Minimal Data Needed for Valid & Accurate Image-Based fMRI Meta-Analysis

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Abstract

Meta-analysis provides a quantitative approach to summarise the rich functional Magnetic Resonance Imaging literature (fMRI). Due to the lack of availability of image data supporting existing literature, the majority of fMRI meta-analysis are coordinate-based. However, when image data is available for each study, the optimal approach is to perform an image-based meta-analysis. A number of approaches have been proposed to perform such meta-analyses including combination of standardised statistics, just effect estimates or both effects estimates and their sampling variance. While the latter is the preferred approach in the statistical community, its properties are only guaranteed in large samples. Additionally, often only standardised estimates are shared, reducing the possible meta-analytic approaches. Finally, because the BOLD signal is non-quantitative care has to be taken in order to insure that effect estimates are expressed in the same units, especially when combining data from different software packages. Given the growing interest in data sharing in the neuroimaging community there is a need to identify what is the minimal data to be shared in order to allow for future image-based meta-analysis. In this paper, we compare the validity and the accuracy of nine meta-analytic approaches on simulated and real data.

Keywords: Meta-analysis, Neuroimaging, Mixed-effects

1. Introduction

Neuroscience research studies typically exhibit low power and therefore have low likelihood that a given result is True [3]. Under low power, both true effects are less likely to be detected and, when an effect is detected, it is less likely to be true [3]. Many authors have advocated for increased sample sizes in neuroimaging (e.g. [24]) which would mechanically increase the power. Aware of this issue, our community has started a couple of years ago to generated large datasets by means of consortia (e.g. ADDREF 1000FC) or dedicated datasets (e.g. ADDREF UK biobank) which are starting to show promising results. Another approach to increase sample size, is to combine the results of previous studies through meta-analyses. Functional Magnetic Resonance Imaging (fMRI), with its ample literature of more than 100 000 articles [ADDREF] is particularly well-suited for this approach. – use mature field with robust well-established methods

Most fMRI meta-analyses are 'coordinate-based", i.e. they proceed by combining information about the locations of local maxima (and optionnally their statistic value). But, the best practice method is an Image-Based Meta-Analysis (IBMA) that combines information from each study at every location in space [1], effectively enabling the discovery of consistent sub-threshold effects that could not be uncover using coordinates only. For a long time, the standard in the fMRI literature was to report a table with coordinates and statistic values of each local maxima. IBMAs were very limited as they required contacting back the original authors of each study of interest to retreive the summary statistic images, an approach that is both time-consuming and inefficient (ADD REF on data availability). With the arrival of tools that facilitate sharing 3D volumes obtained as a result of a statistical analysis, such as NeuroVault [8], IBMAs becomes more and more easy to perform.

IBMAs can be performed with three different types of summary data per study:

- 1. the contrast estimates and contrast variance estimates (E+S).
- 2. the contrast estimates (E).
- 3. the standardized statistic maps (Z).

But all IBMA methods have specific limitations. The first option (E+S), which is considered best practice, leading to more statistically efficient estimates [5], requires the contrasts to be expressed with in the same units. In fMRI, units will depend on the field strength [4] as well as data, model and contrast vector scaling [18]. Also, while often neglected, the statistical methods for this first option rely on asymptotic, large sample results and may have poor performance for

Figure 1: False positive rates of the meta-analytic estimators under the null hypothesis for p < 0.05.

typical number of studies [ADDREF, ADDREF]. The second option (E) assumes homoscedasticity, i.e. constant variance of the study-level contrast estimates, an assumption that is difficult to atteign in meta-analysis where the number of subjects is likely to vary across studies. Finally the third option (Z) corresponds to historical methods that typically assume homogeneity, i.e. no between-study variance.

Given the growing interest in data sharing in the neuroimaging community [24, 19], there is a need to identify what is the best approach for neuroimaging IBMAs and what is the minimal data needed. Here, we compared the use of IMBA using 9 meta-analytic approaches: 2 approaches use E+S's, 2 E's only and 5 Z's. We assessed and compared validity using simulated data under violation of each method's underlying assumptions: in small samples, under heteroscedasticity, under heterogeneity or with contrast using mismatched units. We estimated the accuracy using a real dataset of 21 studies of pain.

2. Methods

2.1. Theory

For study i = 1, ..., k we have contrast estimate $\hat{\beta}_i$, its contrast variance estimate s_i^2 (i.e. squared standard error), its equivalent Z-statistic map Z_i and its sample size n_i .

Combining contrast estimates and their standard error. The recommanded approach is to fit contrast estimates and their standard error with a hierarchical general linear model (GLM) [5], creating a third-level (level 1: subject; level 2: study; level 3: meta-analysis). The general formulation for the study-level data is a meta-regression:

$$\hat{\boldsymbol{\beta}} = X\boldsymbol{\gamma} + \boldsymbol{\epsilon} \tag{1}$$

where $\hat{\boldsymbol{\beta}} = [\hat{\beta}_1 \dots \hat{\beta}_k]^T$ is the vector of contrast estimates, X is the $k \times p$ study-level matrix. While the design matrix X can take any form, typically a one-sample model is used with p = 1 and X comprised of a column of ones. $\boldsymbol{\gamma} = [\gamma_1 \dots \gamma_p]^T$ is a vector of meta-analytic parameters to estimate and $\epsilon \sim \mathcal{N}(0, W)$ is the residual error term.

