# Meta-Analyzing the Multiverse: A Peek Under the Hood of Selective Reporting

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# 10 ABSTRACT

Researcher degrees of freedom refer to arbitrary decisions in the execution and reporting of hypothesistesting research that allow for many possible outcomes from a single study. Selective reporting of results (p-hacking) from this 'multiverse' of outcomes can inflate effect size estimates and false positive rates. We studied the effects of researcher degrees of freedom and selective reporting using empirical data from extensive multi-study projects in psychology (Registered Replication Reports) featuring 211 samples and 14 dependent variables. We used a counter-factual design to examine what biases could have emerged if the studies (and ensuing meta-analyses) had not been preregistered and could have been subjected to selective reporting based on significance of the outcomes in the primary studies. Our results show the substantial variability in effect sizes that researcher degrees of freedom can create in relatively standard psychological studies, and how selective reporting of outcomes can alter conclusions and introduce bias in meta-analysis. Despite the typically thousands of outcomes appearing in the multiverses of the 294 included studies, only in about 30% of studies did significant effect sizes in the hypothesized direction emerge. We also observed that the effect of a particular researcher degree of freedom was inconsistent across replication studies using the same protocol, meaning multiverse analyses often fail to replicate across samples. We recommend hypothesis-testing researchers to preregister their preferred analysis and openly report multiverse analysis. We propose a descriptive index (Underlying Multiverse Variability) that quantifies the robustness of results across alternative ways to analyze the data.

- **Competing interests** The authors have no financial or non-financial competing interests.
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## 14 Author Note

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Researcher Degrees of Freedom (DFs, Simmons et al., 2011) refer to the many arbitrary decisions 23 that need to be made in designing, collecting, analyzing, and reporting research. In the analysis of 24 hypothesis-testing research, the focus of this article, researcher DFs involve decisions such as choosing 25 between different approaches for dealing with missing observations, excluding participants from the 26 analysis depending on different criteria, and a range of other data processing and modelling decisions 27 (for more examples, see Wicherts et al., 2016). Researcher DFs allow for many possible outcomes in 28 a single study where the reported result depends on the specific combination of decisions made. This 29 was illustrated recently by Silberzahn et al. (2018): When 29 independent teams examined the same 30 data with the same research question, the teams' estimated effect sizes (measured as odds ratios) varied 31 from 0.89 to 2.93, with 20 teams finding a statistically significant effect in the expected direction. That 32 different independent teams of researchers reached different estimates shows that there often is no clearly 33 preferable analysis in hypothesis-testing research (see also Botvinik-Nezer et al., 2020; Breznau et al., 34 2021; Huntington-Klein et al., 2021; E.-J. Wagenmakers et al., 2022). 35

The many possible statistical results that are enabled by researcher DFs have been referred to as a 36 'multiverse' of statistical results (Steegen et al., 2016), 'vibration of effects' (Patel et al., 2015), or a 37 specification curve analysis' (Simonsohn et al., 2020). These multiverse style analyses entail (sensitivity) 38 analyses of the robustness of results to researcher DFs and offer insights into potential biases that might 39 emerge if researchers selectively report outcomes from them to present more convincing evidence in favor 40 of a hypothesized effect. The proponents of multiverse style analyses are careful to define 'reasonable' or 41 'arbitrary' decisions in light of substantive, statistical, methodological, and psychometric grounds (Del 42 Giudice & Gangestad, 2021; Patel et al., 2015; Simonsohn et al., 2020; Steegen et al., 2016). In the 43 current study, we use multiverse analysis based on arbitrary choices to demonstrate the potential impact 44 of selective reporting on study-level effects and subsequent meta-analyses of resulting effect sizes. To 45 ensure arbitrariness in our researcher DFs, we consider the effect size computations to be given. That is, 46 we apply researcher DFs that we consider to not change the independent or dependent variables, and we 47 do not add covariates or change the statistical model or constructs of interest. 48

From the perspective of the broader literature the principal concern with researcher DFs is not that they allow multiple statistical results to be computed, but rather that they allow for selective reporting of possibly desirable outcomes. Throughout this article we use 'selective reporting' to refer to cases where multiple statistical results are examined in a study, but some go unreported (Page et al., 2020). We do not include in this definition the special case where no study results are reported and hence do not focus on publication bias of entire studies. Selective reporting is often focused on the significance of outcomes and can be intentional ('*p*-hacking') or happen unintentionally due to hindsight and confirmation biases (Nickerson, 1998; Roese & Vohs, 2012). Selective reporting from the multiverse of statistical results is problematic as it can allow researchers to present statistical evidence even for incredible phenomena (Simmons et al., 2011). Numerous formal approaches and simulation studies have been used to show that selective reporting leads to an overrepresentation of false positive findings (Ioannidis, 2005) and inflated effect size estimates (Ioannidis, 2008) in the literature.

Unfortunately, selective reporting appears common amongst researchers. In psychology, about 50-60% 61 of researchers admit to not reporting all dependent measures in a study (Agnoli et al., 2017; John et al., 62 2012), and in a study registry comparison 70% of studies did not report all outcome variables (Franco et 63 al., 2016). Moreover, there is an extensive literature on selective reporting in the fields of biomedicine, 64 with evidence from, for example, neurology (Fusar-Poli et al., 2014), hematology (Wayant et al., 2017), 65 pediatrics (Rosati et al., 2016), orthopedics (Rongen & Hannink, 2016), obesity (Rankin et al., 2017), 66 and cancer research (Kyzas et al., 2005). A recent study examining results in 67 trials published between 67 October and November 2015 in 5 top journals from general medicine found that 42% of pre-specified 68 outcomes went unreported (Goldacre et al., 2019). Further evidence from the fields of education (Pigott 69 et al., 2013) and studies on partner violence (Madden et al., 2019) suggests the problem of selective 70 reporting is widespread indeed. 71

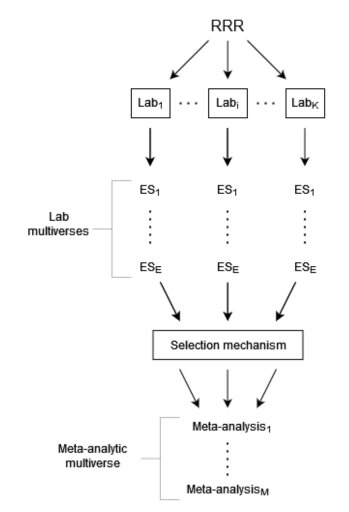
The biases created by selective reporting in primary studies are inherited by meta-analyses that seek 72 to quantitatively review effects or associations across many studies. Each of the studies included in a 73 meta-analysis have their own multiverse. Since the results used for meta-analysis are subsets from these 74 multiverses, meta-analytic result(s) also represent a subset from the multiverse of possible meta-analyses. 75 To avoid that this subset is biased, meta-analytic reporting guidelines such as PRISMA (Moher et al., 76 2009) and MARS (Appelbaum et al., 2018) recommend meta-analysts to evaluate primary studies for 77 selective reporting. We do not consider arbitrary choices made in the context of meta-analyses themselves 78 (i.e., multiverse meta-analysis: Palpacuer et al., 2019; Voracek et al., 2019), but rather vary the analyses 79 in the primary studies while keeping the meta-analytic inclusion criteria and analysis constant (i.e., we 80 meta-analyze multiverses) to study the biasing effects of selective reporting based on researcher DFs in 81 primary studies on meta-analytic outcomes. 82

Such biasing effects have been studied in simulated data for meta-analysis (e.g., Botella et al., 2021; Carter et al., 2019; Friese & Frankenbach, 2020), but simulated data from known distributions may not be representative of actual psychological data that feature unknown (distributional) complexities. Also, the effects of researcher DFs have been studied in observed data of individual studies (Botvinik-Nezer et al., 2020; Breznau et al., 2021; Huntington-Klein et al., 2021; e.g., Silberzahn et al., 2018), but not in meta-analytic context to inform how they might affect cumulative knowledge. We combine these streams of research and study the effects of researcher DFs and selective reporting in observed meta-analytic data, taking advantage of the unique opportunity offered by the open data of ten recent multi-lab direct replication projects in psychology (Registered Replication Reports) that featured a total of 211 samples studying 14 different outcome variables.

Registered Replication Reports (RRRs) each consist of a set of studies (labs) that collected data 93 on an effect in psychology using the same pre-specified research design, decision plan, and materials, 94 collectively known as a 'preregistration.' Each RRR can be seen as making up one (or more) meta-analysis 95 of direct (also called 'exact') replications, where the only difference between included studies is where 96 they collected their data. Even though the preregistrations used in the actual RRRs limited the effect of 97 researcher DFs in the original analyses, the open data from these extensive studies enable us to use a 98 counter-factual design to see what biases could have emerged if the studies (and ensuing meta-analyses) 99 had not been preregistered and could have been subjected to selective reporting based on significance 100 of the outcomes in the primary studies. In doing so, we demonstrate the variability in results that may 101 arise in meta-analytic data in the absence of preregistration, the limitations of multiverse analysis when 102 applied to a single study, and illustrate the entire process of selective reporting, from the researcher DFs 103 in primary studies that enable the practice to the consequences for meta-analysis. 104

## 105 METHODS

Figure 1 summarizes the design of this study. We identify decision points in each RRR where reasonable 106 alternative decisions could have been made (absent any preregistration) and compute all resulting outcomes 107 (create a multiverse) for each included lab. We then combine effect sizes from the lab multiverses in 108 meta-analysis within each RRR. This design allows us to explore the effects of researcher DFs on research 109 output by 1) examining the underlying multiverse variability in effect size estimates at the primary study 110 level, 2) examining the resulting multiverse variability at the meta-analytic level, and 3) examining 111 different mechanisms for selecting effect sizes from primary study multiverses for inclusion in the meta-112 analysis. We refer to the variability due to researcher DFs as the Underlying Multiverse Variability (UMV, 113 statistically defined in 'the multiverses' section). 114



For each RRR, we coded a set of researcher Degrees of Freedom (DFs). We applied the same researcher DFs to all labs within an RRR (Supplement B).

For each lab, we computed all possible outcomes based on our researcher DFs. We show and discuss the resulting underlying multiverse variability (UMV) and other lab statistics (Figures 2 - 4).

To meta-analyze lab results, we selected effect sizes from each lab multiverse either randomly or by emulating selective reporting in 4 different ways (Table 3).

We compared meta-analyses based on our selection mechanisms with original preregistered RRR estimates and the distribution of all possible meta-analytic combinations (Figure 5).

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*Figure 1.* Summary of the study design. For each lab in a Registered Replication Report (RRR) multiverses were computed, analyzed and used for meta-analysis. ES = Effect size. Each RRR consists of *K* labs, lab *i* = 1, 2, 3, ... *K*. Each lab has *E* effect size estimates in its multiverse. There are *M* possible combinations of the *E* effect sizes across labs, resulting in a meta-analytic multiverse of size *M*. We approximate the meta-analytic multiverse by randomly sampling  $10^5$  meta-analyses from the meta-analytic multiverse. For details on how we selected researcher DFs, see Methods under the header 'Selection and justification of researcher DFs.' This figure was created using the website draw.io.

