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- Aggregating evidence from conceptual replication studies using the product Bayes factor
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Abstract

The product Bayes factor (PBF) can synthesize evidence for an informative hypothesis 17 across heterogeneous replication studies. It is particularly useful when the number of 18 studies is relatively low and conventional assumptions about between-studies heterogeneity 19 are likely violated. The present paper introduces a user-friendly implementation of the 20 PBF in the bain R-package, and demonstrates its use in several tutorial examples. The 21 method was validated in a simulation study that manipulated sample size, number of replication samples, and reliability. PBF had a higher overall accuracy when benchmarked 23 against other evidence synthesis methods, including random-effects meta-analysis (RMA). This was primarily due to PBF's greater sensitivity in detecting a true effect. However, PBF had relatively lower specificity. The PBF showed increasing sensitivity and specificity with increasing sample size. With an increasing number of samples, lower sensitivity was traded for greater specificity. Although PBF's overall performance was less susceptible to 28 reliability than the other algorithms, this masked a trade-off between reliability and 29 specificity. PBF thus appears to be a promising method for meta-analysis of heterogeneous 30 conceptual replication studies. Nonetheless, users should be aware of its lower specificity, 31 and the fact that the Bayesian approach to inference addresses a qualitatively different research question than other evidence synthesis methods. 33

34 Keywords: bayes factor, evidence synthesis, bayesian, meta-analysis

Word count: 5356

Aggregating evidence from conceptual replication studies using the product Bayes factor

Recent years have seen a crisis of confidence over the reliability of published results in 37 psychology, and science more broadly (Brembs, 2018). Replication research has emerged as 38 one potential way to address this crisis and derive knowledge that will stand the test of 39 time (see Layelle, 2021). In step with this interest in replication research, research synthesis methods have become increasingly popular. These methods aggregate research findings, and thus enable drawing overarching conclusions across multiple (replication) studies. This paper addresses Bayesian evidence synthesis, a research synthesis method that aggregates evidence for an informative hypothesis, quantified by the Bayes factor, across multiple studies. This method has the potential to provide a more comprehensive and accurate picture of the state of the literature, and to identify areas of consensus and disagreement among studies. We describe the method in detail, benchmark its performance against other commonly used research synthesis methods, and demonstrate its application through a tutorial example analysis. To facilitate uptake of the method by applied 49 researchers, we introduce an implementation of the method in the bain R-package. This implementation enables the use of Bayesian evidence synthesis for many commonly used 51 statistical analyses in R.

A key challenge in quantitative research synthesis is dealing with between-studies
heterogeneity (Higgins, Thompson, & Spiegelhalter, 2009). When studies examine the
same research question in different laboratories, use idiosyncratic methods, and sample
from distinct populations, these between-study differences can introduce heterogeneity in
findings. The most common quantitative research synthesis method is meta-analysis, in
which results of different studies are aggregated to estimate an aggregate effect size
(Borenstein, Hedges, Higgins, & Rothstein, 2009). In meta-analysis, heterogeneity can be
accounted for in four ways (see Van Lissa, 2020). First, if studies are exact replications, one
may assume that no heterogeneity in the outcome exists and a fixed-effect meta-analysis

can be conducted to estimate the common population effect. Second, when heterogeneity
between studies can be assumed to be random, random-effects meta-analysis can be used
to estimate the mean of a distribution of population effects. Third, when there are a few
systematic differences between studies, these can be accounted for using meta-regression.
Finally, when there are many potential variables that cause systematic differences and it is
not known beforehand which are relevant, exploratory techniques like random forest
meta-analysis and penalized meta-regression can be used to identify relevant moderators
(Van Lissa, Van Erp, & Clapper, 2023). However, accounting for moderators requires a
relatively high number of observations per moderator, which may not be available.

Each of the aforementioned approaches makes different assumptions about the nature 71 of heterogeneity (see "Models for meta-analysis" in Van Lissa, 2020). A crucial 72 shortcoming of existing research synthesis methods is that these assumptions may not be 73 tenable when meta-analyzing studies that investigate the same informative hypothesis, but are otherwise very heterogeneous. The situation may arise where each study is uniquely 75 identified by a combination of linearly dependent moderators. In this case, it is no longer possible to synthesize effect sizes while accounting for heterogeneity using statistical methods. It is still possible, however, to quantify the support these studies provide for the underlying informative hypothesis. To this end, Bayesian evidence synthesis (BES) aggregates the evidence for a theoretical relationship across studies, without imposing assumptions about heterogeneity (Kuiper, Buskens, Raub, & Hoijtink, 2013). Although 81 this assumption is not necessary, for the sake of simplicity we assume that this theoretical relationship is evaluated via informative hypothesis  $H_i$  in all studies.

