Willingness to share research data is related to the strength of the evidence and the quality of reporting of statistical results

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Background: The widespread reluctance to share published research data is often hypothesized to be due to the authors’ fear that reanalysis may expose errors in their work or may produce conclusions that contradict their own. However, these hypotheses have not previously been studied systematically.

Methods and Findings: We related the reluctance to share research data for reanalysis to 1148 statistically significant results reported in 49 papers published in two major psychology journals. We found the reluctance to share data to be associated with weaker evidence (against the null hypothesis of no effect) and a higher prevalence of apparent errors in the reporting of statistical results. The unwillingness to share data was particularly clear when reporting errors had a bearing on statistical significance.

Conclusions: Our findings on the basis of psychological papers suggest that statistical results are particularly hard to verify when reanalysis is more likely to lead to contrasting conclusions. This highlights the importance of establishing mandatory data archiving policies.
**Introduction**

Statistical analyses of research data are quite error prone [1,2,3], accounts of statistical results may be inaccurate [4], and decisions that researchers make during the analytical phase of a study may lean towards the goal of achieving a preferred (significant) result [5,6,7,8]. For these and other (ethical) reasons [9], many scientific journals like *PLoS ONE* [10] and professional organizations such as the *American Psychological Association* (APA) [11] have clear policies concerning the sharing of data after research results are published. For instance, upon acceptance for publication of a paper in one of the over 50 peer-reviewed journals published by the APA, authors sign a contract that they will make available data to peers who wish to reanalyze their data to verify the substantive claims put forth in the paper. Nonetheless, the replication of statistical analyses in published psychological research is hampered by psychologists’ pervasive reluctance to share their raw data [1,12]. In a large-scale study Wicherts et al. [12] found that 73% of psychologists publishing in four top APA journals defied APA guidelines by not sharing their data for reanalysis. The unwillingness to share data of published research has been documented in a number of fields [13,14,15,16,17,18,19,20] and is often ascribed in part to the fear among authors that independent reanalysis will expose statistical or analytical errors in their work [21] and will produce conclusions that differ from theirs [22]. However, no published research to date has addressed whether this rather bleak scenario has a bearing on reality.

Here we study whether researchers’ willingness to share data for reanalysis is associated with the strength of the evidence (defined as the statistical evidence against the null hypothesis of no effect) and the quality of the reporting of statistical results (defined in terms of the prevalence of
inconsistencies in reported statistical results). To this end, we followed-up on Wicherts et al.’s requests for data [12] by comparing statistical results in papers from which data were either shared or not, and to check for errors in the reporting of p-values in both types of papers.

**Methods**

In the summer of 2005, Wicherts and colleagues [12] contacted the corresponding authors of 141 papers that were published in the second half of 2004 in one of four high-ranked journals published by the APA: *Journal of Personality and Social Psychology (JPSP), Developmental Psychology (DP), Journal of Consulting and Clinical Psychology (JCCP)*, and *Journal of Experimental Psychology: Learning, Memory, and Cognition (JEP:LMC)*. The data were requested to determine the effects of outliers on statistical outcomes (see Supporting Information for details). Although all corresponding authors had signed a statement that they would share their data for such verification purposes [11], most authors failed to do so. In the current study, we related the willingness to share data from 49 papers published in *JPSP* or *JEP:LMC* to two relevant characteristics of the statistical outcomes reported in the papers, namely the internal consistency of the statistical results and the distribution of significantly reported ($p < .05$) p-values. We restricted the attention to *JPSP* and *JEP:LMC*, because (1) authors in these journals were more willing to share data than authors in the other journals from which Wicherts et al. requested data, (2) no corresponding authors in these two journals declined to share data, because they were part of an ongoing project or because of propriety rights or ethical considerations, and (3) studies in these two journals were fairly homogeneous in terms of analysis and design (mostly lab experiments). We also restricted our
attention to results from null-hypothesis significance testing (NHST) [23]. This procedure is not without its critics [24,25], but remains to be used extensively in psychology [26] and related fields. NHST provides p-values that, if smaller than alpha = .05, are considered by many researchers [27,28] and reviewers [29] to lend support to the hypothesized effects. Psychological research data are often amenable to alternative methods of analysis [6,22,30] that may affect what can be concluded from them (at least within the rules of NHST). The specifics of the analysis will typically have more relevance when statistical results are nearly significant at the alpha = .05 level. Put differently, smaller p-values provide stronger evidence against the null hypothesis of no effect [31]. The strength of the evidence based on Bayes factors from Bayesian t-tests has been found to be strongly inversely related to the p-values of traditional t-tests [32]. If the strength of the evidence so defined plays part in the willingness to share data, then it is to be expected that p-values in papers from which data were not shared lie closer to .05. Because reported p-values are often inconsistent with the given test statistics and degrees of freedom [33], we also check for errors in reporting of statistical results.

