} P(I_{j-1} \le p_{k} \le I_{j} | N_{k}; \hat{\rho}_{F}; \hat{\tau}_{\rho_{F}})}{\sum_{k=1}^{K} P(p_{k} \le I_{J} | N_{k}; \hat{\rho}_{F}; \hat{\tau}_{\rho_{F}})}$$
(5)

with the summation over all *K* significant test statistics. *P* corresponds to the probability that a *p*-value of study *k* (i.e.,  $p_k$ ) is in a certain interval, which depends on the study sample size  $N_k$  and the two estimated parameters of the effect size distribution (i.e.,  $\hat{p}_F$ ,  $\hat{\tau}_{p_F}$ ).

#### 263 Design of simulation study

<sup>264</sup> To examine the potential of the two measures to detect data peeking, their performance was examined

- on simulated data with and without data peeking. We used a two-group between-subjects design with
- <sup>266</sup> 24 participants per group ( $n_k = 24$ ), and compared their means using a *t*-test. The performance of both
- measures was examined as a function of true effect size  $\mu$  (0; 0.2; 0.5; 0.8) and heterogeneity  $\tau$  (0; 0.15).

In the data peeking conditions, data were simulated as follows: means and variances per group were simulated and a two-sample *t*-test was conducted. If this *t*-test was statistically significant (i.e.,  $p \le .05$ ), the *p*-value was stored, otherwise the data peeking procedure was started. In this data peeking procedure, one-third of the original sample size was added to the data before conducting another two-sample *t*-test. This data peeking procedure was repeated until a statistically significant result was obtained or three rounds of additive sampling had taken place (see osf.io/x5z6u for simulation functions). The simulations were stopped if 1,000,000 studies with a *p*-value below .1 were obtained for each combination of  $\mu$  and  $\tau$ .

# 275 RESULTS AND DISCUSSION

In this section, we report the results of our analyses in the following order for the subsets: all reported 276 p-values (258,050 results), exactly reported p-values (69,050 results), p-values erroneously reported as 277 equal to .05 (2,470 results), all recalculated p-values based on exactly reported test statistics (256,393 278 results), recalculated *p*-values based on exactly reported test statistics and exactly reported *p*-values 279 (68,776 results), and the modeling of p-value distributions based on recalculated p-values 0-.00125 and 280 0-.05 (54,561 results and 127,509, respectively). These analyses apply the Caliper test to investigate 281 evidence of a possible bump below .05. Subsequently, the results of the two measures are presented based 282 on all recalculated *p*-values. 283

### 284 Reported p-values

Figure 2 shows the distribution for all reported p-values (i.e., 258,050; white bars) and exactly reported p-values (i.e., 69,050; blue bars). Results of the Caliper test indicate (i) there is a bump just below .05 when

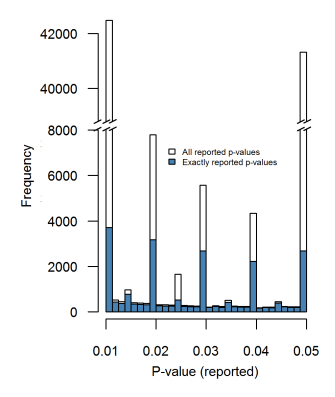
values (i.e., 69,050; blue bars). Results of the Caliper test indicate (i) there is a bump just below .05 when considering all reported *p*-values in bins .03875-.04 versus .04875-.05, N = 45,667, Pr = 0.905, p < .001

and (ii) there is still a bump, but less so, when considering only exactly reported p-values in these bins,

N = 4,900, Pr = 0.547, p < .001. The difference in bumps between these two subsets can be explained

by the amount of *p*-values that are reported as < .05, which is 86% of all *p*-values reported as exactly

<sup>291</sup> equal to .05 and 14% of all reported *p*-values.



**Figure 2.** Distributions of all reported *p*-values (white) and exactly reported *p*-values (blue) across eight psychology journals. Binwidth = .00125.

292

To investigate whether this observed bump below .05 across exactly reported *p*-values originates from

one or multiple journals, we performed the Caliper test on the exactly reported *p*-values per journal. Table 293 3 shows the results for these tests. The results indicate that there is sufficient and reliable evidence for 294 a bump below .05 (i.e., Pr > .5) for the journals DP and JPSP and sufficient evidence, but debatable 295 reliability for JAP, where the results depend on the binwidth. However, the other five journals show no 296 evidence for a bump below .05 in exactly reported *p*-values at all. In other words, the bump below .05 in 297 exactly reported *p*-values is mainly driven by the journals DP, JAP, and JPSP. 298

Binwidth		0.0	0125			0.0	0025		0.005				0.01			
	x	Ν	Pr	р	х	N	Pr	р	x	N	Pr	р	х	Ν	Pr	р
All	2,682	4,900	0.547	< .001	2,881	5,309	0.543	< .001	3,308	6,178	0.535	<.001	4,218	8,129	0.519	< .001
DP	319	531	0.601	< .001	336	567	0.593	< .001	383	653	0.587	< .001	464	843	0.55	0.002
FP	96	193	0.497	0.557	105	227	0.463	0.884	141	304	0.464	0.906	215	458	0.469	0.912
JAP	78	131	0.595	0.018	82	137	0.599	0.013	85	154	0.552	0.113	101	183	0.552	0.092
JCCP	246	517	0.476	0.874	267	562	0.475	0.889	308	641	0.48	0.848	395	823	0.48	0.882
JEPG	147	285	0.516	0.318	159	310	0.513	0.346	195	375	0.52	0.235	258	509	0.507	0.395
JPSP	1,252	2,097	0.597	< .001	1,310	2,207	0.594	< .001	1,408	2,399	0.587	<.001	1,623	2,869	0.566	< .001
PLOS	307	649	0.473	0.921	366	760	0.482	0.854	489	1,000	0.489	0.766	744	1,558	0.478	0.964
PS	237	497	0.477	0.859	256	539	0.475	0.886	299	652	0.459	0.984	418	886	0.472	0.957

**Table 3.** Caliper test for exactly reported *p*-values per journal for different binwidths. *x* = frequency of *p*-values in .05 minus binwidth through .05, N = total frequency of *p*-values across both intervals in the comparison, Pr = x/N, p = p-value of the binomial test. Significant results ( $\alpha = .05$ , one-tailed) indicating excess of *p*-values just below .05 and are reported in bold.