In the most general case of a random-effects meta-analysis, i.e. assuming non-zero between-study variance, we have $W = \operatorname{diag}(\sigma_1^2/n_1 + \tau^2, \dots, \sigma_k^2/n_k + \tau^2)$ where τ^2 denotes the between-study variance and σ_i^2/n_i denotes the variance of the contrast estimate for study i. Eq. (1) can be solved by weighted least squares giving:

$$\hat{\boldsymbol{\gamma}} = (X^T W^{-1} X)^{-1} X^T W^{-1} \hat{\boldsymbol{\beta}} \tag{2}$$

$$\hat{\gamma} = (X^T W^{-1} X)^{-1} X^T W^{-1} \hat{\beta}$$

$$Var(\hat{\gamma}) = (X^T W^{-1} X)^{-1}$$
(3)

Table S1 and S2 provides the WLS estimates and their sampling variances for one and two-sample tests. But in practice, the weight matrix W is unknown and has to be estimated from the data. Given \hat{W} a consistent estimate of W, the feasible generalized least squares (FGLS) estimator is computed as:

$$\hat{\boldsymbol{\gamma}} = (X^T \hat{W}^{-1} X)^{-1} X^T \hat{W}^{-1} \hat{\boldsymbol{\beta}} \tag{4}$$

$$\hat{\gamma} = (X^T \hat{W}^{-1} X)^{-1} X^T \hat{W}^{-1} \hat{\beta}$$

$$Var(\hat{\gamma}) = (X^T \hat{W}^{-1} X)^{-1}$$
(5)

Here we use s_i^2 as an estimate of σ_i^2/n_i and estimate $\hat{\tau}^2$ from the data. FGLS is asymptotically efficient but its finite sample properties are unknown [9]. We used the 'FLAME 1' implementation available in FSL that provides inference by comparing the statistic to a Student distribution with k-p degrees of freedom [34] as depicted in Table 1. This reference approach will be referred to as Mixed-effects (MFX) GLM.

In a fixed-effects meta-analysis, i.e. assuming no or negligible between-study variance, we have $W = \operatorname{diag}(\sigma_1^2/n_1, \dots, \sigma_k^2/n_k)$. We obtain a feasible generalied least squares estimate by approximating σ_i^2/n_i by s_i^2 . We used the 'Simple OLS' implementation available in FSL that provides inference by comparing the statistic to a Student distribution with $(\sum_{i=1}^{k} (n_i - 1)) - p$ degrees of freedom [29] as depicted in Table 1. This reference approach will be referred to as Fixed-effects (FFX) GLM.

	Meta-analysic statistic	Nominal H_0 distrib.	Inputs	Assumptions
MFX GLM	$\left(\sum \kappa_i \hat{\beta}_i\right) / \sqrt{\sum_{i=1}^k \kappa_i} \text{ with } \kappa_i = 1/(\hat{\tau}^2 + s_i^2)$	\mathcal{T}_{k-1}	$\hat{\beta}_i, s_i^2$	IGE, large sample.
RFX GLM	$\left(\sum_{i=1}^k rac{\hat{eta}_i}{\sqrt{k}} ight)/\widehat{\sigma_C^2}$	\mathcal{T}_{k-1}	\hat{eta}_i	IGE; $\tau^2 + \sigma_i^2 = \sigma_C^2 \ \forall i$
Contrast Perm.	$\left(\sum_{i=1}^k rac{\hat{eta}_i}{\sqrt{k}} ight)/\widehat{\sigma_C^2} \ \left(\sum_{i=1}^k rac{\hat{eta}_i}{\sqrt{k}} ight)/\widehat{\sigma_C^2}$	Empirical	$\hat{\beta}_{\pmb{i}}$	ISE.
FFX GLM	$\left(\sum_{i=1}^k rac{\hat{eta}_i}{s_i^2} ight)/\sqrt{\sum_{i=1}^k 1/s_i^2}$	$\mathcal{T}_{(\sum_{i=1}^k n_i-1)-1}$	$\hat{\beta}_i, s_i^2$	$\tau^2 = 0$, large sample.
Fisher	$-2\sum \ln P_i$	$\chi^2_{(2k)}$	Z_i	$\tau^2 = 0$
Stouffer	$\sqrt{k} imesrac{i}{k}\sum_{i}Z_{i}$	$\mathcal{N}(0,1)$	Z_i	$\tau^2 = 0$
Weighted Z	$rac{1}{\sqrt{\sum_i n_i}} \sum_i \sqrt{n_i} Z_i \ \left(\sum_{i=1}^k Z_i\right) / \sqrt{k} \hat{\sigma} \ \left(\sum_{i=1}^k Z_i\right) / \sqrt{k}$	$\mathcal{N}(0,1)$	Z_i, n_i	$\tau^2 = 0$
Z RFX	$\left(\sum_{i=1}^k Z_i\right)/\sqrt{k}\hat{\sigma}$	\mathcal{T}_{k-1}	Z_i	IGE; $1 + \tau^2/\sigma_i^2$ cst.
Z Perm.	$\left(\sum_{i=1}^k Z_i\right)/\sqrt{k}$	Empirical	Z_i	ISE.

Table 1: Statistics for one-sample meta-analysis tests and their sampling distributions under the null hypothesis H₀. Empirical null distributions are determined using permutations with sign flipping. IGE=Independent Gaussian Errors, ISE=Independent Symmetric Errors. Note: P_i = $\Phi(-Z_i)$, σ_C^2 is the unbiased sample variance.