**Data Retrieval**

We extracted from the papers all the t, F, and χ² test statistics associated with NHST, the given degrees of freedom (e.g., F(2,24) = 3.41), the sidedness of tests (1- or 2-tailed), and the reported exact p-value (e.g., p = .03) or the reported level of significance (e.g., p < .05). We considered these tests because these are the most common test statistics of NHST in psychology. Although it was infeasible to determine for each test whether it was in line with the researchers’ predictions, NHST is typically used for the purpose of rejecting the
null hypothesis. We did not consider test statistics that were not associated with NHST (e.g., model fitting or Bayesian analyses). We only included test results that were uniquely reported, complete (i.e., test statistic, degrees of freedom, and p-value were reported), and that were reported as being significant (i.e., \( p < .05 \)) in the main text or in tables in the results sections. T-tests were considered 2-tailed, unless stated otherwise. The exact p-values were computed on the basis of the given test statistic and DF(s) in Microsoft Excel 2008 for Mac, version 12.1.0. A further four papers published in the two journals from which data were requested in 2005 were not included in the follow-up because they did not involve NHST or did not contain significant results on the basis of \( t \), \( F \), of \( \chi^2 \) tests.

Five undergraduates, who were unaware from which papers data were shared also independently retrieved a total of 495 statistics and DFs. We compared these 495 statistics to ours and determined that the accuracy rate in our own data was 99.4%. The three minor errors in our data retrieval were corrected but proved trivial.

**Detecting Reporting Errors**

Inconsistencies between reported p-values (or ranges) and p-values recalculated from the retrieved statistics were detected automatically in Excel as follows. The recomputed p-value was first rounded to the same number of digits as was used in the reported p-value (range). Subsequently, an IF-statement automatically checked for consistency. Next, we determined by hand whether reporting errors were not due to possible errors in extraction (none were found) or to rounding. For example, a test result such as “\( t(15) = 2.3; \ p = 0.034 \)” could have arisen from test statistic that could possibly range from 2.25 to 2.35. Consequently, the correct p-value could range from .033 to .040 and so the
reported value was not seen as inconsistent, although the recomputed p-value is .0362. In the analyses of the p-value distributions, we used the nearest next decimal that attained consistency for these correctly rounded cases (i.e., 2.34 in the example), but used the p-value on the basis of the reported test statistic in other cases. We checked whether over-reported p-values were corrected upwards via procedures like Bonferroni’s or Huyn-Feldt’s, but did not use these corrections in analyzing p-value distributions. As some of the inconsistencies may have arisen from the use of one-sided testing, we made additional searches of the text for explicit references thereof. In one instance, an F-test result was reported explicitly as a one-sided test, but because this result was equivalent to a one-sided t-test we did not consider it erroneous (as suggested by an independent reviewer). As a final check, the three authors independently verified all 49 inconsistencies on the basis of the papers. All documented errors are available upon request.

The use of this method previously revealed quite high errors rates in the reporting of p-values in papers published in Nature, the British Medical Journal [4], and two psychiatry journals [34]. In a recent study covering a fairly representative sample of 281 psychology papers [33], roughly 50% of the papers that involved NHST were found to include at least one such reporting error. As discussed elsewhere [33], likely causes include (1) errors in the retrieval and copying of test statistics, degrees of freedom, and/or p-value (e.g., reporting the total DF instead of the error DF of an F test), (2) incorrect rounding of last decimal (e.g., \( p = .059 \) reported as \( p = .05 \)), (3) the use of one-tailed tests without mentioning this, (4) incorrect use of tests (e.g., dividing the p value of an F or \( \chi^2 \) test by two to report a one-sided p value, whereas the F or \( \chi^2 \) test is already a one-sided test), (5) confusion of = with < (e.g., \( p = .012 \))
reported as $p < .01$), and (6) copy-editing errors (e.g., a failure to alter relevant numbers after the use of “copy-paste” in writing the paper). Although many inconsistencies between reported and recomputed $p$-values in Bakker and Wicherts’ study were minor, roughly 15% of the papers contained at least one result that was presented as being statistically significant ($p < .05$), but that proved, upon recalculation, not to be significant ($p > .05$). Such serious errors in the reporting of results increase the desirability to have the data available for reanalysis.