The Caliper test results for reported *p*-values indicate two things: (i) inexactly reported *p*-values 299 severely distort the *p*-value distribution, and (ii) a bump below .05 is also found when only considering 300 exactly reported *p*-values. Because inexact reporting of *p*-values causes excess at certain points of the 301 *p*-value (e.g., the significance threshold .05; Ridley et al., 2007), we recommend only inspecting exactly 302 reported *p*-values when examing *p*-value distributions. 303

Considering only exactly reported *p*-values, there is sufficient evidence for a bump below .05 in the 304 journals DP, JAP, and JPSP, but not in the remaining five journals (i.e., FP, JCCP, JEPG, PLOS, PS). A 305 tentative explanation of the bump of p-values just below .05 for DP, JAP, and JPSP may be that QRPs that 306 aim to obtain barely significant results are more frequent in the fields of these journals. However, another 307 explanation may be that scientists in these fields are more prone to exactly report p-values just below .05 308 (e.g., to emphasize they are really smaller than .05) than *p*-values considerably smaller than .05. 309

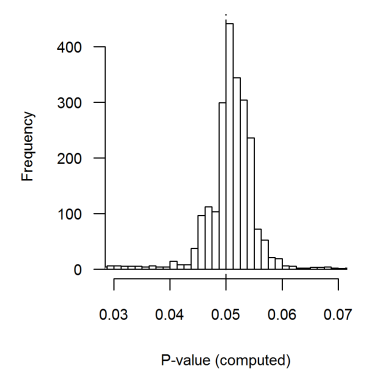
#### Recalculated *p*-value distributions 310

#### Recalculated when reported p = .05 311

Results for reported *p*-values remain inconclusive with regard to the distribution of *p*-values, due to 312 potential rounding or errors (Bakker and Wicherts, 2011; Nuijten et al., 2015; Veldkamp et al., 2014). 313 Rounding and errors could result in an over-representation of p-values  $\leq .05$ . To investigate the plausibility 314 of this notion, we inspected recalculated p-values when p = .05 was reported. Figure 3 indicates that 315 *p*-values that were reported as .05 show remarkable spread when recalculated, which indicates that the 316 reported *p*-value might frequently be rounded or incorrect, assuming that the reported test statistics are 317 correct. More specifically, 67.45% of p-values reported as .05 were larger than .05 when recalculated and 318 319 32.55% were smaller than .05. This percentage does not greatly vary across journals (range 58.8%-73.4% compared to 67.45%). Taking into account rounding possibilities (i.e., widening the range of correct 320 *p*-values to .045-.055), these percentages become 13.81% and 7.85%, respectively, meaning that at least 321 21.66% of the *p*-values reported as .05 was incorrectly reported. In comparison, *p*-values reported as 322 p = .04, p = .03, or p = .02 show smaller proportions of downward rounding when compared to p = .05323 (i.e., 53.33%, 54.32%, 50.38%, respectively compared to 67.45%). When taking into account potential 324 rounding errors in the initial reporting of *p*-values, the discrepancy remains but to a smaller extent (i.e., 325 11.74%, 9.57%, 8.03%, respectively compared to 13.81%). These results provide direct evidence for 326 the QRP "incorrect rounding of p-value" (John et al., 2012), which contributes to a bump or monotonic 327 excess just below .05. 328

The discrepancy between recalculated *p*-values and *p*-values reported as equal to .05 highlights 329 the importance of using recalculated *p*-values when underlying effect distributions are estimated as in 330 *p*-uniform and *p*-curve (van Assen et al., 2015; Simonsohn et al., 2014). When interested in inspecting the 331

*p*-value distribution, reported *p*-values can substantially distort the *p*-value distribution, such that results 332



**Figure 3.** Distribution of recalculated *p*-values where the *p*-value is reported as p = .05. 9.7% of the results fall outside the range of the plot, with 3.6% at the left tail and 6.1% at the right tail. Binwidth = .00125

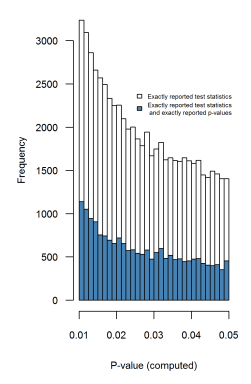
become biased if we rely solely on the reported p-value. Such a discrepancy indicates potential rounding of p-values, erroneous reporting of p-values, or strategic reporting of p-values. The p-value distortions can be (partially) corrected for by recalculating p-values based on reported test statistics. Additionally, potential distortions to the distribution at the third decimal place due to the rounding of p-values to the second decimal (Hartgerink, 2015) is also solved by recalculating p-values. We continue with recalculated p-values in our following analyses.

#### 339 Recalculated p-values

Figure 4 shows the distribution of all recalculated *p*-values (i.e., set of 256,393 results) and of recalculated *p*-values whenever the reported *p*-value is exact (i.e., set of 68,776 results). The recalculated *p*-value distribution is markedly smoother than the reported *p*-value distribution (see Figure 2) due to the absence of rounded *p*-values.