Combining contrast estimates. If the s_i^2 are unavailable, the contrast estimates $\hat{\beta}_i$ can be combined by assuming that the within-study contrast variance σ_i^2/n_i is constant $(\sigma_i^2/n_i = \sigma^2 \ \forall i)$ or negligible in comparison to the between-study variance $(\sigma_i^2/n_i \ll \tau^2)$. Then $W = \text{diag}(\sigma_C^2, \ldots, \sigma_C^2)$ where σ_C^2 combines the within and between-study variances, i.e. $\sigma_C^2 \simeq \tau^2$ or $\sigma_C^2 \simeq \tau^2 + \sigma^2$ (note, however, in this setting we do not separately estimate τ^2 or σ^2). Under these assumptions, Eq. (1) can be solved by ordinary least squares giving:

$$\hat{\gamma} = (X^T X)^{-1} X^T \hat{\beta} \tag{6}$$

$$\hat{\gamma} = (X^T X)^{-1} X^T \hat{\beta}$$

$$\text{Var}(\hat{\gamma}) = (X^T X)^{-1} \sigma_C^2$$
(6)

Given $\hat{\sigma}_C^2$ the unbiased sample variance, we obtain the statistics presented in Table 1 for one sample tests. This approach will be referred to Random-effects (RFX) GLM in the following. Inference can be carried out by comparing the RFX GLM statistic to a Student distribution with k-1 degrees of freedom, this result holds asymptotically as well as in small samples [9].

As an alternative to parametric approaches, non-parametric inference [11, 20] can be performed by comparing the one-sample RFX GLM T-statistic to the distribution obtained with "sign flipping", i.e. randomly multiplying each study's data by 1 or -1, justified by an assumption of independent studies and symmetrically distributed random error. For two-sample tests, the non-parameteric distribution can be obtained by random permutation of the group labels. This approach will be referred to as **Contrast permutation**.

Combining standardised statistics. When only test statistic images are available there are a several alternative approaches available. Fisher's meta-analysis provide a statistic to combine the associated p-values [6]. Stouffer's approach combines directly the standardised statistic [28]. In [36] following [15], the author proposed a weighted method that weights each study's Z_i by the square root of its sample size [3,7]. This approach will be referred to as Weighted Z. All these meta-analytic statistics assumes no or negligible between-study variance and are only suited for one-sample tests. The corresponding statistics and nominal null distributions are presented in Table 1.

As suggested in [1], to get an RFX-like method based on Stouffer's approach, the standardised statistical estimates Z_i can be combined in an OLS analysis. The corresponding estimate, referred as **Z** RFX is also provided in 1. Nonparametric inference [11, 20] can also be obtained by sign flipping on the Z_i 's. This approach will be referred to as **Z** permutation.

2.2. Approximations

In practice, all the methods are either approximate in small samples or rely on assumptions that might not be verified in the context of neuroimaging meta-analyses. Methods based on FGLS (MFX GLM and FFX GLM) have approximate parametric null distributions. The nominal distributions of RFX GLM and two-sample contrast permutations are under the (unrealistic) assumption of homogeneous standard errors over studies; even if all studies are 'clean' and conducted at the same center, variation in sample size will induce differences in s_i^2 's. The fixed-effects methods (Fisher, Stouffer, wieghted Z and FFX GLM) assume homogeneity across studies, i.e. zero between-study variance. All contrast methods (MFX GLM, RFX GLM, Contrast permutation and FFX GLM) require the contrasts to be expressed with in the same units. Finally, RFX Stouffer and Z RFX are ad-hoc methods and their validity has to be investigated.

2.3. Experiments

2.3.1. Validity on null data

We used Monte Carlo simulations to empirically investigate the validity of each estimator. We simulated a set of contrast estimates $\hat{\beta}_i$ and contrast variance estimates s_i^2 according to:

$$\hat{\beta}_i \sim \mathcal{N}(0, \frac{\sigma_i^2}{n} + \tau^2)$$
 (8)

$$s_i^2 \sim \frac{\sigma_i^2}{n} \frac{\chi_{(n-1)}^2}{n-1}$$
 (9)

where σ_i^2 was either constant across studies (homoscedasticity) and taken from $\{n \times [0.25, 0.5, 1, 2, 4]\}$ or varying across studies (heteroscedasticity) such as $\max(\sigma_i^2) = \alpha \min(\sigma_i^2)$ with $\alpha \in \{2, 4, 8, 16\}$ and $\max(\sigma_i^2) = 1$, allowing for 5 different values of σ_i^2 that were linearly spaced and repeated as many times as needed for the specified number of studies k. The between-study variance τ^2 was set to zero (homogeneity) or 1 (heterogeneity). We looked at four different number of studies per meta-analysis $k \in \{5, 10, 25, 50\}$ and two number of subjects $n \in \{20, 100\}$. We set the default number of subjects per studies to n = 20 which is a common sample size in existing neuroimaging studies [23]. A total of 144 parameter sets $(9\sigma_i^2 \times 2\tau^2 \times 4k \times 2n_i)$ was therefore tested and a total of 1 026 000 realisations was performed for each parameter set.