#### 123 Transparency and openness

All our code and data for this project are available on the Open Science Framework (OSF) at osf.io/j8yg2/, and permanently archived at Zenodo (doi.org/10.5281/zenodo.7341292). We refer directly to relevant files on the OSF using brackets and links in the sections below. We registered data cleaning code and the researcher DFs available in each RRR before proceeding to analysis (osf.io/h397y/). We only made minor code corrections and clarifications of researcher DFs after registration, fully detailed in Supplement A (osf.io/xem2y/). We handled all data in R version 4.0.2 (R Core Team, 2020), and cite used packages in

#### 130 the reference list.

#### 131 Data Collection

We included all ten RRRs available at the time of data collection (i.e., published up until 2019-05-15) available in the journals 'Perspectives on Psychological Science' and 'Advances in Methods and Practices in Psychological Science' (see also Olsson-Collentine et al., 2020). Three RRRs (RRR3, RRR5, RRR9) had multiple primary outcome variables (as explicitly identified in the accompanying publications). In total, we included 10 projects containing 14 primary outcome variables that could be meta-analyzed, consisting of 211 unique lab samples and 34,357 participants (Table 1). These values correspond to the sum of the labs and participants of RRR1 - RRR9, as RRR9 and RRR10 used the same sample.

We use the RRRs because the meta-analyses they offer allow us to consider the effects of (selective 139 reporting from) multiverse analyses in relation to a benchmark based on meticulously collected data 140 from multiple labs (and different samples) using the same protocol. We selected the RRRs for our study 141 over other multilab replication initiatives for two reasons. First, we wished to examine researcher DFs 142 within a meta-analytic structure, which is what the RRRs nicely provide. The RRRs have the additional 143 advantage that most of them report average results not significantly different from zero, allowing us to 144 examine the bias from selective reporting under the most problematic circumstances (i.e., when there is 145 no genuine effect) and the percentage of significant outcomes appearing across multiverses (Type I error 146 rate). Second, we wished to allow for researcher DFs to depend on study design. The Many Labs series of 147 replication projects (which we have worked with previously in Olsson-Collentine et al., 2020) consists of 148 many effects studied at the same time in the same samples, meaning (almost) all researcher DFs will be 149 identical across all studied effects. Hence the RRRs allow us to delve deep into the generalizability of the 150 multiverse variability across labs and effects. 151

152 Table 1.

#### 153 Preregistered Multi-Lab Replication Projects

| RP    | Paper                      | Countries | Labs | Effects | Ν    | Sample and Settings  | Description of Effects  |  |
|-------|----------------------------|-----------|------|---------|------|--|---|--|
| RRR1  | Alogna et al.<br>(2014)    | 10        | 31   | 1       | 4832 | 31/32 samples were<br>undergraduate students aged<br>18-25, 1 general population<br>which was also the only online<br>sample.                | Verbal overshadowing 1; Independent<br>two-group experiment. Participants either<br>described a robber after watching a video or<br>listed countries/capitals and after a filler task<br>attempted to identify the robber in a lineup.  |  |
| RRR2  | Alogna et al.<br>(2014)    | 8         | 26   | 1       | 2932 | 22/23 samples were<br>undergraduate students aged<br>18-25, 1 general population<br>which was also the only online<br>sample.                | Verbal overshadowing 2; Different from 1 only<br>in that the filler task took place before the<br>descriptive task instead of after.  |  |
| RRR3  | Eerland et al.<br>(2016)   | 2         | 10   | 3       | 1210 | 11/12 samples were<br>undergraduate students mostly<br>aged 18-25, one of which was<br>online. 1 sample was a broader<br>online sample.      | Grammar's effect on interpretation; Independ<br>two-group vignette experiment with three<br>outcome variables. Participants read about<br>actions either described in imperfect or perfo<br>tense and then rated protagonist's intentions<br>(intentionality/intention attribution/detailed<br>processing). |  |
| RRR4  | Hagger et al.<br>(2016)    | 10        | 24   | 1       | 3127 | All samples consisted of in-lab<br>undergraduate students  | Ego depletion; Independent two-group<br>experiment. Participants either assigned to a<br>cognitively demanding or a neutral task, and<br>performance was then measured in a subsequ<br>cognitive task.  |  |
| RRR5  | Cheung et al.<br>(2016)    | 5         | 16   | 2       | 2279 | All samples consisted of in-lab<br>undergraduate students aged<br>18-25  | Commitment on neglect/exit; Independent<br>two-group experiment with two outcome<br>variables. Participants either primed to think<br>about commitment to or independence from<br>partner.  |  |
| RRR6  | Wagenmakers et al. (2016)  | 8         | 17   | 1       | 2542 | All but one sample explicitly<br>consisted of students and all took<br>place in-lab. The last sample was<br>recruited at university grounds. | Facial feedback hypothesis; Independent<br>two-group experiment. Participants either<br>induced to 'smile' or 'pout' by holding a pen in<br>their mouth differently and simultaneously rates<br>funniness of cartoons.  |  |
| RRR7  | Bouwmeester et al. (2017)  | 12        | 21   | 1       | 3669 | All samples consisted of in-lab<br>undergraduate students aged<br>18-34.   | Intuitive cooperation; Independent two-group<br>experiment. Economic game with money<br>contribution to a common pool either under<br>time pressure or time delay.  |  |
| RRR8  | O'Donnell et al.<br>(2017) | 13        | 40   | 1       | 7041 | All samples consisted of in-lab<br>undergraduate students aged<br>18-25  | Professor priming; Independent two-group<br>experiment. Participants primed with either a<br>'professor' or 'hooligan' stimuli. Outcome w<br>percentage correct trivia answers.   |  |
| RRR9  | McCarthy et al.<br>(2018)  | 13        | 26   | 2       | 6720 | All samples consisted of in-lab<br>students aged 18-25   | Hostility priming; Independent two-group<br>experiment with two outcome variables.<br>Participants descrambled sentences, either 20%<br>or 80% were hostile, then rated an individual<br>and a list of ambiguous behaviors on perceived<br>hostility.   |  |
| RRR10 | Verschuere et al. (2018)   | 12        | 25   | 1       | 3245 | All samples consisted of in-lab<br>students aged 18-25   | Moral reminder; Independent two-group<br>experiment. Participants either recalled the Ten<br>Commandments or books they had read.<br>Outcome was degree of cheating when reporting<br>results.  |  |

Note:

All RRRs published up until 2019-05-15 in the journals 'Perspectives on Psychological Science' and 'Advances in Methods and Practices in Psychological Science'. RP = Replication Project, Countries = number of lab country locations, Effects = number of primary effects studied, N = participants before exclusions, RRR = Registered Replication Report. Table adapted with permission from Olsson-Collentine et al. (2020). Code to reproduce table: osf.io/jehpy/

We downloaded individual level data for all RRRs in Table 1. Summary data of all RRRs were available on the OSF. When the raw lab data were not publicly available via the OSF we contacted authors by email to request them. Only for one lab in RRR1 and 2 and two labs in RRR3 were we unable to acquire their individual level raw data.

<sup>158</sup> For each RRR, we standardized data formatting across labs, fixed minor mistakes (e.g., mislabeled

columns in RRR8), and prepared the datasets for multiverse analysis (osf.io/cf86y/). We prepared the data in the same way as the original RRRs to the extent possible. However, we largely had to write our own code, because the alternative decisions needed to create our multiverses (e.g., exclusion criteria) could not be taken in the code by the original authors. In preparing the datasets for analysis, we only excluded participants due to reported experimenter error or when participants were reported to not have followed instructions or completed the experiment. Note that exclusions based on 'not following instructions' are usually *ad hoc*, and hence are distinct from formalized exclusions based on attention checks.

#### 166 **The Multiverses**

Although multiverse type analyses have been suggested by multiple authors under somewhat different names (Patel et al., 2015; Simonsohn et al., 2020; Steegen et al., 2016) all multiverse analyses consist of identifying points in the research process where multiple reasonable decisions could have been made, identifying what these decisions might be, and examining the impact of these decisions on the study results. A core assumption of multiverse type analysis is that the alternative decisions are all (approximately) equally reasonable (Simonsohn et al., 2020; Steegen et al., 2016; see also Del Giudice & Gangestad, 2021).

It is important that researcher DFs are defined such that these choices are indeed 'reasonable' or 174 'arbitrary' on a priori substantive and methodological grounds (Patel et al., 2015; Simonsohn et al., 175 2020; Steegen et al., 2016). Del Giudice & Gangestad (2021) rightly pointed out that many decisions 176 implemented in published multiverse analyses were not truly arbitrary (are 'non-equivalent') because they 177 can a priori be expected to result in different a) measurement reliability/validity, b) studied psychological 178 effects (e.g., when including a covariate that changes the prediction) or c) power/precision. Only decisions 179 for which this does not hold (i.e., we are either confident they are equivalent or uncertain) should according 180 to Del Giudice & Gangestad (2021) be included in a multiverse analysis. 181

We agree with Del Giudice & Gangestad (2021) that such substantive and methodological issues should be considered when performing multiverse analysis as a sensitivity analysis. Hence, we carefully selected our researcher DFs to reflect choices for which we saw no prior substantive or methodological grounds to expect them to affect the true effect sizes tapped by the different labs. However, we could imagine that others might object to some of those choices and hence we offer a range of supplementary results to assess how alternative choices in designing the multiverse affect our results. We also make our data and code available for re-analysis using alternative multiverse setups.