**Ethical Considerations**

This study has been approved by the Ethics Committee of the Psychology Department of the University of Amsterdam. In light of the purpose of our study, we could not ask the corresponding authors for their informed consent. The Ethics Committee exempted the use of informed consent because all corresponding authors had signed APA publication forms related to data sharing and in light of Article 8.05 of the Ethical Principles of the APA. The documented errors are based on publically available papers and so are considered archival material. The sharing or non-sharing of data is considered to be an unobtrusive observation of professional behavior of the corresponding authors that should not create distress on their behalf, provided that anonymity is assured. To protect the identity of corresponding authors, we are not allowed to make public who did or did not share data with Wicherts et al. However, this information is available upon request to allow others to verify our results through reanalysis. The problems that we highlight are general, and our results should not be used to question the academic integrity of individual researchers. The analyses we report
here were all conducted independently by at least two of us on the basis of the data that all of us have in our possession.

Results

Responses to Data Requests

Of the 49 corresponding authors, 21 (42.9%) had shared some data with Wicherts et al. Thirteen corresponding authors (26.5%) failed to respond to the request or any of the two reminders. Three corresponding authors (6.1%) refused to share data either because the data were lost or because they lacked time to retrieve the data and write a codebook. Twelve corresponding authors (24.5%) promised to share data at a later date, but have not done so in the past six years (we did not follow up on it). These authors commonly indicated that the data were not readily available or that they first needed to write a codebook.

Errors in the Reporting of Statistical Results

The 49 papers contained a total of 1148 test statistics that were presented as significant at $p < .05$. Table 1 presents for each paper the number of significantly reported test results, the number of misreporting errors, and the median and average of all genuinely significant $p$-values (as based on the recalculated values). Forty-nine of these statistics (4.3%) were inconsistent with the reported (range of) $p$-values. In forty-seven of the inconsistent results (95.9%), the reported $p$-value (range) was smaller than the recalculated $p$-value. Figure 1 gives the origin of three types of reporting errors. Although 51.1% (587) of the tests statistics were from papers from which no data were shared, most incorrectly reported $p$-values (36 out of 49; 73.5%) originated from these papers. These errors include quite small ones (e.g., $p = .0002$
reported as $p < .0001$). Twenty-eight of the 32 p-values (87.5%) that were incorrectly reported at the level of the 2nd decimal (e.g., $p = .02$ reported as $p < .01$) were from papers from which no data were shared. Negative binomial regressions (Table 2) that accounted for the number of test statistics and the average p-values in each paper (see below) showed that reluctance to share data was predictive of the prevalence of both types of reporting errors.

We came across a total of ten cases (from seven papers) in which the recomputed p-value was above .05, whilst the result was presented as being significant ($p < .05$). None of the authors of these papers had shared data with Wicherts et al. As a negative binomial regression is infeasible with these data, we tested this relation at the level of papers (includes serious error(s) versus shared) with a 2-by-2 Fisher exact test: $p = .015$ (2-tailed). So the reluctance to share data was particularly evident when the reporting errors concern statistical significance. This corroborates an earlier finding that it took authors considerably longer to respond to queries for data when the inconsistency in their reported results had a bearing on the significance of their results [33].