Three types of analyses were computed: a one-sample meta-analysis testing significance of the mean effect in a group of kstudies, a two-sample meta-analysis testing significance in mean differences between two groups of k studies each and an unbalanced two-sample meta-analysis testing significance in mean differences between two groups of $2/5 \times k$ and $8/5 \times k$ respectively. Two-sample analysis assumed a common between-study variance across groups.

We conducted those simulations to evaluate the validity of each estimator in small samples and under violations of their assumptions, such as inhomogeneity of contrast variances s_i^2 or presence of non-negligible between-study variance. Furthermore, we studied the robustness of contrast-based methods to the presence of mismatched units across studies. To simulate units mismatch, each contrast estimates $\hat{\beta}_i$ and contrast variance estimates s_i^2 was replaced by a rescaled version: $\hat{\beta}_i^* = \hat{\beta}_i a_i$ and $s_i^{2*} = s_i^2 a_i^2$. 2 types of unit mismatched were investigated:

- Mismatched contrast vectors: a_i linearly sampled between 0.4 and 1.6 (mean 1).
- Mismatched data scaling algorithms (simulating data from different analysis sofware): either 20% or 50% of the studies were rescaled with a factor $a_i = 2$ other studies keeping the original scaling.

In the case of two-sample tests, the same mismatch was applied to both groups.

The simulations were implemented in Matlab R2016b [26] on a High Permformance Computing cluster. Code is available at: https://github.com/cmaumet/zmeta_buster and depends on the IBMA toolbox cb5aee [16], SPM12 revision 6196 (RRID:SCR_007037) [7], FSL 5.0.10 (RRID:SCR_002823) [13] and SnPM f254ff (RRID:SCR_002092) [20]. Figures were computed in R [25] jupyter notebooks [22] and depends on ggplot2 [32] and cowplot [33]. The code used to compute the figures displayed in this manuscript is available at: https://github.com/cmaumet/zmeta.

2.3.2. Accuracy on real data

We then used Receiver-Operating-Characteristics (ROC) curves to assess the sensibility of each of the nine metaanalytic estimators on real data. We used a dataset of 21 studies investigating pain of which 10 were computed with SPM and 11 with FSL, this dataset is available at: http://neurovault.org/collections/1425/.

Comparability of contrast estimates depends on equivalent scaling of the data, models, and contrast vectors. Data scaling in FSL is performed by setting the median brain intensity to 10,000 while in SPM the mean brain intensity is set to 100. Furthermore, due to imprections of SPM data scaling algorithm, gray matter values tends to be closer to 200 than 100. We therefore rescaled SPM contrast estimates and standard errors by 2 and FSL contrast estimates and standard errors by 100. Furthermore when contrasts were not constructed to preserve units, with sum of positive elements equal to 1, sum of negative elements equal to -1, we further rescale the data to take into account these undesired effects.

We used Neurosynth [35]'s automated large scale coordinate-based meta-analysis of pain as ground truth of areas that should present an effect (cf. http://neurosynth.org/analyses/terms/pain/). For each meta-analytic approach, we estimated the true positive rate over a range of thresholds and combined these values to the false positive rates computed on simulated data to create the ROC curves.

The real data analyses were implemented in Matlab R2016b on a Mac Book Pro laptop. Code is available at: https://github.com/cmaumet/zmeta_rocs. Software dependencies and repositories for figures are identical as for simulated analyses. Real data figures were computed in Python 3.6.0 [30] jupyter notebooks and depends on nibabel [2], Scipy [14], numpy [31], scikit-learn [21] and matplotlib [12].

2.3.3. Presence of heterogeneity and heteroscedasticity in real data

To investigate the presence of between-study variations we used Cochran's Q test of heterogeneity [10]:

$$Q = \sum_{i=1}^{i=k} w_i (\hat{\beta}_i - \hat{\theta})^2$$
 (10)

where $\hat{\theta}$ is the fixed-effects GLM meta-analytic estimate such that $\hat{\theta} = \frac{\sum_{i=1}^{i=k} w_i \hat{\beta}_i}{\sum_{i=1}^{i=k} w_i}$. The theoretical weights are $w_i = n_i/\sigma_i^2$, but in practice we use $w_i = 1/s_i^2$. The nominal null distribution of Q is χ_{k-1}^2

We also computed the I^2 statistics:

$$I^2 = \frac{\hat{\tau}^2}{\hat{\tau}^2 + \hat{\sigma}^2} \tag{11}$$

We used the between-study variance estimated by FSL's FLAME [27] and computed $\hat{\sigma}^2$ was computed as the average within-study contrast variance across all studies. Using this metric, voxels with values close to 0 present negligible between-study variance and values close to 1 outline appreciable study heterogeneity and the importance of random-effects models.

3. Results

3.1. Robustness to violation of model assumptions

Fig. 2A presents the one-sample simulation results in small samples, i.e. under small number of studies or small number of subjects. We focus here on methods fow which valitidy is only guaranteed in large samples: FFX GLM and MFX GLM, under ideal conditions otherwise (i.e. τ^2 =0 for FFX GLM and τ^2 =1 for MFX GLM). When the number of subjects is small, FFX GLM is invalid for all within-study variances investigated, regardless of the number of studies included in the meta-analysis. On the other hand MFX GLM is conservative for small number of studies and constant within-study variance. More surprinsingly, while MFX GLM is valid for constant within-study variances it is invalid in the presence of large variations in the within-study variances, regardless of the number of subjects included in each study.