The amount of evidence for a hypothesis can be expressed as a Bayes factor, or BF.

The BF can be interpreted as the ratio of evidence for one hypothesis relative to another

hypothesis. Within the scope of this paper, all Bayes factors are the ratio of evidence for

an informative hypothesis  $H_i$  relative to its complement  $H_{!i}$  (see Gu, Mulder, & Hoijtink,

2018). The subscript ! represents the negation operator; in other words,  $H_{!i}$  means "not

 $H_i$ ". This Bayes factor, which we will refer to as  $BF_c$ , represents the ratio of evidence for  $H_i$  divided by evidence against it. A value of  $BF_c = 10$  means that the data provide ten times more support for the hypothesis than against it.

When multiple studies each provide evidence for  $H_i$  in the form of complement Bayes factors, these Bayes factors can be synthesized across studies by taking their product (Kuiper et al., 2013). The resulting product Bayes factor (PBF) summarizes the total evidence for the hypothesis. The only assumption of the PBF is that all study-specific hypotheses provide evidence about the same underlying theoretical relationship. Note that other approaches to BES exist; for instance, it is possible to use the posterior of one study as the prior for a replication study, and thus accumulate evidence across studies (see Heck et al., 2022). Such applications are out of scope of the present paper, which addresses the PBF approach to BES.

Although meta-analysis and BES are both research synthesis methods, they answer 101 different research questions. Meta-analysis estimates the point estimate or distribution of a 102 population effect size. It pools estimates of this effect size across multiple studies to obtain 103 an overall estimate of the effect size. It thus answers questions like: Given certain 104 assumptions about between-studies heterogeneity, what is the average population effect size? BES, on the other hand, aggregates evidence for an informative hypothesis across 106 multiple studies. It thus answers the question: Do all these studies support the hypothesis of interest? Both methods are appropriate for different research questions, and provide 108 complementary information. 109

This paper introduces the first implementation of BES in user-friendly free open source software. A function pbf() was contributed to the bain R-package for Bayesian informative hypothesis evaluation, version 0.2.9. This paper presents a simulation study to validate the method and benchmark it against alternative evidence synthesis methods.

It additionally illustrates several use cases through reproducible examples.

115

## Simulation study

The present simulation study set out to validate the PBF algorithm and benchmark 116 it against other evidence synthesis methods. We simulated a scenario where an informative 117 hypothesis about a correlation between two variables was measured across several 118 independent samples, and the resulting evidence was synthesized across samples using 119 multiple methods. The informative hypothesis, set to be equal across studies, was 120  $H_i: \rho > .1$ . To examine the performance of the different evidence synthesis methods in a 121 range of scenarios, several design factors were manipulated. First was the presence or 122 absence of a true population effect. Given the informative hypothesis of  $H_i: \rho > .1$ , the 123 presence of a true population effect was defined as  $\rho = .2$  and a null effect was defined as 124  $\rho = .1$ . The second design factor was the number of observations per sample 125  $n \in (50, 200, 500, 800)$ . These values were chosen because they correspond to a statistical 126 power to reject a false null hypothesis of  $\beta \in (.10, .30, .60, .80)$  power, respectively, 127 assuming  $\alpha = .05$  and a known effect size of  $\rho = .1$  (Cohen, 1988). Third, we manipulated the number of independent samples (or: replication studies),  $k \in (2,3,10)$ . Fourth, the 129 reliability of the two correlated variables was varied between  $\alpha \in (0.6, 0.8, 1.0)$  to range 130 from questionable to perfect reliability (Nunnally & Bernstein, 2017). Questionable 131 reliability is the lowest level considered to be acceptable in social scientific research, and 132 perfect reliability is what is assumed when analyzing correlations between observed items 133 or scale scores. For all unique combinations of these design factors, the simulation was 134 repeated 1000 times. 135

## 136 Algorithms

The main algorithm of interest was the PBF. As a decision criterion to conclude that  $H_i$  was supported over its complement, we used PBF > 3 - a conventional threshold for
inference using Bayes factors (Jeffreys, 1998).