After inspecting all recalculated p-values, we did not observe a bump just below .05, N = 2,808, Pr =344 .5, p = 0.508. When we analyzed the recalculated *p*-values per journal (Table 4), there is no evidence 345 for a bump below .05 in any of the journals. Additionally, we inspected all recalculated p-values that 346 resulted from exactly reported p-values. For this subset we did observe a bump below .05, N = 809, Pr =347 0.564, p = 0.000165 (blue histogram in Figure 4) for the smallest binwidth (i.e., .00125), but this effect 348 was not robust across larger binwidths, as shown in Table 5. This table also specifies the results for a 349 bump below .05 per journal, with sufficient evidence of a bump only in JPSP. This finding, however, was 350 only observed for binwidths .00125 and .0025, not for larger binwidths. Considering the results from 351 the recalculated *p*-values, there is sparse evidence for the presence of a bump below .05, opposed to 352 widespread evidence (Masicampo and Lalande, 2012; Leggett et al., 2013; Head et al., 2015). Moreover, 353 interpretation of the bump for JPSP is not straightforward; it may also be that authors of JPSP are more 354 prone to report exact test statistics if the *p*-value is just below .05 than whenever *p*-values are considerably 355 smaller than .05. 356





**Figure 4.** Recalculated *p*-values for exactly reported test statistics (white bars), and recalculated *p*-values for exactly reported test statistics where *p*-values are also exactly reported (blue bars). Binwidth = .00125

#### 357 Excessive significance over time

The regression results of the development of a bump below .05 over time, based on recalculated *p*-values, 358 are shown in Table 6. Results indicate that there is no evidence for a linear relation between time in years 359 and the degree to which a bump of *p*-values below .05 is present across the different binwidths (only 360 results for binwidth .00125 are presented; results for the other binwidths available at http://osf.io/96kbc/). 361 Conversely, for PLOS there is some evidence for a minor increase of a bump throughout the years 362 (b = .072, p = .039), but this result is not robust for binwidths .0025, .005, and .01. These results contrast 363 with Leggett et al. (2013), who found a linear relation between time and the degree to which a bump 364 occurred for JEPG and JPSP. Hence, our findings contend the increase of a bump below .05 for the period 365 1965-2005 in psychology (Leggett et al., 2013). In other words, our results of the Caliper test indicate 366 that, generally speaking, there is no evidence for an increasing prevalence of p-values just below .05 or of 367 QRPs causing such a bump in psychology. 368

Binwidth		0.00	0125			0.0	025			0.0	05		0.01			
	х	Ν	Pr	р	х	N	Pr	р	х	Ν	Pr	р	x	Ν	Pr	р
All	1,404	2,808	0.5	0.508	2,808	5,761	0.487	0.973	5,761	11,824	0.487	0.997	11,824	25,142	0.47	> .999
DP	184	382	0.482	0.779	382	829	0.461	0.989	829	1,710	0.485	0.9	1,710	3,579	0.478	0.996
FP	30	69	0.435	0.886	69	172	0.401	0.996	172	376	0.457	0.956	376	799	0.471	0.955
JAP	73	145	0.503	0.5	145	270	0.537	0.124	270	556	0.486	0.765	556	1,168	0.476	0.952
JCCP	160	308	0.519	0.265	308	633	0.487	0.763	633	1,267	0.5	0.522	1,267	2,706	0.468	> .999
JEPG	81	164	0.494	0.593	164	332	0.494	0.608	332	683	0.486	0.778	683	1,535	0.445	> .999
JPSP	640	1,268	0.505	0.379	1,268	2,557	0.496	0.668	2,557	5,174	0.494	0.802	5,174	10,976	0.471	> .999
PLOS	125	260	0.481	0.752	260	541	0.481	0.828	541	1,170	0.462	0.995	1,170	2,544	0.46	> .999
PS	111	212	0.524	0.268	212	427	0.496	0.577	427	888	0.481	0.88	888	1,835	0.484	0.919

**Table 4.** Caliper test for exactly recalculated *p*-values per journal for different binwidths. x = frequency of *p*-values in .05 minus binwidth through .05, N = total frequency of *p*-values across both intervals in the comparison, Pr = x/N, p = p-value of the binomial test. Significant results ( $\alpha = .05$ , one-tailed) indicating excess of *p*-values just below .05 and are reported in bold.

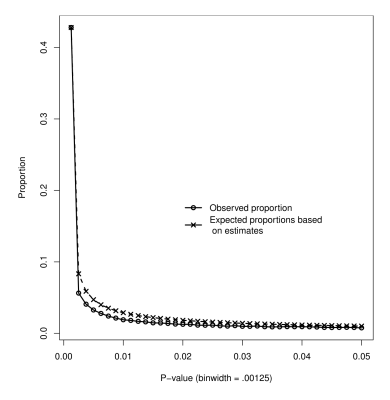
Binwidth		0	.00125			0.0025				0.0	0.005 0.01					
	х	Ν	Pr	р	х	Ν	Pr	р	x	Ν	Pr	р	х	Ν	Pr	р
All	456	809	0.564	< .001	809	1,617	0.5	0.5	1,617	3,403	0.475	0.998	3,403	7,402	0.46	1
DP	46	87	0.529	0.334	87	185	0.47	0.811	185	358	0.517	0.281	358	756	0.474	0.932
FP	15	27	0.556	0.351	27	87	0.31	> .999	87	192	0.453	0.915	192	437	0.439	0.995
JAP	8	20	0.4	0.868	20	29	0.69	0.031	29	65	0.446	0.839	65	141	0.461	0.844
JCCP	43	78	0.551	0.214	78	161	0.484	0.682	161	364	0.442	0.988	364	780	0.467	0.971
JEPG	27	50	0.54	0.336	50	98	0.51	0.46	98	209	0.469	0.834	209	479	0.436	0.998
JPSP	184	305	0.603	<.001	305	547	0.558	0.004	547	1,117	0.49	0.764	1,117	2,451	0.456	> .999
PLOS	76	149	0.51	0.435	149	323	0.461	0.926	323	698	0.463	0.978	698	1,470	0.475	0.975
PS	57	93	0.613	0.019	93	187	0.497	0.558	187	400	0.468	0.912	400	888	0.45	0.999

**Table 5.** Caliper tests for exactly recalculated and exactly reported *p*-values per journal, including alternative binwidths. x = frequency of *p*-values in .05 minus binwidth through .05, N = total frequency of *p*-values across both intervals in the comparison, Pr = x/N, p = p-value of the binomial test. Significant results ( $\alpha = .05$ , one-tailed) indicating excess of *p*-values just below .05 and are reported in bold.