Fig. 2B presents the one-sample simulation results under heteroscedasticity (σ_i^2 varying across studies). We focus here on methods fow which valitidy is only guaranteed under homoscedasticity: RFX and Contrast permutation, in a sample of 25 studies with 20 subjects each under ideal conditions otherwise (i.e. $\tau^2=1$). RFX GLM and contrast permutation are robust to heteroscedasticity for all settings studied. RFX GLM is closer to nominal. For small P-values, Contrast Permutation is conservative as expected due to the discrete nature of its distribution.

Fig. 2C presents the one-sample simulation results under heterogeneity ($\tau^2 > 0$). We focus here on methods fow which valitidy is only guaranteed under homogeneity: Fisher, Stouffer, Weighted Z and FFX GLM, with a sample of k = 25 studies with $n_i = 20$ subjects each. All fixed-effects methods are invalid under heterogeneity.

Similar behaviours are observed for two-sample tests (cf. Supplementary Fig. S1 and Fig. S2).

3.2. Robustness to units mismatch

Fig. 3 presents the simulation results under unit mismatches for one-sample tests. When different scaling algorithm are used (Fig. 3, 2nd and 3rd columns), e.g. with different neuroimaging software packages (provided that differences in scaling targets have been accounted for), Contrast Permutation has a behaviour that is close to nominal. RFX GLM is valid but conservative. MFX GLM is robust to the presence of unit mimatches when the studies are homoscedastic. In the presence of strong heteroscedasticity, MFX GLM remains invalid as when the units are matched due to finite sample inaccuracies (cf. previous section). In the presence of slight heteroscedasticity, unit mismatches can cause invalidity of othewise valid MFX GLM. When the contrast are scaled differently (Fig. 3, 4th column), we observe a very similar pattern than for different scaling algorithm. Similar behaviours are observed for two-sample tests (cf. Supplementary Fig. S3 and Fig. S4).

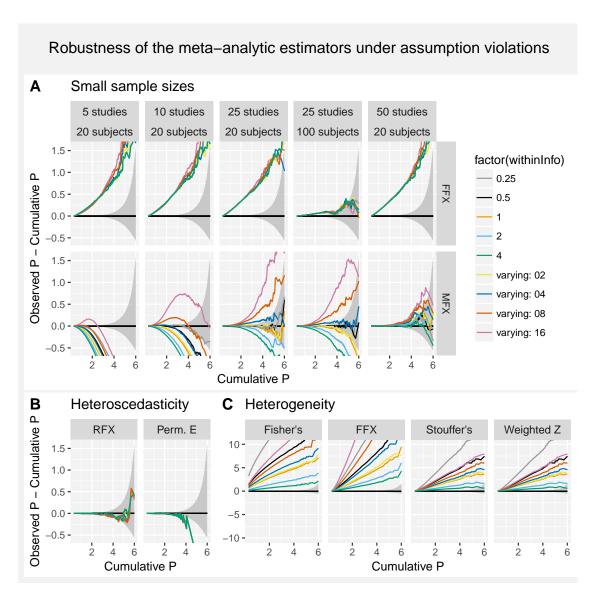


Figure 2: Deviation from theoretical P-values in one-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). P-values are displayed using a negative log₁₀ scale.

3.3. Accuracy on real data

Fig. 4A. displays the ROC curves for all the meta-analytic estimators under varying levels of heterogeneity. As expected, the fixed-effects approaches are the most affected by heterogeneity. Hence, Fisher's method is both the most sensitive method under low heterogeneity and the less sensitive under large heterogeneity. Random-effects approaches are relatively insensitive to the level of heterogeneity. Amongst random-effects approaches the most optimal are Stouffers RFX and Z Permutation that display nearly identical ROC curves, followed by MFX GLM, Contrast Permutation and finally RFX GLM. Differences between MFX GLM and RFX GLM are likely to be non-significant as they both had p-values within the 95% confidence interval.

Fig. 4B. displays the ROC curves for all the meta-analytic estimators under varying levels of heteroscedasticity. Again, the fixed-effects approaches are the most sensitive to heteroscedasticity. This can be explained by the fact that under high heteroscedasticy, some studies will present a low (or high) within-study variance, relatively increasing the between-study variance by comparison to the within-study variance. Amongst the random-effects approaches the most optimal are again Stouffer MFX and Z permutation that display nearly identical ROC curves.

3.3.1. Presence of heterogeneity and heteroscedasticity in real data

Fig. 5 presents the results of Cochran's Q test of heterogeneity on real data. Overall, 88% of the voxels displayed significant heterogeneity at FDR-corrected P < 0.05. Fig. 6 displays the I^2 statistic. 27% of the voxels presented

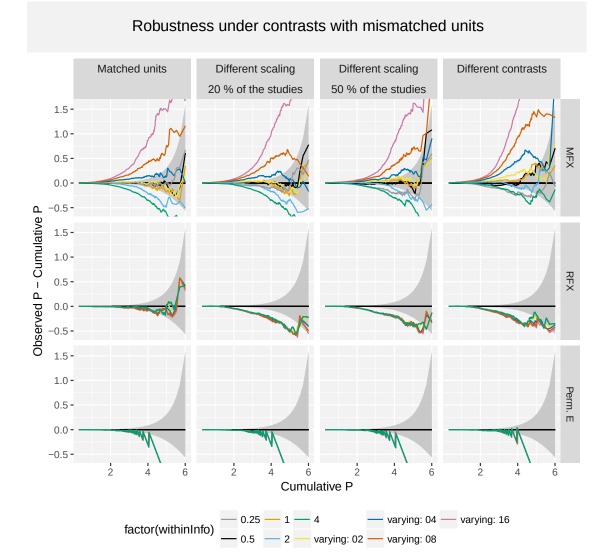


Figure 3: Deviation from theoretical P-values in one-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). Deviation from theoretical Z in two-sample tests with unit mismatch, under ideal circumstances for each statistical approach ($\tau^2 = 1$ and k = 5, 25, 50 with matched ("nominal") or mismatched ("different scaling target", "different scaling algorithm", "different contrast vector scaling") units.