**Strength of Evidence (against the Null Hypothesis)**

P-values from NHST are traditionally interpreted as the strength of the evidence against the null hypothesis of no effect [31]. From the distribution of significant p-values across the two groups of papers in Figure 2, it is clear that higher p-values, like those in the interval between .03 and .05 (which have a low chance of occurring regardless of actual effect sizes [35]), were indeed more common in papers from which no data were shared (16.7%) than in the other papers (9.1%). The individual statistical results are statistically dependent within papers in rather intractable ways, and so we computed the mean of p-values of
all genuinely significant results within each paper. This variable was indeed significantly higher in the 28 papers from which the data were not shared ($M = .0124, SD = .0074, Median = .0114$ vs. $M = .0079, SD = .0046, Median = .0073$, Cohen’s $d = .72$): Wilcoxon’s $W = 413, Z = 2.26, p = .024$ (2-tailed). The use of the median of $p$-values per paper provided similar results ($Z = 2.14, p = .032$).

We also conducted a bootstrap analysis to test this difference between shared and non-shared papers on the basis of individual $p$-values as clustered in the papers. In this analysis, we determined on the basis of 100,000 replications the null distribution of Wilcoxon’s $W$ test for the 1138 statistically dependent $p$-values that were smaller than .05. To this end, we randomly assigned each paper (and hence all $p$-values in it) to either the shared or non-shared category (on the basis of the base rate of $p = 21/49$), and repeated this process 100,000 times to get an empirical null distribution for $W$ on the basis of our data. The $W$ statistic computed on the basis of actual difference between shared and non-shared gave a $p$-value of .0298 (2-tailed) in this bootstrapped null distribution. Hence, the analyses of individual $p$-values corroborated that $p$-values were significantly higher in papers from which data were not shared.

**Discussion**

In this sample of psychology papers, the authors’ reluctance to share data was associated with more errors in reporting of statistical results and with relatively weaker evidence (against the null hypothesis). The documented errors are arguably the tip of the iceberg of potential errors and biases in statistical analyses and the reporting of statistical results. It is rather disconcerting that roughly 50% of published papers in psychology contain reporting errors [33] and
that the unwillingness to share data was most pronounced when the errors concerned statistical significance. Although our results are consistent with the notion that the reluctance to share data is generated by the author’s fear that reanalysis will expose errors and lead to opposing views on the results, our results are correlational in nature and so they are open to alternative interpretations. Although the two groups of papers are similar in terms of research fields and designs, it is possible that they differ in other regards. Notably, statistically rigorous researchers may archive their data better and may be more attentive towards statistical power than less statistically rigorous researchers. If so, more statistically rigorous researchers will more promptly share their data, conduct more powerful tests, and so report lower p-values. However, a check of the cell sizes in both categories of papers (see Supplementary Information) did not suggest that statistical power was systematically higher in studies from which data were shared.

The association between reporting errors and sharing of data after results are published may also reflect differences in the rigor with which researchers manage their data. Rigorously working researchers may simply commit fewer reporting errors because they manage and archive their data more diligently. A recent survey among 192 Dutch psychological researchers highlighted a rather poor practice of data archiving in psychology [36]. When asked whether they archived their research data, only a third of the psychologists responded positively. This is remarkable in light of guidelines of the APA [11] that stipulate that data should be retained a minimum of five years after publication of the study. Even among those psychologists who indicated that they “archive” their data, most did not follow proper archiving standards (e.g., by keeping code books and writing meta-data [37]), but simply stored data on their own (current)
computer (32%), on CDs/DVDs (18%), or on the shelf (20%). Haphazard data management is documented in a number of scientific fields [37,38,39], may result in errors in analyzing and reporting of results, and obviously impedes the sharing of data after results are published.

Regardless of the underlying processes, the results on the basis of the current papers imply that it is most difficult to verify published statistical results when these are contentious. We focused here on NHST within two psychology journals and so it is desirable to replicate our results in other fields and in the context of alternative statistical approaches. However, it is likely that similar problems play a role in the widespread reluctance to share data in other scientific fields [13,14,15,16,17,18,19,20]. Because existing guidelines on data sharing offer little promise for improvement [40], progress in psychological science and related fields would benefit from having research data itself be part of the process of replication [15,16], notably by the establishment by journals, professional organizations, and granting bodies of mandatory data archiving policies.