	T:	Caeffeient	Estimate	<u>C</u> E	4	
	Timespan	Coefficient	Estimate	SE	t	p
All	1985-2013	Intercept	0.007	0.017	0.392	0.698
All		Years (centered)	-0.001	0.001	-0.492	0.627
DP	1985-2013	Intercept	-0.043	0.056	-0.769	0.448
DP		Years (centered)	0.001	0.003	0.193	0.849
FP	2010-2013	Intercept	-0.182	0.148	-1.233	0.343
FP		Years (centered)	0.055	0.079	0.694	0.560
JAP	1985-2013	Intercept	0.041	0.081	0.504	0.619
JAP		Years (centered)	-0.001	0.005	-0.208	0.837
JCCP	1985-2013	Intercept	0.077	0.058	1.315	0.200
JCCP		Years (centered)	-0.006	0.004	-1.546	0.134
JEPG	1985-2013	Intercept	-0.022	0.124	-0.176	0.862
JEPG		Years (centered)	0.001	0.007	0.097	0.924
JPSP	1985-2013	Intercept	-0.002	0.027	-0.062	0.951
JPSP		Years (centered)	0.000	0.002	-0.005	0.996
PLOS	2006-2013	Intercept	-0.382	0.114	-3.344	0.016
PLOS		Years (centered)	0.072	0.027	2.632	0.039
PS	2003-2013	Intercept	0.081	0.078	1.045	0.323
PS		Years (centered)	-0.009	0.013	-0.669	0.520

**Table 6.** Linear regression coefficients as a test of increasing excess of *p*-values just below .05. Intercept indicates the degree of excess for the first year of the estimated timespan (> 0 = excess). Significant results ( $\alpha = .05$ , two-tailed) are reported in bold.

- Results of two measures based on modeling *p*-value distributions
- 370 Data of eight psychology journals
- <sup>371</sup> Figure 5 depicts the observed *p*-value distribution and the expected *p*-value distribution corresponding to
- the fitted effect size distribution based on *p*-values  $\leq$  .00125. Estimates for *p*-values  $\leq$  .05 were effect
- size  $\hat{\rho}_F = 0$  and heterogeneity  $\hat{\tau}_{\rho_F} = .183$ , and  $\hat{\rho}_F = .149$  and  $\hat{\tau}_{\rho_F} = .106$  for *p*-values  $\le .00125$ . Misfit
- between observed and expected *p*-value distribution for  $p \le .00125$  was minor ( $\chi^2 = 4.1$ ), indicating that
- the observed *p*-values  $\leq$  .00125 were well approximated by the estimated effect size distribution.



**Figure 5.** Observed proportions of *p*-values (circles) and expected proportions of *p*-values based on  $\hat{\rho}_F$  and  $\hat{\tau}_{\rho_F}$  estimated from 0-.00125 (crosses).

Our first measure suggests practices leading to a monotonic excess of *p*-values below .05, because 376 the estimated effect size based on all significant *p*-values (i.e., 0) is much smaller than the supposedly 377 unbiased estimate based on only the very small p-values (i.e., .183). Moreover, assuming that effect 378 sizes are normally distributed with  $\rho_F = 0$  and  $\tau_{\rho_F} = .183$ , combined with the degrees of freedom of the 379 observed effects, implies that only 27.5% of all effects would be statistically significant. However, of all 380 reported *p*-values, 74.7% were statistically significant, but this difference may at least partly be caused by 381 other factors such as publication bias. It is highly unlikely that the average true effect size underlying 382 statistically significant results in psychology is truly zero. It remains undecided, however, whether this 383 very low estimate is mainly due to QRPs leading to a downward bias of the effect size estimate, or to a 384 misspecification of the model, an issue we revisit later in the paper. 385

For the second measure that compares the ratio of observed and expected *p*-values below .05, we found D = .701, which does not suggest data peeking but *under*-reporting of *p*-values (29.9%) in the *p*-value interval .00125-.05. The simulation results that follow below, however, demonstrated that the measure *D* performs badly under effect size heterogeneity. Since heterogeneity is underlying the observed data, we conclude that the measure *D* is not useful for investigating evidence of a bump or monotonic excess of *p*-values.

#### 392 Simulation study

Table 7 shows the results of the two measures for data simulated with and without data peeking. The

column headers show the mean (i.e.,  $\mu$ ) and heterogeneity (i.e.,  $\tau$ ) of the simulated conditions, with the

			au = 0				$\tau = .15$			
	<i>p</i> -values		$\mu = 0$	$\mu = .2$	$\mu = .5$	$\mu = .8$	$\mu = 0$	$\mu = .2$	$\mu = .5$	$\mu = .8$
	p-values		$ ho_F=0$	$\rho_{F} = .099$	$\rho_{F} = .247$	$\rho_{F} = .390$	$\rho_F = 0$	$\rho_{F} = .099$	$\rho_{F} = .247$	$\rho_{F} = .390$
Without data peeking	0-1	$\hat{ ho}_F$	0	0.103	0.258	0.413	0	0.103	0.258	0.413
		$\hat{\tau}_{ ho_F}$	0	0	0	0	0.077	0.077	0.077	0.077
	005	$\hat{ ho}_F$	0	0.103	0.258	0.413	0	0.103	0.258	0.413
		$\hat{\tau}_{ ho_F}$	0	0	0	0.001	0.077	0.077	0.077	0.077
		Misfit $\chi^2$	0	0	0	0	0	0	0	0
	000125	$\hat{\rho}_F$	0	0.103	0.258	0.413	0.1	0.107	0.259	0.413
		$\hat{\tau}_{ ho_F}$	0	0	0	0.001	0.025	0.076	0.077	0.077
		Misfit $\chi^2$	0	0	0	0	0	0	0	0
		D	1	1	1	1	1.205	1.006	1.003	1.001
With data peeking	005	$\hat{ ho}_F$	0	0	0.117	0.345	0	0	0.075	0.360
1 0		$\hat{\tau}_{ ho_F}$	0	0	0	0.038	0	0.055	0.137	0.091
		Misfit $\chi^2$	126,267.4	50,298.4	696.6	101.6	14,867.6	1,209.5	576.3	340.6
		Ν	759,812	811,296	936,517	994,974	434,660	525,023	707,650	889,681
	000125	$\hat{\rho}_F$	0	0.075	0.218	0.366	0.066	0.161	0.283	0.402
		$\hat{\tau}_{\rho_F}$	0	0	0	0	0.036	0	0	0.012
		Misfit $\chi^2$	6.9	3.2	7.1	11.8	2	1.9	2.6	2.1
		N	9,729	21,576	95,615	350,482	14,791	34,530	124,991	366,875
		D	1.977	1.976	1.835	1.166	1.628	1.620	1.472	1.164