a larger between-study variance. Both statistics therefore support the presence of study heterogeneity (non negligible between-study variance) in this collection of studies.

Fig. 7 displays the variability of the study parameter estimate across voxels. This plots supports the presence of heteroscedasticity in this dataset.

4. Discussions

With the growing availability of summary image data for published neuroimaging studies, image-based meta-analysis becomes feasible. Here, we investigated the validity and accuracy of nine meta-analytic estimators under conditions that are typically observed in fMRI and which might invalidate the underlying assumptions of each method.

As expected, fixed-effects methods were shown to be invalid in the presence of heterogeneity. On the other hand, homoscedastic methods were shown to be robust to heteroscedasticity which is in line with published literature on group fMRI statistics [17]. More surprisingly, MFX GLM was showed to be invalid in the presence of large heteroscedasticity due to its approximations not being accurate in small samples.

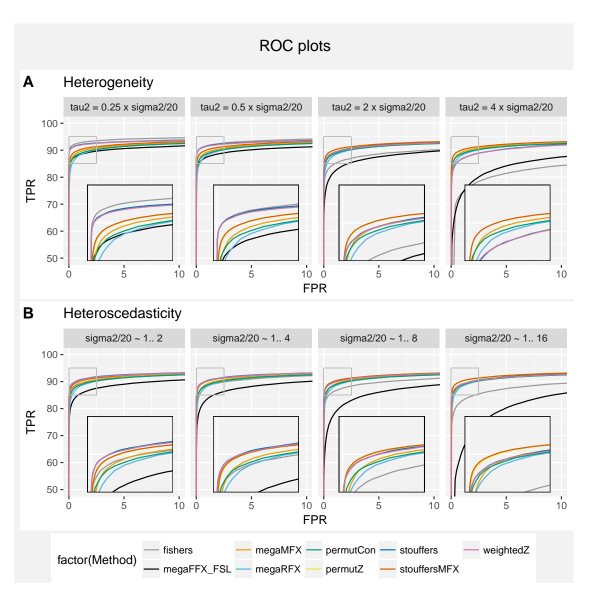


Figure 4: ROC curves of the meta-analytic estimators where true positive rate were computed using a real data meta-analysis of 21 studies of pain and false positive rates using simulated data under various levels of heterogeneity (A) and hetoroscedasticity (B).

In the presence of mismatched units, GLM RFX appeared conservative, while contrast permutation provided the best behaviour, closest to nominal. As confirmed by our real data analysis, we do expect heterogeneity to be present in meta-analytic studies due for instance to variations in the analytic procedure including varying experimental designs, analysis workflows or even due to different imaging instruments. The dataset we used in our real data analysis was created within a single lab and using the same neuroimaging analysis software (FSL), our estimates of heteroscedasticity and heterogeneity are likely to be lower than would be typically observed in a dataset pulling more heterogeneous studies. Similarly, we do expect heteroscedasticity to be present for the same reasons as well as due to varying sample sizes across studies. Finally meta-analysis sample sizes are still relatively small.

Given the still relatively small sample sizes that can be achieved in IBMA as of today, we recommend using RFX GLM, Contrast Permutation, Stouffers MFX or Z Permutation that do not rely on small sample approximations and are robust to both heterogeneity and heteroscedasticity. Unit mismatch across contrasts is likely to be an issue as, even when statistic maps are shared in support of a publication, it is very rare for it to be accompagnied by metadata describing full details on the analysis. Although z-based meta-analyses are suboptimal [5] until full reporting is routinely done, we suggest to use Z-based methods that are insensitive to units.

Because true areas of activations in real data are unknown, we relied on an external source to compute the ground truth activations: the Neurosynth platform which provides very large-scale automatic coordinate-based meta-analysis of the literature. Because this ground truth was determined using a coordinate-based (and not image-based) meta-analysis,

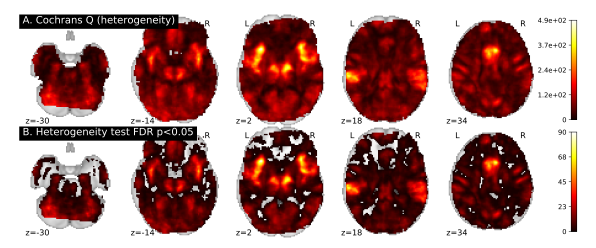


Figure 5: Test of heterogeneity using Cochran's Q test: Q statistic (A) and significant P-values in $-\log_{10}$ at FDR-corrected P < 0.05.

it is very likely to be missing some true activated areas with small effect sizes (that would effectively not pass the threshold at the level of a single study). Our estimated true positive rate are therefore likely to be overestimations.