As a benchmark for comparison, we included several other algorithms that might 140 feasibly be used by researchers who intend to examine whether a hypothesis is true across 141 several independent samples. The first benchmark was vote counting: counting the number 142 of significant effects. Although this method is still in use for aggregating conceptual 143 replications, it is considered bad practice. Three disadvantages are that vote counting 144 disregards sample size, reduces statistical power, and does not quantify the strength of the 145 evidence (Hedges & Olkin, 1980). Our vote counting algorithm summed the number of 146 one-sided z-tests of a null hypothesis corresponding to the informative hypothesis, so 147  $H_0: \rho = .1$  and  $H_a: \rho > .1$ , which corresponds to  $H_i$ . The decision criterion was that the 148 hypothesis was supported in the majority of samples. Thus, for example, if  $H_0$  was rejected 149 in three out of five samples, our vote counting algorithm would find overall support for  $H_a$ 150 (and, by extension,  $H_i$ ).

The second benchmarking algorithm was random-effects meta-analysis (RMA), which is the current gold standard for evidence synthesis (Viechtbauer, 2010). For this algorithm, the null-hypothesis was rejected if a 90% confidence interval excluded the hypothesized value under  $H_i$ . Note that a 90% confidence interval corresponds to a test at  $\alpha = .05$ , because all effects in the simulation are directional.

The third benchmarking algorithm was individual participant data (IPD)

meta-analysis (Riley, Lambert, & Abo-Zaid, 2010). Like classic meta-analysis (RMA), IPD

is a multilevel model, clustered by sample. IPD uses the raw data, which makes it possible

to estimate variance at the first level. By contrast, RMA treats the first-level variance as

known. Note that the PBF can be estimated using either sufficient statistics (as in

meta-analysis) or using raw data (as in IPD). With this in mind, it is informative to

benchmark it against both of these methods. Just as for RMA, a 90% confidence interval

was used for inference.

## 65 Performance indicators

For each algorithm, inferential decisions made using the criteria described above were 166 compared to the population status of the hypothesis (true or false). The resulting 167 confusion matrix gives the number of decisions that were true positives (TP), true 168 negatives (TN), false positives (FP), and false negatives (FN). These quantities were 169 summarized as sensitivity,  $\frac{TP}{TP+FN}$ , the ability to detect an effect given that it was indeed 170 true in the population, and specificity,  $\frac{TN}{TN+FP}$ , the ability to correctly conclude that the 171 informative hypothesis is not supported, given that it was indeed false in the population. 172 The overall performance was captured by the accuracy, which represents the total 173 proportion of correct (true positive and true negative) decisions,  $\frac{TP+TN}{TP+TN+FP+FN}$ . 174

175 Results

We examined overall model performance across conditions. PBF had a higher overall accuracy than other algorithms followed by IPD, then RMA, and finally VC, see Table 1.

This higher accuracy was primarily driven by PBF's greater sensitivity to detect a true effect compared to other algorithms. However, PBF had a lower specificity compared to all other algorithms. This suggests that the PBF trades a loss of specificity for increased sensitivity.

## 182 Effect of simulation conditions

We used ANOVAs to examine the effect of simulation conditions on overall accuracy.

The differences between algorithms were analyzed in analyses that included two-way interactions between design factors and algorithm. As the sample size was very large, significance tests were uninformative. We thus focused on interpreting the effect sizes of the design factors. The performance of PBF was most impacted by sample size n, followed by the number of groups k, and reliability. The differences in the effects of sample size and

number of groups were relatively small between PBF and the two best practice algorithms,
RMA and IPD - but substantial between PBF and the suboptimal VC algorithm. The
reverse pattern occurred for reliability, however: it showed a substantial difference in effect
between PBF and the two best practice algorithms only.

Effect of sample size. Figure 1 indicates that for PBF, both sensitivity and specificity increased with sample size. The other algorithms showed only increasing sensitivity; specificity was limited by a ceiling effect. This difference explains the effect of sample size on the difference between algorithms (see Table 2).

Effect of number of samples. Figure 2 indicates that PBF showed increasing specificity at higher levels of k, while sensitivity was relatively unaffected. RMA and IPD showed a similar pattern, although their specificity was at a ceiling. Only VC showed decreasing sensitivity with an increasing number of groups; this is because the probability of obtaining false negatives increases with the number of groups (as also noted by Hedges & Olkin, 1980). This difference in pattern of effects explains why number of samples had a moderate effect on the difference between algorithms (see Table 2).