More stringent policies concerning data archiving will not only facilitate verification of analyses and corrections of the scientific record, but also improve the quality of reporting of statistical results. Changing policies require better educational training in data management and data archiving, which is currently suboptimal in many fields [36,37,38,39]. On the other hand, technical capabilities for storage are already available. For instance, several trial registers in the medical sciences (like clinicaltrials.gov) enable storage of research data. Rigorous archiving of data involves documentation of variables, meta-data, saving data files in formats that are robust (e.g., ASCII files), and submitting
files to repositories that already require these standards. Best practices in conducting analyses and reporting statistical results involve, for instance, that all co-authors hold copies of the data, and that at least two of the authors independently run all the analyses (as we did in this study). Such double-checks and the possibility for others to independently verify results later should go a long way in dealing with human factors in the conduct of statistical analyses and the reporting of results.

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References

1. Wolins L (1962) Responsibility for raw data. American Psychologist 17: 657-658.

2. Murphy JR (2004) Statistical errors in immunologic research. Journal of Allergy and Clinical Immunology 114: 1259-1264.

3. Strasak AM, Zaman Q, Marinell G, Pfeiffer KP, Ulmer H (2007) The use of statistics in medical research: A comparison of The New England Journal of Medicine and Nature Medicine. American Statistician 61: 47-55.

4. Garcia-Berthou E, Alcaraz C (2004) Incongruence between test statistics and P values in medical papers. BMC Medical Research Methodology 4: 13.

5. Barber TX (1976) Pitfalls in Human Research: Ten Pivotal Points New York: Pergamon Press.

6. Friedlander F (1964) Type I and Type II bias. American Psychologist 19: 198-199.

7. Ioannidis JPA (2005) Why most published research findings are false. Plos Medicine 2: e124.

8. Mahoney MJ (1976) Scientist as Subject: The Psychological Imperative. Cambridge, MA, US: Ballinger Publishing Company.

9. Board on Life Sciences National Research Council (2003) Sharing publication-related data and materials: Responsibilities of authorship in the life sciences. Washington, DC: National Academies Press.

10. PLoS ONE (2011) PLoS Editorial and Publishing Policies. 7. Sharing of Materials, Methods, and Data. Available from: http://www.plosone.org/static/policies.action#sharing. Accessed August 1, 2011.
11. American Psychological Association (2010) Publication Manual of the American Psychological Association. Sixth Edition. Washington, DC: American Psychological Association.

12. Wicherts JM, Borsboom D, Kats J, Molenaar D (2006) The poor availability of psychological research data for reanalysis. American Psychologist 61: 726-728.

13. Savage CJ, Vickers AJ (2009) Empirical study of data sharing by authors publishing in PLoS journals. PLoS One 4: e7078.

14. McCullough BD, Vinod HD (2003) Verifying the solution from a nonlinear solver: A case study. American Economic Review 93: 873-892.

15. Firebaugh G (2007) Replication data sets and favored-hypothesis bias. Sociological Methods & Research 36: 200-209.

16. Freese J (2007) Replication standards quantitative social science - Why not sociology. Sociological Methods & Research 36: 153-172.

17. Piwowar HA, Day RS, Fridsma DB (2007) Sharing detailed research data is associated with increased citation rate. PLoS One 2: e308.

18. Reidpath DD, Allotey PA (2001) Data sharing in medical research: An empirical investigation. Bioethics 15: 125-134.

19. Campbell EG, Clarridge BR, Gokhale M, Birenbaum L, Hilgartner S, et al. (2002) Data withholding in academic genetics: Evidence from a national survey. Journal of the American Medical Association 287: 473-480.

20. Kyzas PA, Loizou KT, Ioannidis JPA (2005) Selective reporting biases in cancer prognostic factor studies Journal of the National Cancer Institute 97: 1043-1055.

21. Ceci SJ, Walker E (1983) Private archives and public needs. American Psychologist 38: 414-423.
22. Nature (2006) A fair share. Nature 444: 653-654.

23. Nickerson RS (2000) Null hypothesis significance testing: A review of an old and continuing controversy. Psychological Methods 5: 241-301.

24. Cohen J (1994) The earth is round (P less-than.05). American Psychologist 49: 997-1003.

25. Wagenmakers E-J (2007) A practical solution to the pervasive problems of p values Psychonomic Bulletin & Review 14: 779-804.

26. Hubbard R, Ryan PA (2000) The historical growth of statistical significance testing in psychology-and its future prospects. Educational and Psychological Measurement 60: 661-681.