<sup>395</sup> corresponding  $\rho_F$  and  $\tau_{\rho_F}$  on the Fisher transformed correlation scale. The first set of rows shows the <sup>396</sup> results for the data simulated without data peeking, of which we discuss the results first.

**Table 7.** Results of parameter estimation of the distribution of effect sizes and measures of data peeking as a function of population effect size  $(\mu, \rho_F)$ , population heterogeneity  $(\tau)$ , and data peeking, for the simulated data. Results are based on all *p*-values 0-1, *p*-values  $\leq .05$ , and  $\leq .00125$ .  $\hat{\rho}_F$  = estimated population effect,  $\hat{\tau}_{\rho_F}$  = estimated population heterogeneity, misfit 0-.05 = misfit of estimates based on *p*-values 0-.05, misfit 0-.00125 = misfit of estimates based on *p*-values 0-.00125 (bold indicates *p* < .05), *N* = number of results included in estimation, *D* = comparison of observed- and expected *p*-value frequencies.

The results for the data without data peeking inform us on (i) whether the effect size distribution parameters can accurately be recovered using only very small ( $\leq$  .00125) or small *p*-values ( $\leq$  .05), and (ii) if both measures accurately signal no data peeking. Note that  $\rho_F$  is slightly overestimated due to categorizing the *p*-value distribution into 40 categories: the estimates based on all *p*-values (i.e.,  $\hat{\rho}_F$ , first row) are slightly larger than the population parameter (i.e.,  $\rho_F$ , column headers).

Answering the first question of accurate parameter estimates, whenever there is no heterogeneity (i.e.,  $\tau_{\rho_F} = 0$ ) both  $\rho_F$  and  $\tau_{\rho_F}$  are accurately recovered. When heterogeneity is non-zero, the parameters were also accurately recovered, but not when  $\rho_F = 0$ . Here,  $\rho_F$  was overestimated (equal to .1) and  $\tau_{\rho_F}$ underestimated (.025 rather than the true .077), while at the same time the misfit was negligible.

The latter result, that the effect is overestimated under heterogeneity when  $\rho_F = 0$ , is explained by the fact that a *p*-value distribution can accurately be modeled with an infinite range of negatively correlated values of  $\rho_F$  and  $\tau_{\rho_F}$ . An increase in  $\rho_F$  yields a more right-skewed distribution, which is hardly distinguishable from the right-skewed distribution caused by an increase in  $\tau_{\rho_F}$ . The similar effects of both parameters on the fitted *p*-value distribution already hint at potential problems for both measures, because performance of these measures is dependent on accurate estimates of these paramaters.

With respect to the second question, whether the measures accurately signal the absence of data 412 peeking, the first measure does so in both homo- and heterogeneous conditions, whereas the second 413 measure correctly signals absence only under homogeneity. The first measure signals data peeking if the 414 estimate of  $\rho_F$  is smaller when based on  $p \leq .05$  than on  $p \leq .00125$ . Previously, we already noted that 415 effect size estimates were identical to population effect sizes under homogeneity, and equal or *larger* 416 when based on  $p \le .00125$  under heterogeneity. This suggests that the first measure behaves well if there 417 is no data peeking (but see the conclusion section). The second measure, D, performed well (i.e., was 418 equal to 1) under homogeneity, but incorrectly suggested data peeking under heterogeneity. For instance, 419 D = 1.205 for  $\rho_F = 0$  and  $\tau = .15$ , which suggests that 20.5% more p-values were observed in the interval 420 .00125-.05 than were expected based on the  $\hat{\rho}_F$  estimate even though no data peeking occurred. The 421 explanation for the breakdown of the performance of D is that the parameters of the effect size distribution 422 were not accurately recovered, overestimating the average effect size and underestimating heterogeneity 423 based on small p-values. This yields a lower expected frequency of higher p-values (between .00125 and 424

425 .05), thereby falsely suggesting data peeking.

The last rows present the results obtained when data peeking does occur. First consider the estimates 426 of  $\rho_F$  and the performance of the first measure of data peeking. The estimates of  $\rho_F$  confirm that data 427 peeking results in underestimation, particularly if the average true effect size is not large (i.e.,  $\mu = .2$  or 428 .5). Moreover, downward bias of  $\rho_F$  decreases when it is estimated on p-values  $\leq .00125$  than on  $\leq .05$ , 429 accurately signaling data peeking with the first measure. For instance, if  $\rho_F = .099$  and  $\tau = 0$ ,  $\hat{\rho}_F = .075$ 430 when based on p-values  $\leq .00125$  and  $\hat{p}_F = 0$  when based on p-values  $\leq .05$ . Together with the good 431 performance of this measure under no data peeking, these results suggest that the first measure may be 432 useful to detect data keeping in practice. 433

Consider the estimates of  $\tau_{\rho_F}$  and the performance of *D*. Similar to conditions under no data peeking, heterogeneity is grossly underestimated when using *p*-values  $\leq .00125$ . Hence *D* cannot be expected to perform well under data peeking. Although *D*-values seem to correctly signal data peeking in all conditions and decrease as expected when the effect size increases, these values do not correspond to the actual values of data peeking. For instance, consider the condition with  $\mu = .5$  and  $\tau_{\rho_F} = .15$ ; of the 582,659 simulated *p*-values in interval .00125-.05, 106,241 *p*-values were obtained through data-peeking, which yields a true D = 1.223, which is very different from the estimated D = 1.472 in Table 7.