Effect of reliability. Figure 3 indicates that the PBF traded sensitivity for specificity. At low levels of reliability, specificity exceeded sensitivity; at high levels of reliability, this pattern was reversed. The other algorithms did not show this pattern, as their specificity was at a ceiling. Their sensitivity increased with higher reliability, however, and therefore, so did their overall performance. This difference in pattern of effects explains why reliability had a moderate effect on the difference between algorithms (see Table 2). Note that, whereas the overall accuracy of the PBF was found to be less susceptible to reliability as compared to other algorithms, this finding masked the trade-off between reliability and specificity.

213 Discussion

This simulation study examined the performance of the product Bayes factor (PBF) 214 as compared against three other common methods for evidence synthesis (IPD, RMA, and 215 VC). The results showed that PBF had a higher overall accuracy than the other 216 algorithms, primarily due to its greater sensitivity to detect a true effect. PBF had lower 217 specificity, however, suggesting that it trades specificity for increased sensitivity. PBF's 218 performance was most impacted by sample size, followed by the number of groups and 219 reliability. Even at the smallest sample size of n = 20, PBF had a superior sensitivity to the other algorithms. This suggests that PBF is more suitable than other methods as a small sample solution. When the number of samples increased, most algorithms showed approximately stable specificity and increasing sensitivity. A notable exception was vote counting; its sensitivity decreased with an increasing number of samples, as has been 224 previously documented (Hedges & Olkin, 1980). With increasing reliability, PBF showed a 225 substantially different pattern of results than the two other best practice algorithms (RMA) 226 and IPD). Although the overall effect of reliability on overall accuracy was smaller for PBF 227 than for other algorithms, this masks a trade-off between reliability and specificity. 228 Whereas most algorithms showed near-stable specificity and increasing sensitivity, PBF 220 showed a clear trade-off between decreasing specificity and increasing sensitivity. This 230 implies that the PBF is more conservative - less likely to detect an effect - in the presence 231 of increasing measurement error. 232

These results have important implications for applied evidence synthesis. For
example, the finding that PBF had a higher overall accuracy due to greater sensitivity
suggests that it may be a better choice than the other algorithms, particularly when
detecting true effects has high priority. However, researchers should be aware that this
increased sensitivity comes at a loss of specificity, which incurs a greater risk of false
positive results. If specificity is a higher priority, other algorithms such as IPD or RMA

239 may thus be more appropriate.

The present study also has some limitations. First, the simulation study makes 240 specific assumptions that may not generalize to all real-world applications. A second 241 important caveat is that most of the algorithms did not reach a level of sensitivity that is 242 considered acceptable from a perspective of statistical power (i.e., greater than .80, J. 243 Cohen, 2013). PBF performed notably better than other algorithms, but its sensitivity still 244 fell below .80 in many conditions. Low power increases the risk of false negatives, or failing 245 to detect a true effect. One reason power was low is that, in conditions where a true effect 246 was present, its value only exceeded the boundary value of the informative hypothesis by .1 247 points. Such small effects are hard to detect. All algorithms will likely perform better when 248 the true effect is larger. The low sensitivity of all algorithms highlights the importance of 249 reticence when interpreting evidence syntheses of studies with small samples and small 250 effect sizes. It may be prudent to avoid generalizing such results to the population, and 251 instead consider them as merely descriptive of the published research. Additionally, 252 sensitivity analyses can be used to assess the robustness of the results to different modeling 253 assumptions and methods.

A third limitation is that the evidence synthesis methods compared here represent different approaches to inference and answer different research questions. Since each of these methods is optimized for a different purpose, the present study should not be considered as a comprehensive assessment of their strengths and weaknesses. We nonetheless compare them because of their similar usage in evidence synthesis. It is up to individual researchers to choose an appropriate method, guided by the research question and the available information. For instance, when raw data is unavailable, IPD cannot be used, and when parameter estimates or effect sizes are not reported, only VC can be used.