27. Hoekstra R, Finch S, Kiers HAL, Johnson A (2006) Probability as certainty: Dichotomous thinking and the misuse of p values. Psychonomic Bulletin & Review 13: 1033-1037.

28. Rosenthal R, Gaito J (1963) The interpretation of levels of significance by psychological researchers. Journal of Psychology 55: 33-38.

29. Mahoney MJ (1977) Publication prejudices: An experimental study of confirmatory bias in the peer review system. Cognitive Therapy and Research 1: 161-175.

30. Blanton H, Jaccard J, Klick J, Mellers B, Mitchell G, et al. (2009) Strong claims and weak evidence: Reassessing the predictive validity of the IAT. Journal of Applied Psychology 94: 567-582.

31. Fisher RA (1958) Statistical Methods for Research Workers. New York: Hafner.

32. Wetzels R, Matzke D, Lee MD, Rouder JN, Iverson GJ, et al. (2011) Statistical evidence in experimental psychology: An empirical comparison using 855 t tests Perspectives on Psychological Science 6: 291-298.
33. Bakker M, Wicherts JM (2011) The (mis)reporting of statistical results in psychology. Behavior Research Methods 43: 666-678.

34. Berle D, Starcevic V (2007) Inconsistencies between reported test statistics and p-values in two psychiatry journals. International Journal of Methods in Psychiatric Research 16: 202-207.

35. Sellke T, Bayarri MJ, Berger JO (2001) Calibration of p values for testing precise null hypotheses. The American Statistician 55: 62-71.

36. Voorbrood C (2010) Archivering, beschikbaarstelling en hergebruik van onderzoeksdata in de psychologie [Archiving, sharing, and reusing of psychological research data]. The Hague, The Netherlands: Data Archiving and Networked Services.

37. Jubb M (2008) To share or not to share: Publication and quality assurance of research data outputs. London, UK: Research Information Network.

38. McCullough BD, McGeary KA, Harrison TD (2006) Lessons from the JMCB Archive. Journal of Money, Credit, and Banking 38: 1093-1107.

39. Nature (2011) Devil in the details. To ensure their results are reproducible, analysts should show their workings. Nature 470: 305-306.

40. Wicherts JM, Bakker M (2009) Sharing: guidelines go one step forwards, two steps back. Nature 461: 1053.
Figure 1

Distribution of reporting errors per paper for papers from which data were shared and from which no data were shared.

Distribution of the number of errors in the reporting of p-values for 28 papers from which the data were not shared (left column) and 21 from which the data were shared (right column) for all misreporting errors (upper row), larger misreporting errors at the 2nd decimal (middle row), and misreporting errors that concerned statistical significance (p < .05; bottom row).
Figure 2

Distribution of p-values reported as being significant in papers from which data were shared or not.

Distribution of p-values reported as being significant (at p < .05) in 21 papers from which data were shared (N = 561; in black) and in 28 papers from which data were not shared (N = 587; in grey), showing that p-values often lie closer to the typical boundary of significance when data are not shared for reanalysis. Frequencies of reporting errors (as given above the bars) reflect higher error prevalence in papers from which no data were shared.
## Table 1