Finally, consider the (mis)fit of the estimated *p*-value distribution. Despite the considerable downward bias in heterogeneity estimate  $\hat{\tau}_{p_F}$ , the simulated *p*-value distribution is mostly well approximated by the expected *p*-value distribution, as indicated by the small values of the  $\chi^2$  statistic for *p*-values in 0-.00125. Hence, good fit again does not imply accurate parameter estimates. The misfit of the estimated distribution for *p*-values  $\leq .05$  is indicated by large  $\chi^2$ -values, particularly when the *p*-value distribution is not monotonically decreasing (which is the case for, e.g.,  $\mu = 0$ ).

To conclude, this simulation study showed that under true homogeneity both measures of data peeking can accurately signal both absence and presence of data peeking. However, under true heterogeneity, heterogeneity is underestimated and the performance of D breaks down, while results suggest that comparing estimates of average effect size, the first measure, may still accurately signal both the absence and presence of data peeking.

# 452 LIMATIONS AND CONCLUSION

Before concluding, some limitations of our method to collect *p*-values need to be addressed. First, 453 statcheck (Epskamp and Nuijten, 2015; Nuijten et al., 2015), the R package used to collect the 454 observed data, extracts all APA test results reported in the text of an article, but not those reported in 455 tables. Hence, our selection of results is potentially not representative of all reported results, but this 456 most likely does not affect results. Second, our analysis assumed that test statistics other than *p*-values 457 were accurately reported. If test statistics and degrees of freedom are incorrectly reported, recalculated 458 459 *p*-values are wrong as well. We identified some erroneous test statistics (e.g.,  $df_1 = 0$  and r > 1), but do not know how often these errors occur and how they may have affected our results. We assumed that 460 *p*-value errors were made due to the overemphasis on them in current day research. 461

In light of conflicting findings and interpretations, we aimed to provide final answers to the questions 462 (1) Does a bump or monotonic excess of p-values below .05 exist in psychology? and (2) Did evidence 463 for a bump increase over time in psychology? Answering these research questions may inform us on 464 the prevalence of QRPs and its development over time in psychology. Using statcheck, we extracted 465 and analyzed 258,050 test results conforming to APA-style across 30,710 articles from eight high impact 466 journals in psychology, and distinguished between results with inexactly reported *p*-values, exactly 467 reported p-values, and recalculated p-values. The basic idea underlying our analyses is that QRPs distort 468 the *p*-value distribution. We argued that only some QRPs yield an excess of *p*-values just below .05, and 469 show that QRPs sometimes yield a bump and sometimes only monotonic excess of *p*-values just below 470 .05. We used the Caliper test to test for a bump, and suggested two measures to examine monotic excess. 471 Starting with the existence of a bump in psychology, we drew the following conclusions. First, 472 inexactly reported p-values are not useful for analyses of p-value distributions. Second, a bump in exactly 473 reported *p*-values indeed exists in psychology journals DP, JAP, and JPSP. QRPs leading to just significant 474 *p*-values can explain these bumps, but we also cannot rule out the explanation that scientists in these 475 476 particular journals are more prone to exactly report *p*-values just below .05 (e.g., to emphasize they are really smaller than .05) than *p*-values considerably smaller than .05. Third, contradicting Leggett et al. 477 (2013), the bump and evidence of a bump in psychology did not increase over the years. Fourth, when 478

analyzing only the *exactly* reported *p*-values equal to .05, clear and direct evidence was obtained for the
QRP "incorrect rounding of *p*-value" (John et al., 2012). Evidence of this QRP, which contributed to
the bump in exactly reported *p*-values in psychology, was found in all psychology journals. Fifth, after
removing reporting errors and analyzing the *recalculated* reported *p*-values, evidence of a bump was
found only for JPSP. Again, this may have been caused by QRPs or by scientists being more prone to
report all test statistics when *p*-values are just below .05 than if they are considerable smaller than zero.

The conclusions obtained with the two measures investigating monotonic and non-monotic excess are 485 not satisfactory. First, performance of both measures is dependent on accurately recovering parameters 486 of the effect size distribution, which turned out to be difficult; estimates of effect size heterogeneity 487 and average effect size are highly correlated and unstable when based on only statistically significant 488 findings. Second, simulations show that one of the measures, D, does not accurately assess the QRP data 489 peeking when effect sizes are heterogeneous. Third, even though performance of the second measure 490 (i.e., difference between effect sizes based on contaminated and supposedly uncontaminated *p*-values) 491 is affected by estimation problems, it correctly signaled data peeking in the simulations. Fourth, when 492 493 applying the second measure to the observed distribution of significant *p*-values in psychology, the measure found evidence of monotonic excess of *p*-values; the average effect size estimate based on all 494 these p-values was 0, which seems very unrealistic, and suggests the use of QRPs in psychology leading 495 to *p*-values just below .05. 496