Aside from the aforementioned fact that the PBF answers a different research question than the other algorithms, it is worth noting limitations of the interpretation of

the PBF. The PBF renders support for one specific informative hypothesis versus its 265 complement. If the informative hypothesis is supported, this does not necessarily mean 266 that it is also true. Consider the hypothetical example that the informative hypothesis that 267 the earth is flat was supported with BF = 3.01. Although the data support this hypothesis 268 over its complement, the hypothesis is clearly wrong (the earth is spherical). If we would 269 have evaluated another hypothesis, e.g., the earth is shaped like an American football, it 270 would have received much more support, e.g. BF = 1000, even though it is also wrong. A 271 high Bayes factor thus does not mean that the hypothesis is true. Conversely, a low Bayes 272 factor merely indicates that the informative hypothesis is not supported, and does not 273 provide information about the true state of affairs. A related limitation is that our 274 simulation study used an arbitrary - albeit conventional - threshold for inference (Jeffreys, 275 1998). In applied research, it is more sensible to evaluate the weight of evidence, rather than resorting to a rule of thumb. 277

Tutorial

This tutorial demonstrates how to synthesize evidence for an informative hypothesis 279 across heterogeneous replications using the Product Bayes Factor (PBF). We assume that 280 users have installed the free open source statistical programming language R (R Core 281 Team, 2021). The R-package bain version 0.2.9 or later is required, which can be installed 282 by running install.packages("bain") in the R console. The data used in this tutorial 283 are included in the bain package, and have been simulated based on the data presented in 284 (Leeuwen, Van Lissa, Papakonstantinou, Petersen, & Curry, 2022). A more detailed 285 description of the datasets is found in (Leeuwen et al., 2022); additionally, the dataset documentation is accessed by running ?synthetic us, ?synthetic dk or ?synthetic nl in the R console. Van Leeuwen and colleagues conducted a theory-driven, preregistered 288 study to address the research question whether political orientation and moral dispositions 289 are associated. Suitable data were collected in three countries: the United states of 290

America, Denmark, and the Netherlands. Each sample contained multiple measures of 291 political orientation and moral dispositions. In the original publication, the PBF was used 292 to aggregate evidence across scales and countries to obtain an overall measure of support 293 for the central hypothesis. This tutorial follows the same rationale, but uses only one effect 294 size per sample, and varies the way this effect size is computed to illustrate the more typical 295 use case where the same informative hypothesis has been studied in different ways in 296 multiple studies. We will examine the informative hypothesis that self-reported importance 297 of family morality is positively associated with a conservative socio-political orientation. 298 We load the bain library and assign the data to three objects with convenient names: 290

```
library(bain)

NL <- synthetic_nl

DK <- synthetic_dk

US <- synthetic_us</pre>
```

How to use bain. We briefly introduce the basic use of the bain() function, and
how to interpret its output. We must estimate a model suitable for evaluating our
informative hypothesis. Because both scales consist of multiple items, we can use structural
equation modeling (SEM) to perform latent variable regression (see Van Lissa et al., 2020):

```
con ~ beta * fam"

# Estimate the model in lavaan

results_nl <- sem(model = model_nl, data = NL)</pre>
```

The informative hypothesis in this tutorial is  $H_i: \beta > .1$ , where  $\beta$  (beta) is the 304 standardized regression coefficient. Instead of a conventional null hypothesis,  $H_0: \beta = 0$ , the value of .1 was used as a minimal effect size of interest. The code below illustrates how 306 to obtain a Bayes factor for this informative hypothesis, using the output of the SEM 307 analysis above. We can refer to the parameter beta by name because we labeled it in the 308 lavaan syntax; if we had not done so, we could find the names of all model parameters by 309 running get estimates (results nl, standardize = TRUE). The results indicate that 310 the hypothesis is supported when compared to its complement. For a more in-depth 311 tutorial on bain(), see Hoijtink, Mulder, Lissa, and Gu (2019), and for further guidance on 312 the use of bain() for SEM, see Van Lissa et al. (2020). 313

```
# Test that the effect labeled 'beta' is positive
bf_nl <- bain(results_nl, hypothesis = "beta > .1", standardize = TRUE)
bf_nl
```

```
## Bayesian informative hypothesis testing for an object of class lavaan:
314
   ##
315
   ##
          Fit
                       BF.u
                              BF.c
                                      PMPa
                                            PMPb
                                                   PMPc
                 Com
   ## H1 0.959 0.500 1.918 23.246 1.000 0.657 0.959
317
   ## Hu
                                             0.343
318
   ## Hc 0.041 0.500 0.082
                                                   0.041
319
   ##
320
   ## Hypotheses:
321
```

```
322 ## H1: beta>.1
323 ##
```