#### 189 Selection and justification of researcher DFs

<sup>190</sup> We selected our researcher DFs to correspond to normative researcher behavior that is at risk of selective

reporting in the fields of the RRRs that make up our data. These RRRs belong to the fields of social

and cognitive psychology (Olsson-Collentine et al., 2020). Although there are many researcher DFs 192 before analyzing the data (Wicherts et al., 2016), due to using already collected data we were only able 193 to vary post data collection decisions. Moreover, because our focus was on researcher DFs in primary 194 studies and their consequences for downstream meta-analysis, we only varied decisions in data processing 195 (the data multiverse, Steegen et al., 2016) and not the statistical models used in data analysis (the model 196 multiverse). Consequently, several chosen researcher DFs concerned using different exclusion criteria 197 (which we prepend with 'E'), although we also varied how the composite score was computed from 198 multiple indicators (researcher DFs prepended with 'S'). 199

When creating our list of researcher DFs, we proceeded in two steps: We 1) set up a list of 'common' 200 researcher DFs, and then 2) set up a list of researcher DFs unique to each RRR. These were then combined 201 to create our final list of researcher DFs for each RRR, which we registered before analyzing any data 202 (Supplement B). Because all labs in an RRR used the same design, it was only necessary to identify 203 decisions and create associated options once for each RRR and not for each lab/study separately. Our 204 coded researcher DFs each consisted of a decision that needs to be made and several associated potential 205 options for that decision. When defining the researcher DFs, we explored the data in the sense of 206 examining whether potential DFs could be applied (i.e., whether the variables existed and how they were 207 defined) but did not examine what effect applying them would have. 208

We created our list of common researcher DFs based on recommendations in statistical textbooks, 209 common decisions by applied researchers as reported in research literature, data analytic decisions made 210 by the included RRRs, and our own experience of decisions encountered in the literature. Table 2 provides 211 an overview of all common researcher DFs. We considered option a) across decisions to be the default 212 option, corresponding to no scale adjustments or participant exclusions (although for some researcher 213 DFs, an active decision must be made; S2, E1 Table 2). In Supplement C we detail how we selected 214 each researcher DF and its options. We acknowledge that many additional multiverses could be run in 215 these and other studies, but we consider our setup typical of researcher DFs that could be used in practice 216 across a range of psychological studies, and as such useful to study the influence of selective reporting. 217 Researchers instead interested in using multiverse analysis as a sensitivity analysis for a particular effect 218 should carefully consider the advice of Del Giudice & Gangestad (2021) on equivalent pathways before 219 applying any of the researcher DFs in Table 2. 220

#### Table 2.

Note:

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List of common researcher degrees of freedom applied to Registered Replication Reports

| Decision                           | Options  | Short Explanation   |  |  |  |
|------------------------------------|--|---|--|--|--|
| S1. Post hoc scale length          | <ul><li>a) No adjustment</li><li>b) Drop the item with the lowest item-rest correlation</li><li>c) Drop the two items with the lowest item-rest correlations</li></ul>                                       | It is unclear how common it is to post hoc drop items 'that<br>don't work' from a scale, but dropping more than a few seems<br>unlikely. In the research we are looking at (experimental) ther<br>are rarely long scales. Excluding 1-item scales, the average<br>scale length in a large sample of psychological research in 201-<br>was 6.87 (SD = 7.18) (Flake et al., 2017)   |  |  |  |
| S2. Composite score                | <ul><li>a) Unweighted average score</li><li>b) Sum score</li><li>c) PCA score: Varimax rotation, force two components and pick the first, requires at least 3 items.</li></ul>                               | For Likert-type scales with multiple items. Other DVs e.g.,<br>reaction time variability (RRR3), dichotomous correct/incorrec<br>(RRR1&2), continuous measures (RRR7), single item DVs<br>(RRR10) may need more unique choice options. We chose<br>Varimax rotation to maximize variance between outcomes.  |  |  |  |
| E1. Missingness DV                 | <ul> <li>a) Any missing items -&gt; list-wise deletion</li> <li>b) If ≤25% items missing then pair-wise deletion of missing items. Otherwise list-wise.</li> </ul>   | List-wise deletion appears to be by far the most common<br>approach to missing data. In van Ginkel et al.'s (2010) review<br>of personality psychology 97% used list-wise deletion for<br>missing data and several reviews in medicine have also found it<br>to be an extremely common method (Eekhout et al., 2012;<br>Rombach et al., 2016; Burton & Altman, 2004). Nonetheless,<br>we see that for example RRR6 used pair-wise deletion (option<br>b) which may seem reasonable to some researchers, in<br>particular with a longer scale. |  |  |  |
| E2. Missingness E3-E4 variables    | a) No exclusion<br>b) Missing on any variable -> list-wise deletion  | RRRs that excluded data based on a E3-E4 variable (e.g., ag<br>did list-wise deletion when data was missing. For other<br>non-DV variables we make no exclusions based on<br>missingness, unless this was explicitly done by the project (e.<br>'task completion' RRR10).   |  |  |  |
| E3. Age                            | a) No exclusion<br>b) Not 18-24<br>c) Not 18-23<br>d) Not 18-22<br>e) Not 18-21  | Used by 9/10 RRRs for exclusions. Across 25 cohorts of Dutch<br>bachelor psychology students 96.7% of students were below 25<br>92.7% below 24, 86.6% below 23 and 77.8% below 22<br>(Wicherts et al, 2012). The oldest students in this dataset were<br>25. We choose a set of age ranges based on these data that we<br>believed might go unremarked if used as exclusion criteria in<br>psychological literature with an accompanying motivation such<br>as 'we only included young adults'.   |  |  |  |
| E4. Language / Student / Ethnicity | a) No exclusion<br>b) Exclude participants not belonging to the dominant category  | Used by 3/10 RRRs. Demographic variables which are<br>sometimes used for exclusions. Language includes variables<br>such as 'native speaker' which may have a yes/no response.<br>Ethnicity includes similar variables such as 'country of birth' o<br>'race'. If multiple of these demographic variables are available<br>they are treated as separate exclusion criteria.   |  |  |  |
| E5. Attention check                | <ul> <li>a) No exclusion</li> <li>b) Exclude if failed &gt;50% of attention check items (i.e., with two items, must fail both, e.g., RRR7)</li> <li>c) Exclude if failed any attention check item</li> </ul> | Attention checks are common in psychology, as evidenced by<br>the more than 1500 citations of Oppenheimer et al. (2009) whi<br>introduced 'instructional manipulation checks'. Curran (2016<br>suggests' conservative' exclusions based on 50% failed<br>attention checks when multiple items are used. This category<br>does not include manipulation checks which vary more in<br>format.   |  |  |  |
| E6. Univariate outliers            | <ul> <li>a) No exclusion</li> <li>b) DV score &gt; 2 SD from mean</li> <li>c) &gt; 3 SD from mean</li> <li>d) &gt; 1.5 times the interquartile range</li> </ul>  | Used by 1/10 RRRs. Commonly recommended cutoffs (Bakker & Wicherts, 2014). Test for outliers across groups.   |  |  |  |
| E7. Multivariate outliers          | <ul> <li>a) No exclusion</li> <li>b) Mahalanobi's distance with p &lt; .001</li> </ul>   | If the outcome variable is a correlation. Tabachnik, Fidell and Ullman (2007) recommend using Mahalanobi's distance with cutoff of $p < .001$ for detecting multivariate outliers. Outliers tested within groups as recommended by Tabachnik et al.   |  |  |  |

S. = Degree of Freedom (DF) affecting Scale composition, E. = Exclusion DF, DV = Dependent Variable, SD = Standard Deviation, PCA = Principal Component Analysis, RRR = Registered Replication Report. Code to reproduce table: osf.io/jehpy/.

In addition to the list of common researcher DFs, which we applied to all RRRs, each RRR had several unique researcher DFs. These arise from the uniqueness of each research topic and design and consisted of different exclusion criteria. We coded between 2 (RRR3) and 10 (RRR7) unique researcher DFs for each RRR, each decision with 2-6 associated options. Due to the large number of unique researcher DFs we do not describe them all in detail here, but provide only a broad overview and refer interested readers to Supplement B.

We can separate between two types of 'unique' researcher DFs: either 1) the RRRs excluded partici-

pants based on some variable that was not defined in our list of common decisions or 2) an RRR measured

variables (not in our list of common decisions) that they could have used for exclusions. As an example of the first case, in RRR4 (ego-depletion) participants with less than 80% correct on the main task were excluded. However, 80% is a largely arbitrary number, and someone might also consider values such as 75%, 85%, 90%, or many others, in addition to no exclusions. In cases like these, when there are an infinite number of possible values to choose from, we have elected only a maximum of six possible values that we believe an applied researcher would reasonably pick.

As an example of where an RRR measured variables they could have used for exclusions: RRR5 (commitment to romantic partner), amongst other things, asked participants whether they lived within 60 miles of their partner (yes/no) but did not use this variable in their analysis. However, another researcher might have found it relevant to only consider participants (not) living close to each other and used this variable for exclusions. Collecting data on a variable with no clear purpose thus adds researcher DFs and increases the risk of selective reporting, which we in this case used to create our multiverses.

#### 243 Applying researcher DFs to the RRRs

After registering the coding protocol for 'common' researcher DFs, we coded the applicability of each common researcher DFs to each RRR, which differed depending on, for example, how the outcome variable was measured (binary vs. continuous, one item vs. a scale) and how projects coded their data. Because some labs within RRRs pre-screened their participants for the original RRR exclusion criteria, it was not always possible to apply all exclusion criteria to all labs in an RRR. Nonetheless, we still included such labs, prioritizing the inclusion of more labs over the possibility of less multiverse variation. The coded common and unique researcher DFs for all RRRs are available in Supplement B (osf.io/wj38n/).

We computed resulting effect sizes from all possible combinations of decisions for each lab in an 251 RRR (osf.io/zhdrx/). Incompatible decision combinations were not applied. For example, if we wished 252 to drop two items from a scale (Table 2; S1c) but required at least three items in the scale for Principal 253 Component Analysis (Table 2; S2c), this decision combination was inapplicable to scales with fewer 254 than 5 items. We standardized mean differences (Cohen's d, p. 226, Borenstein, 2009) and computed 255 log odds ratios for RRR1 and RRR2. Effect sizes were originally analyzed unstandardized in all RRRs 256 except for RRR4, and if certain researcher DF lower the within sample variance, as is highly likely, then 257 standardized effect sizes will appear larger. However, because certain of our researcher DFs change the 258 dependent variable, and we wanted to draw conclusions across RRRs, it was necessary to standardize 259 effect sizes. As most meta-analyses use standardized effect sizes and we are interested here in the biasing 260 effects of selective reporting on typical unregistered meta-analyses, we do not consider standardization of 261 effect sizes a major concern for our analysis. 262

To prevent including lab multiverses with an unrealistically small number of participants, we only

included lab multiverses with at least 24 participants per experimental group, the median sample size in
psychology (Bakker et al., 2012), in our primary analyses. Three labs in RRR2 (L09, L17, L26) and one
lab in RRR8 (L24) had smaller sample sizes than required in all conditions and were excluded from these
analyses. We present the results of our analyses also without this sample size restriction in Supplement D.