Because true areas of no activations in real data are unknown, we relied on simulated data to compute the false positive rates. We only had a single real dataset for meta-analysis and therefore only computed true positive rates once and combined them with the different false positive rates computed by simulations. In practice the sensibility is very likely to vary under different level of heterogeneity, heteroscedasticity and one would need a larger real dataset to fully investigate sensitivity. The most faithful are the ones that correspond to the settings that are closer to the one observed in the real dataset, i.e. is sigma2/20 1...16.

Finally, in our simulations we investigated heteroscedasticity due to varying within-study variances but not due to varying sample sizes. We do expect the results to be consistent (ignoring the X^2 dof) but would have to be double checked.

5. Conclusion

We have compared nine meta-analytic approaches in the context of one-sample test. Through simulations, we found the expected invalidity of fixed-effects approaches in the presence of study heterogeneity, but also of FFX GLM even with no between-study variation. In a real dataset of 21 studies of pain, there was evidence for substantial between-study variation that supports the use of random-effects meta-analytic statistics. When only contrast estimates are available, RFX GLM was valid. When only standardised estimates are available, permutation is the preferred option as the one providing the most faithful results.

6. Acknowledgements

We gratefully acknowledge the use of the pain dataset from the Tracey pain group, FMRIB, Oxford. The majority of this work was conducted while TEN and CM were at the University of Warwick and used the High Performance Computing cluster of the Department of Statistics, University of Warwick.

7. References

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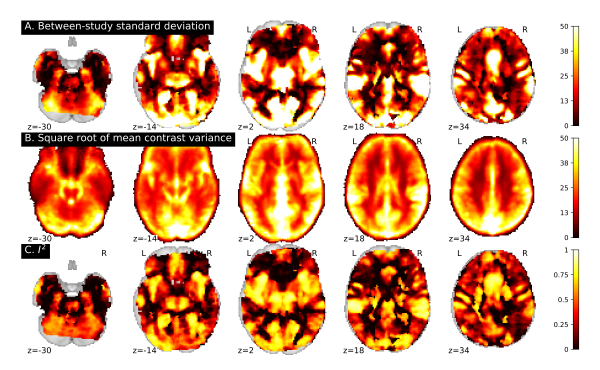


Figure 6: Estimated between-study variance (A), estimated average within-study contrast variance (B) and ratio of the estimated between study-variance onto total variance (i.e. I^2) (C).

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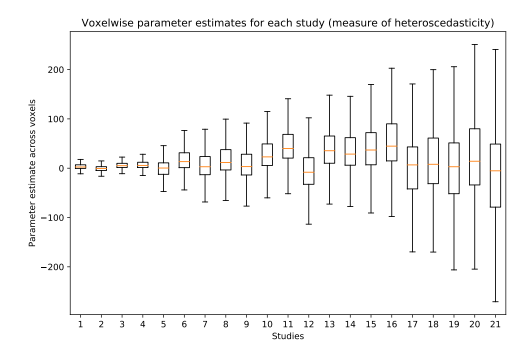


Figure 7: Boxplot of the parameter estimates across voxels by studies. For each box, the orange line corresponds to the median and the top and bottom lines of the black square are the upper and lower quartiles of the distribution. The whiskers cover the data points that are located up to 1.5 times the inter-quartile distance. Points falling out of the whiskers are not displayed. Studies are sorted from smaller to larger interquartile ranges.

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Random effects
$$\left(\sum \eta_i \hat{\beta}_i\right) / \left(\sum \eta_i\right)$$
 with $\eta_i = 1/(\tau^2 + \sigma_i^2/n_i)$ $1/\sum \eta_i$
Fixed effects $\left(\sum \hat{\beta}_i \times n_i/\sigma_i^2\right) / \left(\sum n_i/\sigma_i^2\right)$ $1/(\sum n_i/\sigma_i^2)$

Table S1: One-sample weighted least squares (WLS) estimates and their sampling distributions for random-effects and fixed-effects metaanalyses. The FGLS estimates and assumed sampling distributions are obtained by substituing: $\tau^2 \leftarrow \hat{\tau}^2$ and $\sigma_i^2/n_i \leftarrow s_i^2$.

$$\frac{\hat{\gamma_1} - \hat{\gamma_2}}{\text{Random effects}} \frac{\hat{\gamma_1} - \hat{\gamma_2}}{\left(\sum_{i \in G_1} \eta_i \hat{\beta}_i\right) / \left(\sum_{i \in G_2} \eta_i \hat{\beta}_i\right) / \left(\sum_{i \in G_2} \eta_i \hat{\beta}_i\right) / \left(\sum_{i \in G_2} \eta_i\right) \text{ with } \eta_i = 1 / \left(\tau^2 + \sigma_i^2 / n_i\right) \frac{1}{\sum_{i \in G_1} \eta_i + 1 / \sum_{i \in G_2} \eta_i}{\left(\sum_{i \in G_1} \hat{\beta}_i \times n_i / \sigma_i^2\right) / \left(\sum_{i \in G_2} \hat{\beta}_i \times n_i / \sigma_i^2\right) / \left(\sum_{i \in G_2} n_i / \sigma_i^2\right) \frac{1}{\sum_{i \in G_1} \eta_i + 1 / \sum_{i \in G_2} \eta_i}{\left(\sum_{i \in G_1} \hat{\beta}_i \times n_i / \sigma_i^2\right) / \left(\sum_{i \in G_2} \hat{\beta}_i \times n_i / \sigma_i^2\right) / \left(\sum_{i \in G_2} n_i / \sigma_i^2\right) \frac{1}{\sum_{i \in G_2} \eta_i / \sigma_i^2} \frac{1}{\sum_{$$