 $_{324}$  ## Note: BF.u denotes the Bayes factor of the hypothesis at hand versus the unconstrained

Aggregating evidence across studies. As mentioned before, suitable data were collected to evaluate the substantive hypothesis in three countries. There are differences between countries that prevent analyzing these data as a multilevel model, however. For instance, conservatism was measured using different scales. This is an appropriate situation to use the PBF to aggregate evidence across countries. Below, we estimate a latent regression model for the remaining two countries, taking care to use the same label for the parameter of interest in all samples. Then, we bind all three SEM-models in a list, and call PBF to evaluate the hypothesis of interest on all models and aggregate the evidence. As the BF in all three samples is positive, the resulting PBF is very large. We can thus conclude that the central hypothesis receives overwhelming support across samples.

```
# Specify the models for DK and US
model_dk <- "
fam =~ fam_1 + fam_2 + fam_3
con =~
sepa_soc_1 + sepa_soc_2 + sepa_soc_3 + sepa_soc_4 + sepa_soc_5 +
sepa_eco_1 + sepa_eco_2 + sepa_eco_3 + sepa_eco_4 + sepa_eco_5
con ~ beta * fam"
model_us <- "
fam =~ fam_1 + fam_2 + fam_3
con =~
secs_soc_1 + secs_soc_2 + secs_soc_3 + secs_soc_4 + secs_soc_5 +
secs_soc_6 + secs_soc_7 +
secs_eco_1 + secs_eco_2 + secs_eco_3 + secs_eco_4 + secs_eco_5</pre>
```

```
# Estimate the model in lavaan
results_dk <- sem(model = model_dk, data = DK)
results_us <- sem(model = model_us, data = US)

# Bind the models into a list
results <- list(results_nl, results_dk, results_us)
# Test the hypothesis that the effect size labeled 'beta' is positive
pbf(results, hypothesis = "beta > .1", standardize = TRUE)
```

```
PBF Sample.1 Sample.2 Sample.3

## H1: beta>.1 1.013928e+27 23.24645 1.903063e+12 2.291908e+13
```

Using bain objects. The pbf() function also accepts multiple bain objects. This
makes it possible to, for example, evaluate different sets of hypotheses on different data sets
before using the resulting bain objects to aggregate the evidence for all common hypotheses
across datasets. The example below illustrates this use case. As before, all analyses share
one hypotheses in common  $(H_i: \beta_{fam} > .1)$ , but the Dutch sample now contains a
sample-specific hypothesis regarding the effect of group morality, namely that  $\beta_{grp} < .1$ .
The pbf() function is called on a list of bain objects. Note that, in this case, pbf() does
not require an argument hypothesis, as the hypotheses are contained in the bain objects.

```
PBF Sample.1 Sample.2 Sample.3

## H1: beta>.1 2845266 1301.179 985.1867 2.219562
```

As can be seen, the results are equivalent to the results in the previous example. The sample-specific hypothesis has been left out, and common hypotheses are retained and aggregated. If there are no common hypotheses across all objects, pbf() throws an error.

Using sufficient statistics. A third use case occurs when the raw data from
different samples are not available. This may happen, for example, when aggregating
findings from the published literature (similar to meta-analysis). In this case, one can use
the default interface of bain, as explained in (Hoijtink et al., 2019). This function requires
four arguments: A named vector of parameter estimates, their asymptotic covariance
matrix, the original sample size, and the number of within-group and between-group

parameters. Note that, when analyzing a single parameter per sample, the standard error is sufficient to construct the asymptotic covariance matrix. Thus, this method can be applied to data that have been prepared for classic meta-analysis (effect sizes and their sampling variances). Importantly, unlike meta-analysis, the present method is suitable for conceptual replications. It does not require uniform effect size measures across studies.

The example below illustrates how to aggregate evidence for one hypotheses across three studies that each used different methods.

The present use case evaluates the following hypothesis: There is a positive

association between family morality and political conservatism. This conceptual hypothesis

is evaluated differently in the three samples, resulting in three different types of statistics

and distinct sample-specific hypotheses:

- 1. A t-test was performed using the NL data; using Cohen's D gives  $H_i^{NL}: \delta_{conservative>liberal} > 0, \text{ where } \delta \text{ is the mean difference between groups.}$
- 2. A bivariate regression coefficient was calculated using the DK data, giving  $H_i^{DK}:\beta_{fam}>0$
- 37. A correlation coefficient was calculated using the US data, giving  $H_i^{US}: \rho_{fam,con} > 0$ ,
  where  $\rho$  is the correlation between family morality and conservatism.