#### 268 Analysis

A consequence of assuming that the alternative decisions in the multiverse are equally reasonable is that 269 under the null hypothesis that no researcher DF has a systematic effect (i.e., is an actual moderator of 270 the effect) we can consider the distribution of effect sizes in the multiverse as random variability around 271 a true score. We refer to the variability underlying a given set of researcher DFs as the Underlying 272 Multiverse Variability (UMV) and define it as the standard deviation (SD) in effect size estimates that are 273 in the multiverse of the same study. A different set of researcher DFs will reveal different UMV. Other 274 researchers have focused on the distribution of p-values (Simonsohn et al., 2020; Steegen et al., 2016) or 275 on the range of effect sizes in the multiverse (e.g., Patel et al., 2015), but we consider it more useful to 276 treat multiverse variability in terms of the standard deviation of effect size estimates, in line with how 277 sampling error is defined. The UMV should be seen as a descriptive tool that highlights some degree of 278 variability that might have a relation with bias due to selective reporting over and beyond sampling error, 279 rather than a well-defined statistic. 280

To demonstrate the effects of researcher DFs on research output, we 1) examined the variance in 281 effect size estimates at the lab level (lab multiverses) and 2) compared meta-analytic average effect size 282 estimates based on how lab outcomes were selected from their multiverses. To examine the variance in 283 effect size estimates at the lab level, we created funnel plots, computed UMV, and standard deviations 284 in effect size resulting from variation across the options within a single researcher DF. For the funnel 285 plots, we plotted all effect size estimates at the lab level using either the standard error (for log odds 286 ratios; RRR1/RRR2) or sample size as the y-axis (for standardized mean difference effect sizes; RRR3 -287 RRR10). We used total sample size (N) on the y-axis for all standardized mean difference (SMD) effect 288 types since most of our coded researcher DFs affected sample size. 289

To examine how large the effects of applying a single researcher DF can be and the relative impact of our different researcher DFs, we computed the standard deviation in a lab's estimated effect size across the options associated with each decision. For each researcher DF, we computed the standard deviation in effect size when all other researcher DFs were set to their default value (corresponding to option 'a' for each researcher DF, see Table 2 and Supplement B). In addition to examining standard deviations for labs within RRRs, we also disaggregated these lab estimates across RRRs and then aggregated them across common and unique researcher DF categories. In doing so, we treated all unique researcher DFs as one <sup>297</sup> category.

Due to computational limitations, and because it is often the case that some researcher DF must be 298 applied before another (e.g., outliers cannot be removed before the composite score has been computed), 299 we only applied the researcher DFs in a single fixed order. That is, if we have three researcher DFs (1, 2, 300 3) then we always applied them in the order 1, 2, 3 regardless of chosen option, rather than also varying 301 the order (e.g., 2, 1, 3). This fixed order may affect results when removing items with the lowest item-rest 302 correlation from a scale, or excluding participants based on outlier criteria, although we see no reason to 303 expect a systematic interaction between these two and any other researcher DFs. The fixed order also 304 makes it impossible to compute the impact of a single researcher DF across all possible researcher DF 305 combinations, although it remains possible to compute its impact when not applying any other researcher 306 DFs (see previous paragraph). 307

When comparing meta-analytic average estimates, we compared a) the original (preregistered) RRR estimates, with b) an estimate of the distribution of all possible meta-analytic combinations, c) randomly selected lab effect sizes, and d) lab effect sizes selected by one of four biased selection mechanisms (see below). We ran all meta-analyses as random-effects models with the restricted maximum likelihood estimator for estimating the between-study variance using the R-package 'metafor' (Viechtbauer, 2010).

The huge number of possible effect size combinations across labs for each RRR, the smallest consisting 313 of 697×10<sup>33</sup> possible meta-analyses, made it impossible to compute the full distributions of possible 314 meta-analytic outcomes. Instead, we drew large random samples to approximate the distributions. For 315 each RRR (or outcome variable when an RRR contained multiple primary outcomes), we proceeded as 316 follows: We drew one random effect size from all possible effect sizes from lab 1, one random effect 317 size from all possible effect sizes from lab 2, one from lab 3, one from lab 4, and so on until we had 318 drawn one effect size from all labs in an RRR. The drawn effect sizes across labs were then combined 319 using a meta-analysis. We repeated this procedure, sampling with replacement from each lab's multiverse 320 of effect sizes, until we had sampled 100,000 effect sizes from each lab, and consequently computed 321 100,000 meta-analyses for each RRR. These samples of meta-analyses constituted our approximation of 322 the distribution of possible meta-analyses for each RRR (or outcome variable when an RRR contained 323 multiple primary outcomes). The means of these distributions (and the means of the estimated lower/upper 324 95% Confidence Intervals; CIs) constituted our random sample of estimates. 325

When selectively reporting results, researchers may exhibit different behavior. We included four types of biased selection mechanisms (Table 3: 'Most significant,' 'Below alpha,' 'Random significant,' Bounded significant') with different motivations. All selection mechanisms focused on statistical significance, and we used a two-tailed test with  $\alpha = .05$  for hypothesis testing. First, we selected the effect

size with the lowest *p*-value in each lab. This allowed us to examine the most extreme selection of results 330 possible due to p-hacking ('most significant'). We included this scenario as a worst-case scenario. Second, 331 selective reporting may sometimes result in a 'bump' just below p = .05 when aggregating p-values across 332 selectively reported studies ('below alpha'). This is most likely in the case of incremental p-hacking 333 approaches such as optional stopping (e.g., Hartgerink, 2017). To compare what a meta-analysis of such 334 data might look like, in the 'below alpha' condition, we selected, for each lab in an RRR, the outcome 335 with a *p*-value closest below .05 (or, if there were no *p*-values below .05, the lowest value). These two 336 approaches ('most significant,' 'below alpha') attempted to select a single result from the multiverse, but 337 it may be that several effect sizes have equivalent *p*-values due to being based on exactly the same sample. 338 If so, we picked the effect size with the fewest researcher DFs deviating from their default option a. 339

Third, we represent a *p*-hacker who is satisfied with any significant effect size they encounter (in 340 the expected direction), by picking a random effect size out of those that were statistically significant 341 ('random significant'). If no effect sizes were significant, the effect size with the lowest *p*-value was 342 picked. Fourth and finally, when a *p*-hacking researcher tries multiple analyses, they might choose to 343 report the analysis that resulted in the smallest *p*-value. However, selecting the result with the smallest 344 *p*-value across the full multiverse suggests that the *p*-hacker systematically explored the full multiverse to 345 find the strongest possible effect, whereas reality probably consists of a more ad hoc and limited search. 346 Hence, we represent a 'bounded' search by 1) randomly drawing 100 possible outcomes and 2) out of 347 these 100 outcomes selecting the one with the smallest *p*-value. 348

With all our biased selection mechanisms (i.e., excluding the random draw and original meta-analytic 349 results, see Table 3), we applied a "hypothesized direction filter." That is, when selecting an effect size at 350 the lab level, we excluded all effect sizes that were in the opposite direction of the originally predicted 351 effect (osf.io/r2dum). If there were no effect sizes in the predicted direction, we excluded all significant 352 effect sizes in the 'wrong' direction and selected outcomes from the remainder. We added this filter 353 because we believe researchers who apply selective reporting in reality are unlikely to be agnostic about 354 the direction of their effect, and our focus in this study is on selective reporting, not Hypothesizing After 355 Results are Known (HARKing, Kerr, 1998) 356

### Table 3.

358

Summary of outcome selection mechanisms

| Selection mechanism    | Hypothesized<br>Direction Filter | Single outcome | Description   |  |  |
|------------------------|----------------------------------|----------------|---|--|--|
| Pre-registered         | No                               | Yes            | The original RRR meta-analytic average effect size with pre-registered decisions.   |  |  |
| Random draw            | No                               | No             | The average point estimate and upper/lower 95% CI from $10^5$ meta-analyses randomly sampled from all possible meta-analyses.   |  |  |
| Most signficant Yes    |                                  | Yes            | Select the effect size in the multiverse with the smallest <i>p</i> -value.   |  |  |
| Below alpha Yes        |                                  | Yes            | Select the effect size in the multiverse with a $p$ -value closest below $p = 0.05$ . If no $p$ -value below the cutoff, pick the smallest.   |  |  |
| Random significant Yes |                                  | No             | Identical to the random draw, but with effect size first limited to only significant effect sizes.  |  |  |
| Bounded significant    | Yes                              | No             | We drew 100 effect sizes from a lab's multiverse,<br>and selected the effect size with the lowest p-value<br>This was repeated $10^5$ times, resulting in $10^5$<br>values per lab. These were then meta-analyzed and<br>summarized as above for the random draw. |  |  |

Note:

Description of different implemented selection mechanisms for selecting effect sizes at the lab-level to metaanalyze. 'Hypothesized Direction Filter' = exclude effect sizes not in the predicted direction (yes/no), 'Single outcome' = selection mechanism resulting in a single meta-analytic result (yes/no). Code to reproduce table: osf.io/jehpy/.

Publication bias, the complete suppression of a study being published, and selective reporting (selection of reported results amongst multiple possibilities) are closely related, and it is intuitively appealing to believe correcting for publication bias may be sufficient for generally removing biases in the meta-analytic data (e.g., Kvarven et al., 2019). We applied three publication bias correction methods [PET-PEESE, 3PSM, and *p*-uniform\*; Stanley & Doucouliagos (2014); Vevea & Hedges (1995); van Aert & van Assen (2020)] to examine their applicability to selective reporting in the absence of publication bias.

# 365 **RESULTS**

After excluding conditions that resulted in fewer than 24 participants per experimental group, 8/14 RRR 366 multiverses decreased in size, as can be seen in Table 4. The absolute decrease was largest for the largest 367 multiverses (RRR05 and RRR07), with RRR07 showing the largest absolute decrease and decreasing 368 from 2,621,440 to 525,680 (an 80% decrease) potential outcomes. However, the proportionally largest 369 decrease was seen in RRR08, which decreased from 115,200 to 19,200 (83% decrease) potential outcomes. 370 More importantly, as evidenced by the median number of multiverses per lab, even in the RRRs with 371 relatively few researcher DFs, the researcher DFs jointly created thousands of alternative outcomes per lab. 372 Nonetheless, many labs found zero significant (at p = .05) effect sizes within their multiverses. Across all 373

- studies (counting labs with multiple DVs as separate studies), 205 / 294 (70%) encountered no significant
- effect sizes in the hypothesized direction in their multiverses and 134 / 294 (46%) found no significant
- effect sizes in any direction. The size of the multiverse was strongly correlated with the number of studies
- $_{377}$  that encountered significant effect sizes in the hypothesized direction within their multiverses (Pearson's r
- <sup>378</sup> = 0.63, 95% CI [0.14, 0.87]).