Notwithstanding the outcome of the second measure, suggesting QRPs that cause monotonic excess, 497 we do not consider it as direct evidence of such QRPs in psychology. Lakens (p.3; 2015) suggests that "it is 498 essential to use a model of p-value distributions before drawing conclusions about the underlying reasons 499 for specific distributions of *p*-values extracted from the scientific literature." We explicitly modeled the 500 effect size distribution and by using the degrees of freedom of test results also model the effect sizes 501 power and the *p*-value distribution. But we fear this is not and cannot be sufficient. First of all, we could 502 not accurately recover the effect size distribution under heterogeneity in our simulation study, even if all 503 assumptions of our model were met. This rendered measure D unfruitful when there is heterogeneity, 504 and severely limits the usefulness of the second measure that compares estimated average effect sizes. 505 Second, devising other models may yield other results and thereby other interpretations (Benjamini and 506 Hechtlinger, 2014; Goodman, 2014; Lakens, 2015a; de Winter and Dodou, 2015). 507

Results of all the aforementioned models are most likely not robust to violations of their assumptions. 508 For instance, we assume a normal distribution of true effect sizes. This assumption is surely violated, since 509 the reported *p*-values arise from a mixture of many different types of effects, such as very large effects 510 (manipulation checks), effects corresponding to main hypotheses, and zero effects ('control' variables). 511 Additionally, consider the QRPs themselves; we examined the effect of only one QRP, data peeking, in 512 one of its limited variants. Other QRPs exist that also increase the prevalence of *p*-values just below .05, 513 such as multiple operationalizations of a measure and selecting the first one to be significant. Other QRPs 514 even increase the frequency of very small p-values (van Aert et al., 2015). We deem it impossible to 515 exhaustively model QRPs and their effects, considering the difficulties we show for a single QRP that is 516 clearly defined. To conclude, we fear that Gelman and O'Rourke (2014) may be right when suggesting 517 that drawing conclusions with regard to any QRP based on modeling *p*-value distributions obtained from 518 automatically extracted results is unfruitful. 519

On the other hand, we do recommend modeling effect size and *p*-value distributions of results 520 that all intend to test the same hypothesis, to prevent contamination by irrelevant test results (Bishop 521 and Thompson, 2015; Simonsohn et al., 2015). Examples of methods that focus on similar results are 522 p-uniform (van Assen et al., 2015) and p-curve (Simonsohn et al., 2014), which model statistically 523 significant statistics pertaining to one specific effect and estimate the effect size based on these statistics 524 while correcting for publication bias. Further research should reveal if both methods can also be used 525 to detect and correct for *p*-hacking in the context of estimating one particular effect size. Preliminary 526 results suggest, however, that detection and correcting for *p*-hacking based on statistics alone is rather 527 challenging (van Aert et al., 2015). 528

## 529 **REFERENCES**

American Psychological Association (1983). *Publication manual of the American Psychological Associa- tion*. American Psychological Association, Washington, DC, 3rd edition.

- American Psychological Association (2001). *Publication manual of the American psychological associa- tion*. American Psychological Association, Washington, DC, 5th edition.
- American Psychological Association (2010). *Publication manual of the American Psychological Associa- tion*. American Psychological Association, Washington, DC, 6th edition.
- *tion.* American Psychological Association, Washington, DC, 6th edition.
   Armitage, P., McPherson, C. K., and Rowe, B. C. (1969). Repeated significance tests on accumulating
- data. Journal of the Royal Statistical Society. Series A, 132(2):235–244.
- Asendorpf, J. B., Conner, M., De Fruyt, F., De Houwer, J., Denissen, J. J. A., Fiedler, K., Fiedler, S.,
- <sup>539</sup> Funder, D. C., Kliegl, R., Nosek, B. A., Perugini, M., Roberts, B. W., Schmitt, M., van Aken, M. A. G.,
- Weber, H., and Wicherts, J. M. (2013). Recommendations for increasing replicability in psychology.
   *European journal of personality*, 27(2):108–119.
- Bakker, M. and Wicherts, J. M. (2011). The (mis)reporting of statistical results in psychology journals.
   *Behavior research methods*, 43(3):666–678.
- <sup>544</sup> Bakker, M. and Wicherts, J. M. (2014). Outlier removal, sum scores, and the inflation of the type I error
- rate in independent samples t tests: the power of alternatives and recommendations. *Psychological methods*, 19(3):409–427.
- Benjamini, Y. and Hechtlinger, Y. (2014). Discussion: An estimate of the science-wise false discovery
   rate and applications to top medical journals by jager and leek. *Biostatistics*, 15(1):13–16.
- Bishop, D. V. and Thompson, P. A. (2015). Problems in using text-mining and p-curve analysis to detect
   rate of p-hacking. Technical Report e1550, PeerJ PrePrints.
- <sup>551</sup> Chamberlain, S., Boettiger, C., and Ram, K. (2015). rplos: Interface to the search 'API' for 'PLoS' <sup>552</sup> journals.
- de Winter, J. C. and Dodou, D. (2015). A surge of p-values between 0.041 and 0.049 in recent decades (but negative results are increasing rapidly too). *PeerJ*, 3:e733.
- <sup>555</sup> Epskamp, S. and Nuijten, M. (2015). statcheck: Extract statistics from articles and recompute p-values.
- Ferguson, C. J. (2015). 'everybody knows psychology is not a real science': Public perceptions of psychology and how we can improve our relationship with policymakers, the scientific community, and
- the general public. *The American psychologist*, 70(6):527–542.
- Franco, A., Malhotra, N., and Simonovits, G. (2014). Publication bias in the social sciences: Unlocking
   the file drawer. *Science*, 345(6203):1502–1505.
- García-Berthou, E. and Alcaraz, C. (2004). Incongruence between test statistics and P values in medical papers. *BMC medical research methodology*, 4:13.
- Gelman, A. and O'Rourke, K. (2014). Discussion: Difficulties in making inferences about scientific truth
   from distributions of published p-values. *Biostatistics*, 15(1):18–23.
- Gerber, A. S., Malhotra, N., Dowling, C. M., and Doherty, D. (2010). Publication bias in two political behavior literatures. *American Politics Research*, 38(4):591–613.
- Ginsel, B., Aggarwal, A., Xuan, W., and Harris, I. (2015). The distribution of probability values in medical abstracts: an observational study. *BMC research notes*, 8(1):721.
- Goodman, S. N. (2014). Discussion: An estimate of the science-wise false discovery rate and application
- to the top medical literature. *Biostatistics*, 15(1):23–7.
- <sup>571</sup> Hartgerink, C. H. J. (2015). Reanalyzing head et al. (2015): No widespread p-hacking after all?
- Head, M. L., Holman, L., Lanfear, R., Kahn, A. T., and Jennions, M. D. (2015). The extent and
   consequences of p-hacking in science. *PLoS biology*, 13(3):e1002106.
- <sup>574</sup> Ioannidis, J. P. A. (2014). Discussion: Why "an estimate of the science-wise false discovery rate and <sup>575</sup> application to the top medical literature" is false. *Biostatistics*, 15(1):28–36.
- Jager, L. R. and Leek, J. T. (2014). An estimate of the science-wise false discovery rate and application to the top medical literature. *Biostatistics*, 15(1):1–12.
- John, L. K., Loewenstein, G., and Prelec, D. (2012). Measuring the prevalence of questionable research practices with incentives for truth telling. *Psychological science*, 23(5):524–532.
- Krawczyk, M. (2015). The search for significance: A few peculiarities in the distribution of p values in
   experimental psychology literature. *PloS one*, 10(6):e0127872.
- Kühberger, A., Fritz, A., and Scherndl, T. (2014). Publication bias in psychology: A diagnosis based on
   the correlation between effect size and sample size. *PloS one*, 9(9):e105825.
- Lakens, D. (2015a). On the challenges of drawing conclusions from p-values just below 0.05. *PeerJ*, 3:e1142.
- Lakens, D. (2015b). What p-hacking really looks like: A comment on masicampo and LaLande (2012).