Note that we intentionally manipulate the data to illustrate these different analyses; for example, we compute mean scale scores and dichotomize the continuous conservatism scale to conduct a t-test. We do not advocate these practices for applied research.

First we obtain the relevant parameter estimates and their sampling variances, which allows us to evaluate the specific hypotheses in bain:

```
# Create mean scale scores

NL <- data.frame(
    family = rowMeans(NL[c("fam_1", "fam_2", "fam_3")]),</pre>
```

```
conservative = rowMeans(NL[c("sepa soc 1", "sepa soc 2", "sepa soc 3",
                                "sepa soc 4", "sepa soc 5", "sepa eco 1",
                                "sepa eco 2", "sepa eco 3", "sepa eco 4",
                                "sepa eco 5")]))
DK <- data.frame(</pre>
  family = rowMeans(DK[c("fam_1", "fam_2", "fam_3")]),
  conservative = rowMeans(DK[c("sepa_soc_1", "sepa_soc_2", "sepa_soc_3",
                                "sepa soc 4", "sepa_soc_5", "sepa_eco_1",
                                "sepa eco 2", "sepa_eco_3", "sepa_eco_4",
                                "sepa eco 5")]))
US <- data.frame(</pre>
  family = rowMeans(US[c("fam 1", "fam 2", "fam 3")]),
  conservative = rowMeans(US[c("secs_soc_1", "secs_soc_2", "secs_soc_3",
                                "secs_soc_4", "secs_soc_5", "secs_soc_6",
                                "secs soc 7", "secs eco 1", "secs eco 2",
                                "secs eco 3", "secs eco 4", "secs eco 5")]))
# NL: Conduct t-test using Cohen's D
NL$group <- cut(NL$conservative, breaks = 2,
                labels = c("liberal", "conservative"))
sample_sizes <- table(NL$group)</pre>
sds <- tapply(NL$family, NL$group, sd)</pre>
pooled_sd <- sqrt(sum((sample_sizes - 1) * sds) / (sum(sample_sizes) - 2))</pre>
NL est <- diff(tapply(NL$family, NL$group, mean)) / pooled sd
NL var <- (sum(sample sizes) / prod(sample sizes)) +
  (NL est^2 / (2*sum(sample sizes)))
```

```
# DK: Conduct bivariate regression

DK_fit <- lm(conservative ~ family, data = DK)

DK_est <- coef(DK_fit)["family"]

DK_var <- vcov(DK_fit)["family", "family"]

# US: Correlation coefficient

US_est <- cor(US)[1, 2]

US_var <- (1 - US_est^2)^2 / (nrow(US) - 1)

# Name the estimates so hypotheses will be the same

names(NL_est) <- names(DK_est) <- names(US_est) <- "parameter"</pre>
```

Then, we use bain.default() to evaluate the central hypothesis on each parameter estimate. The pbf() function can be called on a list of the resulting bain objects.

```
Sigma = matrix(US var, 1, 1),
                n = nrow(US),
                hypothesis = "parameter > 0",
                joint parameters = 1)
# Aggregate evidence using pbf()
pbf(list(US_bain, DK_bain, NL_bain))
```

```
##
                                 PBF
                                          Sample.1 Sample.2 Sample.3
380
   ## H1: parameter>0 9.873547e+20 2.291908e+13 540773.7 79.66368
381
```

The results suggest substantial evidence for the hypothesis that there is a positive 382 association between family morality and political conservatism. Although each study used a different method to assess this hypothesis, their evidence can be synthesized using pbf().