### 379 Table 4.

380

Multiverse sizes before and after filtering out outcomes with < 24 participants per experimental group

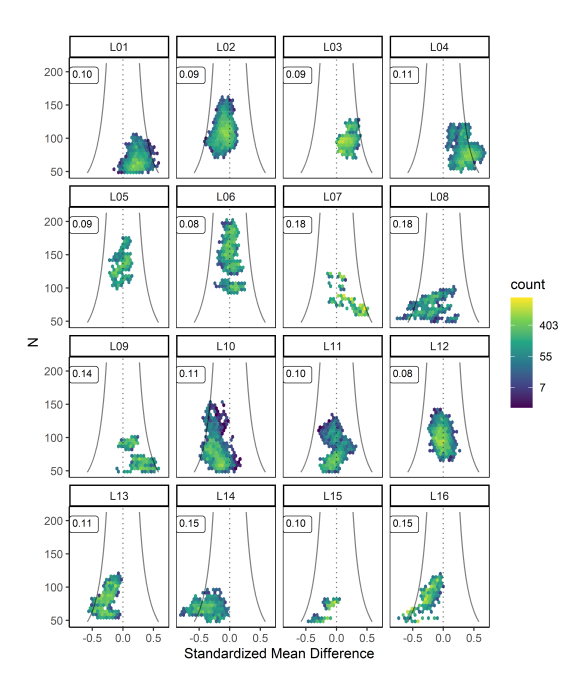
| Meta-Analysis     | Common<br>DFs | Unique<br>DFs | N [lq, uq]     | Multiverse size<br>before exclusion | Multiverse size<br>after exclusion | Labs after<br>exclusion | Labs with any<br>sig. | Labs with hyp. sig. |
|-------------------|---------------|---------------|----------------|-------------------------------------|------------------------------------|-------------------------|-----------------------|---------------------|
| RRR01             | 5             | 5             | 116 [107, 125] | 3,840                               | 3,840 (100%)                       | 31                      | 4 (13%)               | 4 (13%)             |
| RRR02             | 5             | 5             | 88 [84, 98]    | 3,840                               | 3,840 (100%)                       | 23                      | 10 (43%)              | 10 (43%)            |
| RRR03 Attribution | 6             | 2             | 84 [82, 84]    | 3,840                               | 3,816 (99%)                        | 10                      | 5 (50%)               | 3 (30%)             |
| RRR03 Intention   | 6             | 2             | 84 [82, 84]    | 3,840                               | 3,840 (100%)                       | 10                      | 3 (30%)               | 1 (10%)             |
| RRR03 Process     | 7             | 2             | 84 [83, 85]    | 7,680                               | 7,680 (100%)                       | 10                      | 2 (20%)               | 0 (0%)              |
| RRR04             | 5             | 3             | 76 [68, 90]    | 23,040                              | 20,160 (88%)                       | 24                      | 14 (58%)              | 7 (29%)             |
| RRR05 Exit        | 6             | 9             | 82 [70, 94]    | 2,488,320                           | 1,503,904 (60%)                    | 16                      | 12 (75%)              | 7 (44%)             |
| RRR05 Neglect     | 6             | 9             | 82 [70, 94]    | 2,488,320                           | 1,540,176 (62%)                    | 16                      | 15 (94%)              | 10 (62%)            |
| RRR06             | 7             | 5             | 96 [77, 111]   | 122,880                             | 61,440 (50%)                       | 17                      | 9 (53%)               | 5 (29%)             |
| RRR07             | 3             | 10            | 74 [63, 95]    | 2,621,440                           | 525,680 (20%)                      | 21                      | 18 (86%)              | 10 (48%)            |
| RRR08             | 8             | 4             | 79 [65, 102]   | 115,200                             | 19,204 (17%)                       | 39                      | 31 (79%)              | 16 (41%)            |
| RRR09 Behavior    | 8             | 4             | 169 [114, 218] | 46,080                              | 46,080 (100%)                      | 26                      | 16 (62%)              | 5 (19%)             |
| RRR09 Hostility   | 8             | 4             | 168 [114, 218] | 46,080                              | 46,080 (100%)                      | 26                      | 13 (50%)              | 10 (38%)            |
| RRR10             | 5             | 5             | 90 [77, 107]   | 11,520                              | 3,808 (33%)                        | 25                      | 8 (32%)               | 1 (4%)              |

Note:

Meta-analytic distributions and estimates after excluding analytic choices that resulted in <24 particants per experimental group at the study-level. Three labs in RRR2 (L09, L17, L26) and one lab in RRR8 (L24) always had fewer than 24 participants and were excluded. 'Labs with any sig.' = number of labs (%) with any significant (at p = .05) effect size in their multiverse, 'Labs with hyp. sig.' = number of labs (%) with any significant (at p = .05) effect size in their multiverse. DFs = Researcher Degrees of Freedom, 'Common DFs' = DF from a common list of potential DFs, 'Unique DFs,' = study-unique DFs,' = Median study supplies across their multiverse. Code to reproduce table: osf.io/jehpy/.

## 381 Lab multiverses

- <sup>382</sup> There can be substantial variation in effect sizes within labs due to researcher DFs. Figure 2 shows
- effect sizes across the multiverses for 16/24 labs in RRR04. Similar plots for all RRRs (or outcome
- variables when an RRR contains multiple), including for all labs in RRR04, can be found in Supplement
- 385 E (osf.io/2htc6/).



386

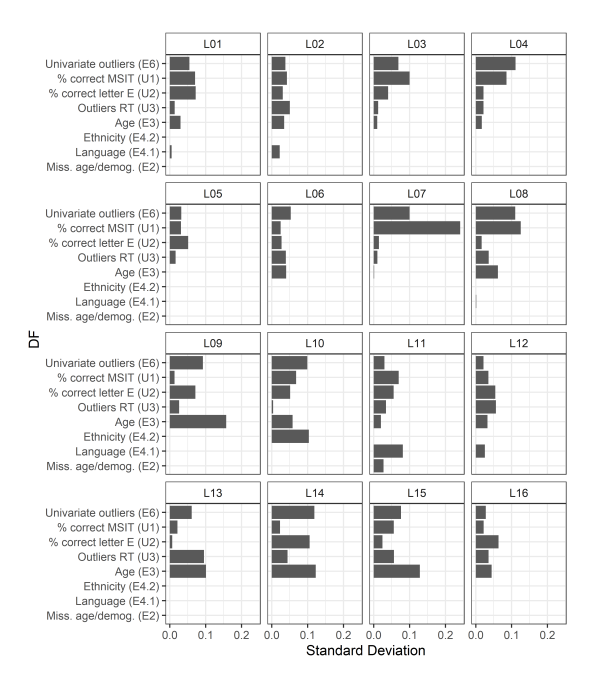
Figure 2. Arbitrary decisions in research cause underlying multiverse variability (UMV) in effect size 387 estimates. Funnel plots showing the effect sizes based on the multiverses in 16 labs for RRR04, after 388 removing cases where n < 24 in either experimental group. Values in upper left corner of each facet are 389 UMV for each lab. For legibility, 16/24 RRR04 labs are shown; the figure including all labs is available 390 in Supplement E (osf.io/2htc6/). L01 – L16 are lab indicators. Solid lines are funnel lines based on the 391 *t*-distribution. Effect sizes falling outside the funnel lines are statistically significant at  $\alpha = .05$  using a 392 two-tailed test. Dotted lines indicate zero effect size. Colors in the funnel plots indicate the frequency of 393 occurrence of an effect size. Brighter colors indicate that an effect size occurred more often. N = total 394

sample size. Code to reproduce figure: osf.io/thuyk/.

Overall in Figure 2, statistically significant observations (indicated by observations falling outside the funnel lines) were rare (median = 0.87%, interquartile range = 0 - 3%). There were labs with a higher proportion of significant outcomes (L14 = 25%, L04 = 24%, L16 = 17%), but in only one case were these in the hypothesized direction (L04). The median underlying multiverse variability (UMV) across labs in Figure 2 was 0.1SD, interquartile range (IQR) = 0.09 - 0.15. Effect sizes could change by as much as d =0.97 (L08). Pearson's correlation based on the 16 labs in Figure 2 between UMV and sample size before applying researcher DFs was r = -0.51.

Across RRRs, the median lab UMV was 0.11SD (IQR = 0.08 - 0.14) for SMD effect sizes and 0.07SD (IQR = 0.04 - 0.12) for log OR, but researcher DFs could change effect sizes in a lab by as much as d =1.27 (RRR05 Neglect, L10) and log OR = 1.31 (RRR01, L05). We expected the lab UMV, just as the standard error, to be generally negatively correlated with (original) sample size. However, the median correlation between lab UMV and sample size (before applying researcher DFs) within RRRs was r =0.09 (IQR = -0.11 - 0.37). Hence, a large sample size does not ensure a small UMV.

There can be substantial variation between labs also in which researcher DF leads to variability in 409 effect sizes (Figure 3). Figure 3 shows the standard deviation (SD) in effect size within the labs from 410 Figure 2 when applying a single researcher DF. Despite identical study design across labs and the same 411 researcher DF being applied, no two bar plots look identical and labs differ in which researcher DF 412 creates the most variation. For example, in Lab 7 (L07) excluding participants based on different accuracy 413 criteria for the main DV (U1) resulted in the largest SD, whereas in Lab 6 (L06) using different criteria 414 for defining and excluding outliers (E6) led to the most variation in estimated effect size. Figures 2 and 415 3 together demonstrate that multiverse analyses should not necessarily be expected to be replicable in 416 new data, because across labs the same researcher DFs can yield different degrees of effect size variance 417 (Figure 2) and this variance can be primarily caused by different researcher DFs (figure 3). Note that the 418 effect sizes in Figure 2 arose from all possible combinations of the researcher DFs and not by applying 419 them separately as in Figure 3 (see 'analysis' methods section), which explains the larger range of effect 420 sizes in Figure 2. 421

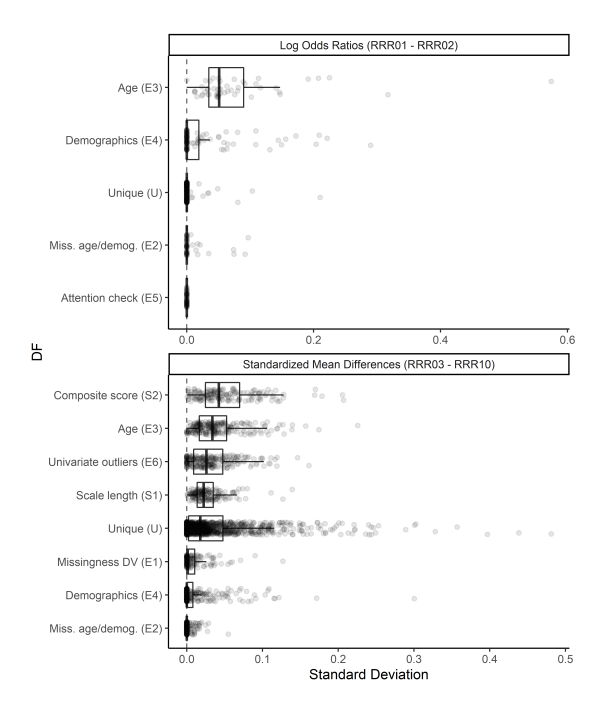


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*Figure 3.* The same arbitrary decisions have a different effect in comparable studies. Standard deviation (SD) in effect size estimates in 16 labs in RRR04 resulting from applying different researcher degrees of freedom (DFs) individually, after removing cases where n < 24 in either experimental group. For legibility, 16/24 RRR04 labs are shown; the figure including all labs is available in Supplement E (osf.io/2htc6/). L01 – L16 are lab indicators. Indicators in parentheses on the *Y*-axis (E2, E3, E4\_1, E4\_2, E6, U1, U2, U3) refer to DF codes for each coded DF (Supplement B osf.io/wj38n/). The *Y*-axis is ordered by median SD across labs. Code to reproduce figure: osf.io/thuyk/