- <sup>587</sup> *Quarterly journal of experimental psychology*, 68(4):829–832.
- Leggett, N. C., Thomas, N. A., Loetscher, T., and Nicholls, M. E. R. (2013). The life of p: "just significant" results are on the rise. *Quarterly journal of experimental psychology*, 66(12):2303–2309.
- <sup>590</sup> Masicampo, E. J. and Lalande, D. R. (2012). A peculiar prevalence of p values just below .05. *Quarterly* <sup>591</sup> *journal of experimental psychology*, 65(11):2271–2279.
- <sup>592</sup> Nuijten, M. B., Hartgerink, C. H. J., Van Assen, M. A. L. M., Epskamp, S., and Wicherts, J. M. (2015).
- <sup>593</sup> The prevalence of statistical reporting errors in psychology (1985-2013). *Behavior Research Methods*.
- <sup>594</sup> Panel on Scientific Responsibility and the Conduct of Research (1992). *Responsible science, volume I:*
- *Ensuring the integrity of the research process*. National Academies Press (US), Washington, DC.
- Pashler, H. and Wagenmakers, E.-J. (2012). Editors' introduction to the special section on replicability in
   psychological science: A crisis of confidence? *Perspectives on Psychological Science*, 7(6):528–530.
- Pautasso, M. (2010). Worsening file-drawer problem in the abstracts of natural, medical and social science
   databases. *Scientometrics*, 85(1):193–202.
- Ridley, J., Kolm, N., Freckelton, R. P., and Gage, M. J. G. (2007). An unexpected influence of widely
   used significance thresholds on the distribution of reported p-values. *Journal of evolutionary biology*,
   20(3):1082–1089.
- <sup>602</sup> 20(3):1082–1089. <sup>603</sup> Simmons, J. P., Nelson, L. D., and Simonsohn, U. (2011). False-positive psychology: Undisclosed
- flexibility in data collection and analysis allows presenting anything as significant. *Psychological science*, 22(11):1359–1366.
- Simonsohn, U., Nelson, L. D., and Simmons, J. P. (2014). P-curve: A key to the file-drawer. *Journal of experimental psychology: General*, 143(2):534–547.
- Simonsohn, U., Simmons, J. P., and Nelson, L. D. (2015). Better p-curves. *Journal of Experimental Psychology: General*, 144:1146–1152.
- <sup>610</sup> Ulrich, R. and Miller, J. (2015). p-hacking by post hoc selection with multiple opportunities: Detectability
- <sup>611</sup> by skewness test?: Comment on simonsohn, nelson, and simmons (2014). *Journal of experimental* <sup>612</sup> *psychology. General*, 144(6):1137–1145.
- van Aert, R. C. M., Wicherts, J. M., and van Assen, M. A. L. M. (2015). Conducting meta-analyses
- on p-values: Reservations and recommendations for applying p-uniform and p-curve. *Manuscript*
- 615 *submitted for publication.*
- van Assen, M. A. L. M., van Aert, R. C. M., and Wicherts, J. M. (2015). Meta-analysis using effect size
   distributions of only statistically significant studies. *Psychological methods*, 20:293–309.
- Veldkamp, C. L. S., Nuijten, M. B., Dominguez-Alvarez, L., van Assen, M. A. L. M., and Wicherts, J. M.
   (2014). Statistical reporting errors and collaboration on statistical analyses in psychological science.
- 620 *PloS one*, 9(12):e114876.
- Vermeulen, I., Beukeboom, C. J., Batenburg, A., Avramiea, A., Stoyanov, D., de Velde, B. v., and
- Oegema, D. (2015). Blinded by the light: How a focus on statistical "significance" may cause p-value misreporting and an excess of p-values just below .05 in communication science. *Communication*
- 624 *methods and measures*, 9(4):253–279.
- Wagenmakers, E.-J. (2007). A practical solution to the pervasive problems of p values. *Psychonomic bulletin & review*, 14(5):779–804.
- <sup>627</sup> Wagenmakers, E.-J., Wetzels, R., Borsboom, D., van der Maas, H. L. J., and Kievit, R. A. (2012). An
- agenda for purely confirmatory research. *Perspectives on psychological science: a journal of the*
- Association for Psychological Science, 7(6):632–638.