**Summary statistics for the 49 papers**

| Journal | DOI | Pageno. | No. of stats. | Mean of ps | Median of ps | Reporting. errors |
|---------|-----|---------|--------------|-----------|--------------|--------------------|
| jep:lmc | 10.1037/0278-7393.30.5.947 | 947–959 | 7 | 0.006636 | 0.00295 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.969 | 969–987 | 13 | 0.027302 | 0.02936 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.988 | 988–1001 | 33 | 0.010325 | 0.00482 | 3 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1002 | 1002–1011 | 25 | 0.004257 | 0.00001 | 1 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1012 | 1012–1025 | 83 | 0.003054 | 0.00000 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1026 | 1026–1044 | 30 | 0.007286 | 0.00189 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1045 | 1045–1064 | 19 | 0.005587 | 0.00073 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1065 | 1065–1081 | 22 | 0.001672 | 0.00009 | 3 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1082 | 1082–1092 | 9 | 0.001089 | 0.00010 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1093 | 1093–1105 | 21 | 0.011132 | 0.00115 | 1 1 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1106 | 1106–1118 | 16 | 0.002213 | 0.00001 | 2 2 1 |
| jep:lmc | 10.1037/0278-7393.30.5.1119 | 1119–1130 | 10 | 0.007128 | 0.00095 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.5.1131 | 1131–1142 | 21 | 0.003256 | 0.00098 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1147 | 1147–1166 | 8 | 0.008461 | 0.00036 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1167 | 1167–1175 | 8 | 0.011841 | 0.00231 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1176 | 1176–1195 | 32 | 0.005418 | 0.00006 | 1 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1196 | 1196–1210 | 37 | 0.004050 | 0.00000 | 1 1 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1211 | 1211–1218 | 11 | 0.019460 | 0.01967 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1219 | 1219–1234 | 39 | 0.016008 | 0.01084 | 7 6 1 |
| jep:lmc | 10.1037/0278-7393.30.6.1235 | 1235–1251 | 23 | 0.004993 | 0.00096 | 1 1 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1252 | 1252–1270 | 46 | 0.010496 | 0.00058 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1271 | 1271–1278 | 20 | 0.002645 | 0.00001 | 1 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1290 | 1290–1301 | 35 | 0.013469 | 0.00475 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1302 | 1302–1321 | 30 | 0.013727 | 0.00680 | 0 0 0 |
| jep:lmc | 10.1037/0278-7393.30.6.1322 | 1322–1337 | 37 | 0.006148 | 0.00094 | 0 0 0 |
| jpsp | 10.1037/0022-3514.87.5.557 | 557–572 | 33 | 0.016946 | 0.01104 | 1 1 0 |
| jpsp | 10.1037/0022-3514.87.5.573 | 573–585 | 15 | 0.011696 | 0.00597 | 1 0 0 |
| jpsp | 10.1037/0022-3514.87.5.586 | 586–598 | 21 | 0.019989 | 0.01519 | 4 4 3 |
| jpsp | 10.1037/0022-3514.87.5.599 | 599–614 | 24 | 0.009036 | 0.00263 | 0 0 0 |
| jpsp | 10.1037/0022-3514.87.5.615 | 615–630 | 27 | 0.003605 | 0.00000 | 3 | 0 | 0 |
|------|---------------------------|--------|----|-----------|----------|---|---|---|
| jpsp | 10.1037/0022-3514.87.5.631 | 631–648 | 6  | 0.008074 | 0.00385 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.5.649 | 649–664 | 16 | 0.012216 | 0.00510 | 4 | 4 | 0 |
| jpsp | 10.1037/0022-3514.87.5.665 | 665–683 | 23 | 0.016715 | 0.00179 | 2 | 1 | 1 |
| jpsp | 10.1037/0022-3514.87.6.733 | 733–749 | 24 | 0.023442 | 0.00000 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.750 | 750–762 | 5  | 0.000002 | 0.00000 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.763 | 763–778 | 29 | 0.007420 | 0.00000 | 1 | 1 | 0 |
| jpsp | 10.1037/0022-3514.87.6.779 | 779–795 | 9  | 0.025925 | 0.03231 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.817 | 817–831 | 20 | 0.007695 | 0.00011 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.832 | 832–844 | 8  | 0.021422 | 0.02079 | 4 | 4 | 1 |
| jpsp | 10.1037/0022-3514.87.6.845 | 845–859 | 48 | 0.009394 | 0.00380 | 2 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.860 | 860–875 | 28 | 0.019047 | 0.01104 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.876 | 876–893 | 27 | 0.011934 | 0.00598 | 1 | 1 | 1 |
| jpsp | 10.1037/0022-3514.87.6.894 | 894–912 | 8  | 0.009142 | 0.00092 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.913 | 913–925 | 7  | 0.018208 | 0.00783 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.926 | 926–939 | 9  | 0.011442 | 0.01224 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.940 | 940–956 | 36 | 0.009620 | 0.00314 | 2 | 2 | 0 |
| jpsp | 10.1037/0022-3514.87.6.957 | 957–973 | 45 | 0.006310 | 0.00020 | 0 | 0 | 0 |
| jpsp | 10.1037/0022-3514.87.6.974 | 974–990 | 30 | 0.018801 | 0.01527 | 1 | 1 | 0 |