430 Some researcher DFs generally contribute more to the UMV than others, and thus constitute larger

risk factors when considering selective reporting. Figure 4 shows the SDs in estimated effect sizes in labs 431 resulting from applying each researcher DF individually, disaggregated across RRRs and then aggregated 432 into their respective categories. How the composite score was computed from a scale with multiple 433 items had the largest median effect amongst SMD effects (d = 0.04, lower panel top row, Figure 4), and 434 in supplement F we demonstrate how removing (the source of variation in) this researcher DF (S2.c; 435 computing the composite score based on PCA) decreases overall UMV, showing how the removal of 436 sources of variation effectively decreases the risk of selective reporting. Figure 4 also show that excluding 437 participants on age had a relatively strong effect (median upper panel,  $\log OR = 0.05$ , lower panel, d =438 0.03. In supplement G we show that this effect is driven by the large degree of exclusions across the 439 options of the age researcher DF (E3), by comparing it with a version with more broad inclusion criteria. 440 That is, when researcher DFs result in datasets with less overlap (i.e., that are less correlated), the UMV 441 increases. Hence, we can predict that certain researcher DFs are of more serious concern under selective 442 reporting, although the observed effect in any given sample will depend on random fluctuations in the 443 sample. 444



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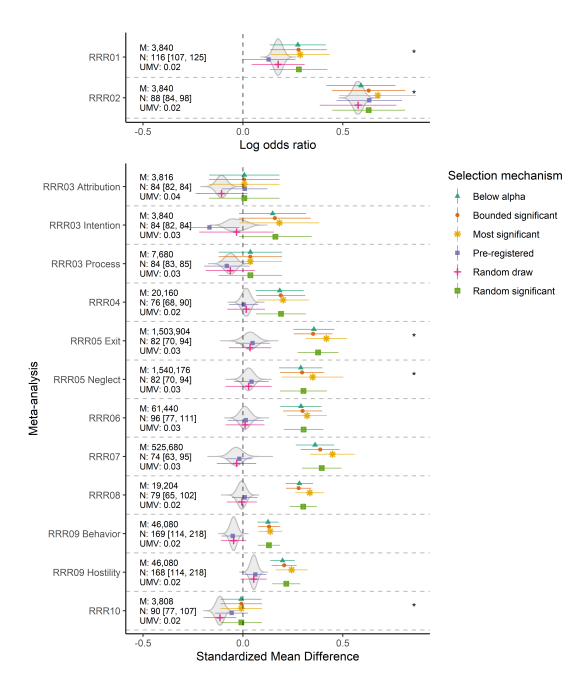
Figure 4. Some arbitrary decisions tend to create more effect size variability than others. Standard 446 deviation (SD) in effect size estimates in labs resulting from different researcher degrees of freedom 447 (DFs) applied individually. The top panel shows results for Registered Replication Reports (RRRs) with 448 an outcome measured as log odds ratio, and the lower panel for RRRs measured as standardized mean 449 differences. The Y-axis is ordered by median effect size SD. Data are after removing cases where n < 1450 24 in either experimental group, disaggregating DFs across Registered Replication Reports (RRRs) and 451 aggregating into categories. Indicators in parentheses on the Y-axis (S1, S2, E1, E2, E3, E4, E5, E6) refer 452 to DF codes in Table 2 or (U) to DFs coded as unique for each research project. The Unique (U) category 453

454 was aggregated across all distinct unique DF. Code to reproduce figure: osf.io/thuyk/.

Unique researcher DFs only had the fifth highest median SD (d = 0.02) for SMD effects (lower panel 455 Figure 4), likely due to many unique researcher DFs having little effect. However, they also show the 456 largest range in possible outcomes. For example, within RRR07, choosing to exclude participants based 457 on whether they complied with the set time limit or not (U2) resulted in the largest median effect size 458 SD of all researcher DFs for that RRR (SD = 0.21, see also Supplement E). This same researcher DF in 459 RRR07 resulted in 4 out of the 5 highest effect size SDs in Figure 4. The remaining observation (third 460 from the right) belonging to RRR09 Hostility, lab 1, and resulted from choosing whether to exclude 461 participants based on their study major. Unique researcher DFs show less impact in the log odds ratio 462 effects (upper panel Figure 4), which may be due to fewer labs/researcher DFs, and most unique researcher 463 DFs in RRR01/02 only being applicable to a few of the constituent labs (see Supplement B). For example, 464 only three labs included a comprehension check (U1), and only three (different) labs coded 'familiarity 465 with effect' (U2). 466

#### 467 Meta-analytic multiverses

Variability in effect sizes within labs due to researcher DFs implies that many different meta-analytic 468 outcomes are possible. How and which effect sizes were selected in labs will change meta-analytic 469 results. Figure 5 shows multiple meta-analytic average effect size estimates for all outcome variables, 470 depending on how effect sizes were selected in the constituent labs. The grey density curves indicate the 471 empirical distributions of meta-analytic point estimates across multiverses for each outcome. UMV in 472 point estimates ranged from 0.02SD (RRR09 Hostility) to 0.04SD (RRR03 Intention) for standardized 473 mean differences and for log odds ratios rounded to 0.02 SD for both RRR01 and RRR02. When outcomes 474 were selected through a preregistered decision procedure (purple squares, Figure 5), meta-analytic mean 475 estimates were generally close to the mean of the estimated multiverse distributions, and matched the 476 random draw estimates well (pink crosses, Figure 5). The point estimate of the random draws is by 477 definition identical to the mean of the multiverse distribution, whereas the lower/upper bounds are the 478 average 95% CIs of these draws. 479



480

Figure 5. Selective reporting in labs results in overestimates in meta-analysis. Meta-analytic distribu-481 tions and estimates after excluding analytic decisions that resulted in n < 24 participants per experimental 482 group at the study-level. Selection mechanism = how effect sizes were selected at the study-level, either 483 by p-hacking ("Most significant," "Bounded significant," "Random significant," "Below alpha,"), preregis-484 tered decisions ("preregistered"), or random selection ("Random draw"). "Random draw" and "Random 485 significant" are averages across  $10^5$  draws from the meta-analytic multiverse, whereas other selection 486 mechanisms are a single outcome. M = Median study multiverse size, N [lower quartile, upper quartile] 487 = Median study sample sizes across their multiverses, \* = effect size sign changed (RRR01, RRR02, 488

RRR05 Exit, RRR05 Neglect, RRR10) so that hypothesized effect size (and *p*-hacking direction) was
 positive for all meta-analyses. Code to reproduce figure: osf.io/thuyk/.

When researcher DFs were combined with biased selection of effect sizes in labs (*p*-hacking), meta-491 analytic mean estimates were also more extreme in the predicted direction (Figure 5). As expected, 492 493 selecting the most significant effect size in each lab (yellow stars) and then meta-analyzing resulted in the most extreme mean estimates. Other *p*-hacking approaches (in Figure 5: turquoise triangles, red circles, 494 and green squares) resulted in similar estimates. This similarity in outcome between biased selection 495 mechanisms can be mostly attributed to the low number of significant results across labs and our biased 496 selection procedures resulting in the same results if there were no significant outcomes in a lab (205 / 294 497 studies across all RRRs, counting labs with multiple DVs as separate studies). 498

There is a tendency for projects with larger multiverses (e.g., RRR05, RRR06, and RRR07, as can be 499 seen in Figure 5) to have more extreme estimated effect sizes when p-hacked. The difference between the 500 average random draw (pink crosses) and the estimates based on the most significant effect sizes (yellow 501 stars) ranged from 0.1 to 0.48 for SMDs and was about 0.1 for log OR. The correlation between effect 502 size inflation and multiverse size was r = 0.77 for SMD effect sizes. The most extreme case corresponded 503 to RRR07 (Figure 5), where the difference in meta-analytic average effect size estimate between the 504 average random draw (pink cross, d = -0.03, 95% CI [-0.13, 0.07]) and the estimate based on the most 505 significant effect sizes (yellow star, d =0.45, 95% CI [0.34, 0.56]) was an increase of almost 0.5SD in the 506 predicted direction. Applying publication bias correction methods (PET-PEESE, 3PSM and *p*-uniform\*) 507 did not lead to improvements in estimated average effect size estimates (Supplement H), in line with other 508 research that has shown publication bias correction methods as unlikely to be useful in correcting for 509 selective reporting (Carter et al., 2019; van Aert et al., 2016). 510

#### 511 DISCUSSION

In this article, we performed multiverse analysis across multiple direct replication studies, using empirical 512 data from ten Registered Replication Reports (RRRs). Even though the preregistrations used in the 513 actual RRRs limited the effect of researcher DFs in the original analyses, the open data from these 514 extensive studies enabled us to use a counter-factual design to see what biases could have emerged 515 if the studies (and ensuing meta-analyses that included them) had not been preregistered and could 516 have been subjected to selective reporting based on significance of the outcomes in the primary studies 517 (p-hacking). We identified researcher DFs based on common decisions in the associated literature for 518 each outcome variable, computed all possible outcomes across direct replications, and examined the 519 variance in these so-called multiverses. We then combined effect sizes from the multiverses of each 520

direct replication in meta-analysis and examined the consequences of different mechanisms for selecting 521 effect sizes for inclusion. Our analyses highlight that multiverse analyses typically yielded thousands of 522 different outcomes within single studies, that multiverse patterns of variation differed across labs using 523 the same protocol, and that selective reporting of outcomes in primary studies could bias meta-analytic 524 results, despite their status as a 'gold-standard' of evidence. We also found that original sample size 525 correlates at most weakly with potential for selective reporting (as measured by UMV), suggesting a larger 526 sample size does not protect against selective reporting. Yet interestingly, 205 / 294 studies (counting labs 527 with multiple DVs as separate studies) did not contain any significant (measured as  $p \le .05$ ) results in 528 their multiverses, suggesting *p*-hacking null results into significance may be more difficult than expected 529 considering the sheer number of potential outcomes per study. We discuss these results and the limitations 530 of our own study in the remainder of the discussion. 531

#### 532 Defining the multiverse

Creating a multiverse is an inherently subjective endeavor given that researchers might disagree about 533 which decisions are (approximately) equally reasonable (Steegen et al., 2016). For example, although we 534 created our researcher DFs based on common practice in the associated literature, there are methodological 535 arguments to carefully consider the meaning and impact of outliers, or use outlier robust statistics (e.g., 536 Rousseeuw & Hubert, 2011) rather than excluding them based on rules of thumb. In the same vein, it may 537 often be preferable to perform multiple or maximum likelihood imputation (e.g., Jakobsen et al., 2017) of 538 missing data points rather than excluding them. For this reason, we have endeavored to structure our data 539 such that a disagreeing reader familiar with R can explore the consequence of only including those of our 540 researcher DFs they consider reasonable, as demonstrated in Supplement F. 541

Del Giudice & Gangestad (2021) critically discussed what it means for a researcher DF to be 542 'reasonable.' They rightly argued why some of the researcher DFs as used in previously published 543 multiverse analyses might not be equivalent on prior grounds or might not show equivalence for reasons 544 yet unknown. We admit that our researcher DF S1, which deletes items in measurement scales based 545 on the lowest item-rest correlations, could perhaps be non-equivalent on psychometric grounds. On 546 the one hand, deletion of the item with lowest item-rest correlation could heighten the reliability of the 547 scale, and hence increase genuine effects, if the deleted item were poorly performing. On the other hand, 548 deletion would have little effect on the reliability and hence on actual effects if the item performed as 549 well as other items in the scale. Hence, if there were some item(s) consistently performing poorly (in 550 the psychometric sense) across labs in an RRR, this researcher DF could increase true effects and be 551 of principled non-equivalence, at least for that RRR. Similarly, the researcher DF related to composite 552 scores (S2) could create genuinely different effects under some conditions. In Supplement F, we repeated 553

<sup>554</sup> our analyses without the PCA option of this researcher DF which showed less variation in outcomes, as <sup>555</sup> expected if indeed the way composite scores are computed makes a difference. However, like all our DFs, <sup>556</sup> we included this researcher DF because we feel that it might be used in practice in unregistered studies.