Conclusion

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In conclusion, this study evaluated the performance of the product Bayes factor as a 386 method for evidence synthesis, and compared it against other commonly used evidence 387 synthesis methods under different simulation conditions. Compared to the other methods, 388 PBF had the highest overall accuracy. This was primarily due to its greater sensitivity. However, PBF had lower specificity than all other algorithms, suggesting a trade-off 390 between sensitivity and specificity. The other algorithms showed ceiling effects in specificity, limiting their sensitivity. The performance of the PBF was most strongly affected by sample size, followed by the number of samples and reliability. We introduced a 393 user-friendly implementation of the PBF in the bain R-package, and demonstrated its use 394 with various analysis techniques in R, as well as with sufficient statistics that are already 395 routinely coded for meta-analysis (i.e., effect sizes and their sampling variance). This 396

means that researchers can now use the PBF to aggregate evidence in situations where 397 classic meta-analytic methods are less suitable. For example, when one informative 398 hypothesis has been evaluated in several replication studies, but these replication studies 399 are quite heterogeneous because they sample from different populations and use different 400 methods or analysis techniques. Especially when the number of replication studies is too 401 small to adequately account for these sources of between-study heterogeneity, the PBF may 402 be a useful method to aggregate evidence for the common informative hypothesis. 403 Researchers should be aware that the PBF trades off increased sensitivity for decreased 404 specificity, and that it addresses a different research question than other research synthesis 405 methods. This highlights the importance of careful interpretation of the results, and 406 consideration of the research question when selecting an aggregation method. In sum, our 407 results suggest that PBF is a useful evidence synthesis method, which is now broadly 408 accessible due to its inclusion in the bain R-package.

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 $\label{eq:marginal} \begin{tabular}{ll} Table 1 \\ Marginal \ confusion \ matrix \ metrics. \\ \end{tabular}$ 

Metric	PBF	IPD	RMA	VC
sensitivity	0.76	0.35	0.32	0.05
specificity	0.76	0.99	0.99	1.00
accuracy	0.76	0.67	0.66	0.52

Table 2

Partial eta squared of the effect of each design factor on accuracy for each algorithm and for the difference between PBF and all other algorithms (e.g., vs RMA).

condition	IPD	RMA	VC	PBF	vs IPD	vs RMA	vs VC
k	0.35	0.40	0.13	0.32	0.01	0.02	0.23
n	0.60	0.58	0.29	0.62	0.01	0.00	0.19
reliability	0.62	0.61	0.23	0.04	0.27	0.25	0.01

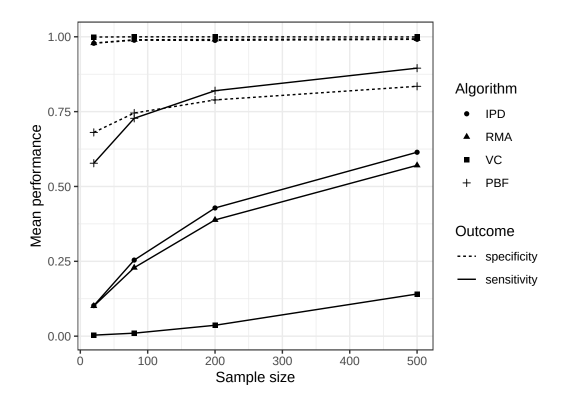


Figure 1. Mean performance by sample size

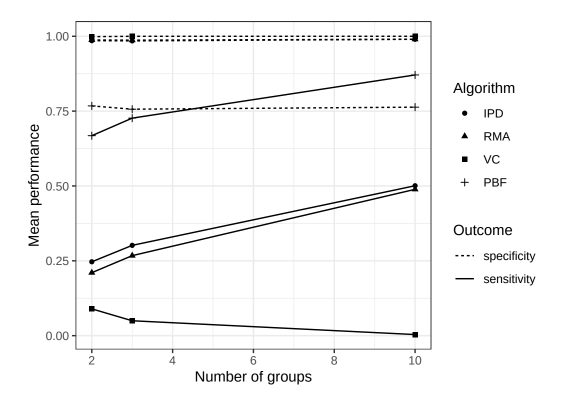


Figure 2. Mean performance by number of groups

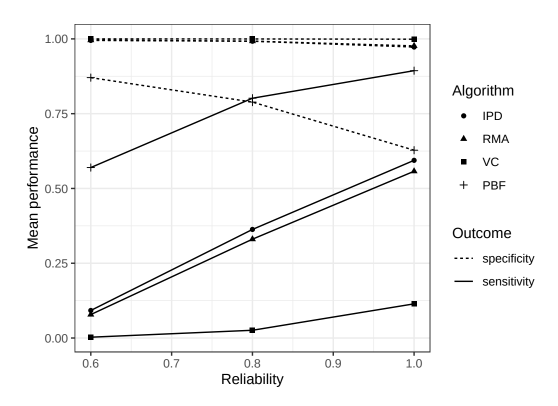


Figure 3. Mean performance by reliability