Note: Ethical considerations preclude the inclusion of "shared vs. non-shared" in this table, but this information is available upon request. JEP:LMC vol. 30, JPSP vol. 87. Correlational design; mixed correlational/experimental design, remaining papers involve experimental designs.
### Table 2

**Results of negative binomial regressions of the number of reporting errors per paper**

| Predictor                                      | Parameter (SE) | Wald $\chi^2$ (DF=1) | p     |
|------------------------------------------------|----------------|----------------------|-------|
| All reporting errors (range: 0-7)              |                |                      |       |
| (Intercept)                                    | -2.76 (1.30)   | 4.53                 | .033  |
| Data shared (1) or not (0)                     | -0.83 (0.38)   | 4.84                 | .028  |
| Square root (Average of p-values)              | 4.39 (6.13)    | 0.51                 | .473  |
| Log (No. of test statistics)                   | 0.85 (0.41)    | 4.19                 | .041  |
| Neg.Binomial parameter                         | 0.83 (0.46)    |                      |       |
| Large reporting errors at the second decimal (range: 0-6) |                |                      |       |
| (Intercept)                                    | -4.10 (1.78)   | 5.30                 | .021  |
| Data shared (1) or not (0)                     | -1.20 (0.52)   | 5.39                 | .020  |
| Square root (Average of p-values)              | 17.17 (9.42)   | 3.32                 | .069  |
| Log (No. of test statistics)                   | 0.71 (0.45)    | 2.53                 | .112  |
| Neg.Binomial parameter                         | 1.41 (0.84)    |                      |       |

Negative binomial regressions (N = 49; with a log link) of the number of misreporting errors per paper on the log of the number of test statistics, square root of the average p-value of genuinely significant results within the papers, and whether or not the data were shared for reanalysis. Analyses were estimated in SPSS 18.0 (The Zelig package in R gave similar results) with a robust variance estimator to deal with the possibility that errors were dependent within papers. Natural log and square root transformations were used to improve predictors' normality.
Supporting Information

The request email was based specifically on Standard 8.14 of APA’s Ethical Principles of Psychologists and Code of Conduct (http://www.apa.org/ethics/code/index.aspx) and ran as follows:

Dear Dr. [Author’s name],

As a student at the Psychology Department of the University of Amsterdam I am currently conducting a study supervised by Drs. Wicherts and Borsboom. For this research I am gathering data from studies, which are published in several APA-journals in 2004. The aim of our study is to verify the substantive claims made in published research through reanalysis of data. Our specific aim is to assess the general impact of outliers on effect sizes and correlations reported in psychological research. Therefore I would like to ask you if you would be so kind as to send me the raw data of the [study/studies] reported in your paper ‘[paper title]’, published in [journal name]. Of course I realize that it will take some time and effort to fulfill this request. Please note, however, that it is not necessary to completely rewrite the databook. You can just send me the file with the essential data and I will contact you later if something appears to be unclear. I would greatly appreciate it if you would help me by letting me use your data. Of course, the confidentiality of the participants in your study is guaranteed. Thank you for considering my request and I am looking forward to your answer.

Yours sincerely,

Mrs. J. Kats – Psychological Methods, Department of Psychology, University of Amsterdam

It is possible that the studies from which data were shared have larger sample sizes than the studies from which no data were shared. This may have led to
difference in statistical power and hence the difference in p-values. This
possibility can be studied by comparing the papers on the basis of the typical cell
size as based on the degrees of freedom from t tests and particular F tests. So
we considered the denominator DFs from F tests that had DF = 1 in the
numerator and the DFs of all the t tests in the papers. We computed the mean of
the error DF as an indicator of typical cell size in the analyses for each of the 48
papers that reported such statistics. The mean of this variable in 27 papers that
were not shared (M = 199, SD = 656, Md = 57) was somewhat higher than the
mean in the 21 papers from which data were shared (M = 77, SD = 107, Md =
48), albeit not significantly so: Wilcoxon’s W = 441, Z = 1.52, p = .127 (2-
tailed). Hence, the cell sizes in these analyses did not suggest that statistical
power was systematically higher in studies from which data were shared.