One could also argue that multiverses with varying samples sizes are non-equivalent in the statistical 557 sense of sampling variability (Del Giudice & Gangestad, 2021). Smaller sample sizes generally result in 558 less statistical power to detect effects and hence larger p-values, which makes it more difficult to directly 559 compare *p*-values across multiverses differing in sample size. We do not consider the issue of power to 560 be a major concern for our study, as many of our analyses focus on effect size estimates and variability, 561 and because most RRRs in our sample (apart from RRR1/RRR2, based on our reading of the original 562 articles) appear to study null effects. In addition, although researcher DFs that lead to smaller sample 563 sizes would increase variability of outcomes regardless of true effect size, the strong sample overlap 564 between multiverse samples in a given study creates covariation between multiverses that diminishes total 565 sampling variability. In the end, which multiverses are considered reasonable will depend not only on 566 individual researchers' beliefs, but also on which decisions their research community considers acceptable 567 in terms of theoretical, methodological, and empirical standards (Del Giudice & Gangestad, 2021). 568

Notwithstanding selective reporting, researchers should recognize that researcher DFs, or decisions 569 treated as if they were researcher DFs, by themselves create another layer of uncertainty in study estimates; 570 we found a median UMV of 0.1SD in our SMD data, although this will differ depending on research 571 field and which DFs researchers find reasonable. As such, we advise researchers doing hypothesis-testing 572 research to 1) preregister the (single) analysis they believe is optimal for testing their hypothesis, motivate 573 why this is the case and report uncertainty estimates (e.g., confidence or credibility interval), and 2) include 574 a multiverse analysis as a sensitivity analysis (following the advice of Del Giudice & Gangestad, 2021) 575 and report their UMV. Although some may argue that preregistering a preferred analysis is contradictory 576 if the options in the multiverse analysis are considered equivalent, pragmatically, we believe that most 577 researcher DFs will not be exactly equivalent, and that most researchers will have a preferred analysis that 578 it would be useful to accompany with a sensitivity analysis. The goal of these recommendations is for the 579 research process to be transparent so that results act as credible evidence despite the potential effect of 580 researcher DFs on outcomes. 581

More generally, multi-lab collaboration, or regular, projects may wish to consider incorporating the multiverse perspective already in the design of their studies, identifying which of their decisions are largely arbitrary and collecting data on alternatives. Preregistration of research is likely to be helpful from this perspective, in addition to its transparency enhancing properties, which are helpful when evaluating a study for selective reporting.

#### 587 Exploring the multiverse

The unique design of our project enabled us to examine the effect of researcher DFs (i.e., perform 588 multiverse analysis) across multiple direct replication studies. We observed that 1) the same researcher 589 DFs applied to direct replication studies resulted in widely varying distributions of effect sizes, and 2) 590 which researcher DF caused the variability within a study differed between direct replications. That 591 is, the effect of researcher DFs both within and across direct replication studies appeared unsystematic. 592 These results demonstrate that results of multiverse analysis in any single study, like other exploratory 593 analyses, are not necessarily replicable in new data. We believe this point is underappreciated among many 594 multiverse analysts. In addition, some researchers may be tempted to directly interpret the existence of 595 researcher DFs and resulting UMV as evidence of 'hidden moderators' (Van Bavel et al., 2016); currently 596 unknown moderators that explain why effect sizes differ between studies. However, the existence of the 597 multiverse does not by itself imply moderators as substantial variability and apparent moderating effects 598 may be found through sampling variance alone. 599

That the effect of researcher DFs both within and across direct replication studies was generally nonsystematic also corroborates previous findings of ours (Olsson-Collentine et al., 2020) that differences in study results in social and cognitive psychology show little to no between-study heterogeneity, and supports the conclusion that the best explanation for differences between effect sizes in (direct) replication studies is typically the joint effect of sampling error and researcher DFs, possibly in combination with selective reporting.

When we have a substantive researcher DF that we suspect of being a moderator, it may be most 606 useful to examine it from an empirical meta-analytic perspective. If we have a researcher DF at the 607 study level (e.g., measurement scale) with sufficient variation between primary studies, it is possible to 608 examine it as a moderator using meta-regression (e.g., Houwelingen et al., 2002). However, individual 609 level researcher DFs (e.g., age) are preferably examined in individual participant data (IPD) meta-analysis 610 to avoid the ecological fallacy (e.g., Stewart & Tierney, 2002). In the case of multiple dependent variables, 611 which might also be a researcher DF, potential systematic differences could be examined in multivariate 612 meta-analysis (e.g., Jackson et al., 2011). As with multiverse analysis, such moderator analyses should 613 primarily be considered exploratory and hypothesis-generating. 614

Researcher DFs in primary studies also add a layer of uncertainty to meta-analysis when those studies are meta-analyzed. Researcher DFs in primary studies can change both point estimates and the associated standard errors and can do so across multiple studies. Consequently, in meta-analysis they can influence not only the meta-analytic point estimate but also the between-study variance. That said, the standard deviations in point estimates over the meta-analytic multiverses in our meta-analyses were quite small,

with an UMV of at most 0.04SD amongst SMD effects. This is unsurprising: when researcher's decisions 620 are truly random and in the absence of selective reporting and publication bias, researcher DFs in primary 621 studies can be expected to cancel out across a sufficiently large number of studies. As such, researcher 622 DFs in primary studies (and resulting multiverses) are not a major concern for meta-analysts in the 623 absence of selective reporting. Unfortunately, the availability of such ideal data is not expected in most 624 meta-analyses. Even with ideal data, in a meta-analysis with a small number of primary studies, as is 625 common in medicine (Davey et al., 2011), researcher DFs are less likely to balance each other out and 626 meta-analytic UMV may be a larger concern. 627

Encouragingly, in our data, the meta-analytic point estimates based on preregistered studies often 628 fell close to the center of the meta-analytic distributions. Preregistration may generally lead to less 629 effect size inflation (Schäfer & Schwarz, 2019) by decreasing the risk of selective reporting through 630 increased transparency (although preregistrations are of varying quality or not always suitable, Bakker et 631 al., 2020; Pham & Oh, 2021). However, the connection between multiverse analysis and preregistration 632 may have been enhanced by the nature of our data: large collaborative projects, including researchers with 633 adversary hypotheses, which may have resulted in a 'wisdom of the crowd' selection of decisions amongst 634 researchers DFs. Alternatively, if no decision within a multiverse has a systematic effect, results from 635 any preregistered set of decisions from that multiverse would also be expected to coincide approximately 636 with the mean of the multiverse distribution when analyzed across samples. Regardless, to account for 637 UMV preregistered multi-lab collaborations (e.g., Moshontz et al., 2018) may offer a way forward in the 638 absence of more concrete theory (Fried, 2020), although as we saw in our results even such data is not a 639 guarantee for a point estimate at the center of the meta-analytic multiverse distribution. It is important 640 to be aware that a preregistered set of decisions nonetheless only represents a single universe from the 641 multiverse. 642

The extent of UMV in any given field depends on the multiverse created, and our estimates in this study may only apply to our non-random sample of social and cognitive psychology research. UMV in other fields could be either larger or smaller, but is unlikely to be non-existent, and it may be worthwhile to study the UMV in different subfields to examine their susceptibility to selective reporting given normative research behavior. Generally, the extent of bias introducible by selective reporting will depend on the multiverse size and the UMV, and researcher DFs that affect these two factors to a larger extent will hence contribute more risk of bias to a study.

#### **Selecting from the multiverse**

<sup>651</sup> We do not mean to imply that exploration of researcher DFs are problematic per se. We view it as <sup>652</sup> important to study the robustness of conclusions in the context of a sensitivity analysis, of which a

multiverse analysis can be seen as an extensive (systematic) variant. Exploring factors that truly moderate 653 an effect or association can be valuable as long as the exploration is transparently reported and employs 654 rigorous statistical controls to guard against overfitting. What is problematic is the selective or incomplete 655 reporting from the multiverse of statistical results. Hence, it is important to evaluate studies for risk of 656 selective reporting when using them to make decisions (e.g., about setting up future research), or including 657 them in systematic reviews (Appelbaum et al., 2018; as recommended by e.g., PRISMA and MARS: 658 Moher et al., 2009). Both preregistrations and multiverse analyses will facilitate evaluation of a study's 659 selective reporting risk by making research decisions more transparent, and there are many selective 660 reporting protocols available (e.g., Page et al., 2018) that may assist in the evaluation. 661

There is a risk that researchers exploit (intentionally or not) researcher DFs to selectively report those results from the multiverse that most strongly support their hypothesis. In extreme cases researcher DFs and *p*-hacking can provide evidence for any desired conclusion; in one lab the effect size estimate changed by as much as d = 1.27. More realistically, we found a median UMV of 0.1SD amongst 294 studies (counting labs with multiple DVs as separate studies). Nonetheless, a median UMV of 0.1SD in a field still implies that selective reporting can turn a statistically non-significant effect into a significant effect.

For many studies in our data this was not a concern. We found that in our data and given our researcher 668 DFs, about 70% of study multiverses did not contain a single significant result (measured as p < .05) in 669 the hypothesized direction. As most RRRs that made up our data had overall effect size estimates (based 670 on preregistered outcomes) not significantly different from zero, this coincides with previous findings 671 of ours that on average null results also tend to have very little heterogeneity (Olsson-Collentine et al., 672 2020). This suggests that it may be more difficult to p-hack null results into significance than many expect. 673 We caution that this observation may no longer hold when applying other types of researcher DFs than 674 we were able to do (e.g., this may not apply when researcher DF options are less correlated, as in the 675 case of outcome switching), and that 30% of labs did contain multiverses with a mix of significant and 676 non-significant effect size estimates. 677

Relatedly, we found a correlation not significantly different from zero between sample size and potential for selective reporting (as measured by UMV). This implies that sample size should not be taken to be protective against selective reporting, as also corroborated by a simulation study by Stefan & Schönbrodt (2022). We do note, however, that for genuine effects, larger sample sizes would increase power thereby lowering the need to selectively report outcomes based on the multiverse and hence less ensuing bias in estimated effects (Bakker et al., 2012).

<sup>684</sup> Our counter-factual design allowed us to see what biases could have emerged if the studies (and <sup>685</sup> ensuing meta-analyses) had not been preregistered and could have been subjected to selective reporting

based on significance of the outcomes in the primary studies, as is the case for most meta-analyses. Our 686 analyses demonstrate the substantial bias in the hypothesized direction that may be incorporated into 687 meta-analytic effect size estimates due to selective reporting in primary studies. The possible inflation 688 of average effect size will depend on the proportion of meta-analyzed studies at risk of bias and the 689 strength of this bias. Evidence from Kvarven et al. (2019) based on social and cognitive psychology 690 research suggests meta-analyses may sometimes estimate effect sizes to be as much as a third larger 691 than in comparable multi-lab projects, although this also includes publication bias. Larger or smaller 692 differences may be more typical in other fields. 693

For meta-analysts using retrospective data, minimizing the risk of bias in their included data (i.e., by 694 only including preregistered data and evaluating it for selective reporting) may be the best option until the 695 practices of multiverse analysis or the sharing of raw data become widespread. Our results corroborate 696 recommendations from meta-analysis reporting protocols such as PRISMA and MARS (Appelbaum et al., 697 2018; Moher et al., 2009) to always evaluate primary studies for risk of bias (here, selective reporting), 698 and we advise meta-analysts to study differences in outcome between studies identified as at high risk of 699 selective reporting bias and those at low risk. In line with previous research on the topic (Carter et al., 700 2019; van Aert et al., 2016), our supplemental results show that existing publication bias methods should 701 not be relied on to correct for *p*-hacking. 702

The large variance in impact of researcher DFs across direct replication studies demonstrates that the 703 study-specific effect of a researcher DF, and related bias induced by selective reporting, is difficult to 704 predict and dependent on sampling error. Stefan & Schönbrodt (2022), who simulated the effects of many 705 different *p*-hacking methods in single studies, reach a similar conclusion in their simulations: "Apart from 706 the aggressiveness of *p*-hacking itself, our simulations showed that across all strategies, the severity of 707 *p*-hacking also depends on the environment in which *p*-hacking takes place, for example, the correlation 708 structure in the data" (p. 46). Our results using the RRR data indeed show that the correlations between 709 multiverses create variation that is generally smaller than the sampling variation one would expect under 710 independent sampling. 711

That is not to say that we cannot draw some conclusions about the expected (average) impact of *p*-hacking different researcher DFs. The potential for effect size bias is larger when studies allow more analyses to be run and when more variance is created by included researcher DFs. Supplement G demonstrates that more overlapping (sub)samples created using alternative exclusions based on age created less subsequent UMV, which is expected given that more overlap creates higher correlations between alternative outcomes. As such, when considering effect size bias, we should (typically) be more concerned about researcher DFs in which the options are less correlated, although high false positive rates <sup>719</sup> are possible in either case (see discussion by Friese & Frankenbach, 2020).

It may be insightful to do more complex modelling of selective reporting from a multiverse perspective, 720 including of non-intentional selective reporting, and we hope our data will also be useful to other 721 researchers interested in more complex modelling of research bias. Our modelling of it in this study was 722 relatively straightforward and we only attempted to model the outcomes of intentional selective reporting 723 (p-hacking). Nonetheless, our biased selection methods applied to empirical RRR data are similar to those 724 used in the simulations of a recent compendium of p-hacking methods (Stefan & Schönbrodt, 2022) and 725 are on par with other recent simulation studies of p-hacking in meta-analysis (Botella et al., 2021; Friese 726 & Frankenbach, 2020). 727

Contrary to these two simulation studies (Botella et al., 2021; Friese & Frankenbach, 2020) we found 728 that p-hacking with actual data and using fairly generic researcher DFs could cause substantial inflation 729 of meta-analytic average effect sizes also when the average effect appears to be null. Both Friese & 730 Frankenbach (2020) and Botella et al. (2021) run extensive simulation studies of the effect of p-hacking 731 across many conditions and Friese & Frankenbach (2020) consider how it interacts with publication bias, 732 something we did not do. We believe the difference in results is due the choice in both of these simulation 733 studies to p-hack results based on the common assumption that p-hacking leads to a peak of p-values 734 below 0.05 (Hartgerink, 2017). In the case of Friese & Frankenbach (2020), results were p-hacked to 735 a distribution with a mode of p = 0.049, and Botella et al. (2021), similarly p-hacked studies into the 736 region .025 . We know from previous studies that the type of p-hacking matters; incremental737 methods such as optional stopping that result in a peak below p = .05 have little effect on effect size 738 inflation, whereas methods such as outcome reporting bias have a large effect and do not result in a 739 peak (Francis, 2012; Kirkham et al., 2010; Stefan & Schönbrodt, 2022). As such, differences between 740 our results and these simulation studies are likely explained by the incremental p-hacking methods used 741 in these simulation studies as compared to our methods of selecting the lowest *p*-values or randomly 742 selecting one of the significant outcomes. 743

Unfortunately, there exists little evidence on which method of selection researchers use in practice. Incremental *p*-hacking can still lead to concerning numbers of false positive results, as discussed by both Friese & Frankenbach (2020) and Botella et al. (2021), and it is important to discuss that not all types of *p*-hacking lead to concerning levels of effect size inflation. Nonetheless, our results show that suggesting selective reporting is not a concern for meta-analytic results is inaccurate when considering non-incremental *p*-hacking based on researcher DFs that we consider to be widely applicable across a range of empirical studies.

<sup>751</sup> Under some assumptions related to correlational structure of the overlapping data across multiverses,

we can be confident that the UMV and hence the potential bias due to selective reporting in a study is less than the reported standard error. The effect sizes in a multiverse are dependent because they are based on the same sample. Due to this dependence, the UMV will normally be smaller than the standard error in a study for a fixed sample size and statistical model, since the variability based on independent data is larger than of dependent data. In other words, if we know that the statistical model and sample size have not changed in a study, and that there is no publication bias, then we can be confident that the UMV in that study is less than its standard error estimate.

Consequently, at a fixed standard error the possible bias is always larger with publication bias than 759 with selective reporting due to the dependency between effect sizes in the multiverse. This suggests 760 that while *p*-hacking is likely more common than publication bias in the literature, being more resource 761 efficient, the distortion in the literature may be larger from publication bias when it does occur. Finally, 762 we note that the correlation between effect sizes within a multiverse also means that the independent 763 sample false positive rate (typically .05) should be expected to be lower when sampling effect sizes within 764 a dataset. We can observe this in our funnel plots, where substantially fewer than 1/20 effect sizes are 765 significant for most labs (i.e., fall outside the funnel lines). 766

#### 767 Limitations and constraints on generality

Although we have attempted to accompany all our claims and findings in this article together with their 768 caveats, we wish to make explicit the limits of generalizability of claims based on the data and design 769 of our study. The included effects are neither a representative nor a random sample of effects from 770 psychology. We expect our conclusions to be robust for effects in social and cognitive psychology, but 771 specific values that we report (e.g., median UMV of 0.1 among SMD effect sizes) may not generalize 772 beyond our sample. The RRRs in our data overwhelmingly reported average results not significantly 773 different from zero (the exception being RRR1/RRR2). This allowed us to examine selective reporting 774 in its most critical context (i.e., in the likely absence of genuine effects), but means it would be good to 775 focus future research efforts on studying multiverses with non-null effects, as these likely create more 776 heterogeneity in results across labs (see Olsson-Collentine et al., 2020), and hence we would also expect 777 larger overall variability and UMV due to researcher DFs (see also Friese & Frankenbach, 2020). During 778 the process of this project new RRRs have been published some of which report non-null effects and could 779 be used for such further analyses (e.g., Elliott et al., 2021). In addition, we are aware of several projects 780 currently in progress with similar designs that will collect new data and provide additional evidence on 781 the impact of standard researcher DFs in different fields. 782

Researchers interested in the theoretical implications of the specific effects studied here should
 carefully consider which researcher DFs they find reasonable (see Del Giudice & Gangestad, 2021)

before drawing conclusions. Our researcher DFs were chosen to match standard decisions in social and 785 cognitive psychology treated as researcher DFs, meaning they were not guided by the substantive theory 786 of the studied effects, except as reflected by decisions made by the RRRs in data collection and analysis. 787 Moreover, if for instance age is theoretically expected to moderate an effect it should preferably be tested 788 formally instead of being used in a multiverse analysis. van Aert (2022) demonstrated how this can be 789 done by looking at the effect of age across labs in RRR9 (McCarthy et al., 2018) using IPD meta-analysis, 790 finding a small positive interaction (p = .038). Individual studies often do not have the power to detect 791 moderating effects, which also affects multiverse analyses. 792

We found that in only about 30% of studies did significant effect sizes in the hypothesized direction 793 emerge in their multiverses, despite the typically thousands of analyses in every study. This finding 794 suggests it is more difficult to turn apparent null results into significant results than might be expected 795 but is dependent on our selection of researcher DFs. Although we implemented an extensive number of 796 researcher DFs that we consider representative of researcher DFs that could be used in practice across a 797 range of social and cognitive psychological studies, our use of secondary (real) data means there were 798 many researcher DFs (e.g., Wicherts et al., 2016) that we could not apply but that might be applied in 799 real situations (e.g., outcome switching, which is known to have a large impact, or changing the analytic 800 model). As such, our selection of DFs and the resulting multiverse variances are unlikely to represent a 801 worst-case scenario. It is feasible that in real life more extreme statistical results are found. Generally, we 802 can expect researcher DFs with lower correlation between options, because of less sample overlap and/or 803 weaker correlations between (in)dependent variables, to result in larger multiverse variance. 804

Finally, it may be informative to analyze other multi-lab replications studies than those we included in our study such as Many Labs 1 - 5 (e.g., Klein et al., 2018). The studies in our sample (mostly) studied a single effect across multiple labs, whereas the Many Labs projects study many effects at the same time across multiple labs. We examined the RRRs to be able to apply study-unique researcher DFs, but the Many Labs design would allow examining the impact of applying a single set of researcher DFs on a large sample of effects from social and cognitive psychology.

#### 811 Conclusion

We have shown that researcher degrees of freedom offer a wide array of potential outcomes in relatively standard psychological studies and demonstrated how selective reporting based on these researcher degrees of freedom creates bias in meta-analytic effect size estimates that may undermine the credibility of many meta-analyses. Preregistration is a methodological solution to researcher degrees of freedom enabling selective reporting, whereas a statistical solution is to perform multiverse analysis of results. These two transparency-enhancing practices can also be applied together, although our analyses of multiverses across direct replications highlight that multiverse analyses in single studies should not necessarily be expected to replicate in new data. Due to dependencies between effect sizes within multiverses, exploring multivariate approaches to multiverse analysis may be a useful next step in helping to address uncertainties and biases in primary studies due to researcher degrees of freedom.

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