Table S2: Two-sample weighted least squares (WLS) estimates and their sampling distributions for random-effects and fixed-effects metaanalyses. The FGLS estimates and assumed sampling distributions are obtained by substituing: $\tau^2 \leftarrow \hat{\tau}^2$ and $\sigma_i^2/n_i \leftarrow s_i^2$.

$$\begin{array}{c|c} & \hat{\gamma_1} & \operatorname{Var}(\hat{\gamma_1}) \\ \hline \text{RFX GLM} & \sum \hat{\beta_i}/k & \sigma_C^2/k \\ \text{Z MFX} & \sum Z_i/k & \sigma_C^2/k \end{array}$$

Table S3: For the meta-analytic approaches based on Ordinary Least Squares (OLS), one-sample meta-analytic estimates and sampling variances. Note that σ_C^2 will be different for each row of this table.

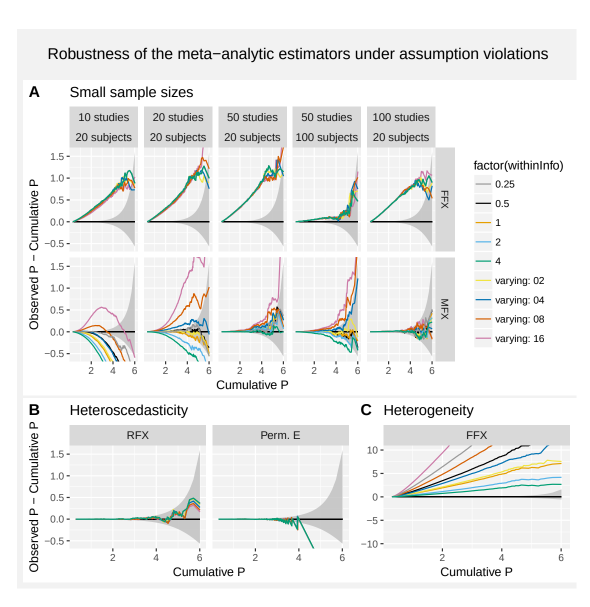


Figure S1: Deviation from theoretical P-values in two-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). P-values are displayed using a negative \log_{10} scale.

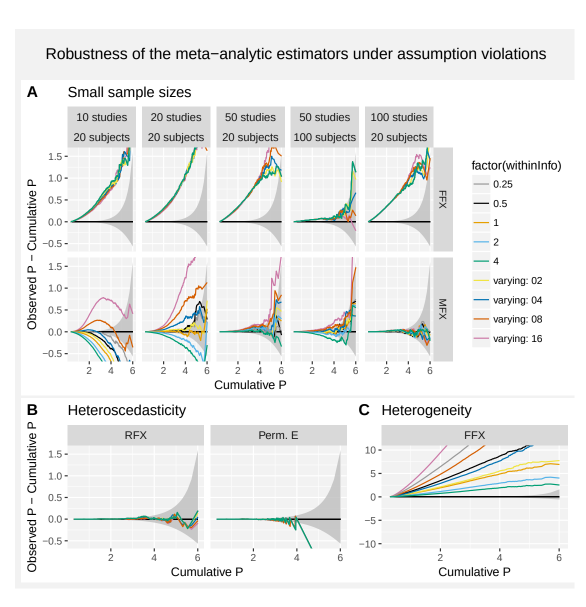


Figure S2: Deviation from theoretical P-values in unbalanced two-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). P-values are displayed using a negative \log_{10} scale.

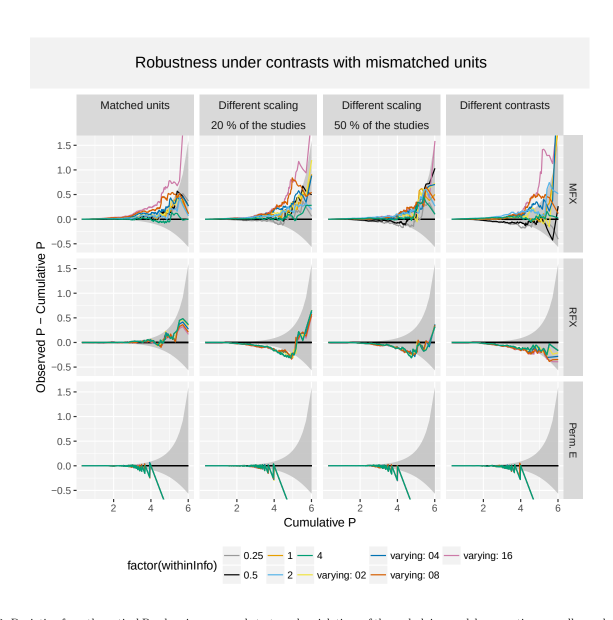


Figure S3: Deviation from theoretical P-values in one-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). Deviation from theoretical Z in two-sample tests with unit mismatch, under ideal circumstances for each statistical approach ($\tau^2 = 1$ and k = 5, 25, 50 with matched ("nominal") or mismatched ("different scaling target", "different scaling algorithm", "different contrast vector scaling") units.

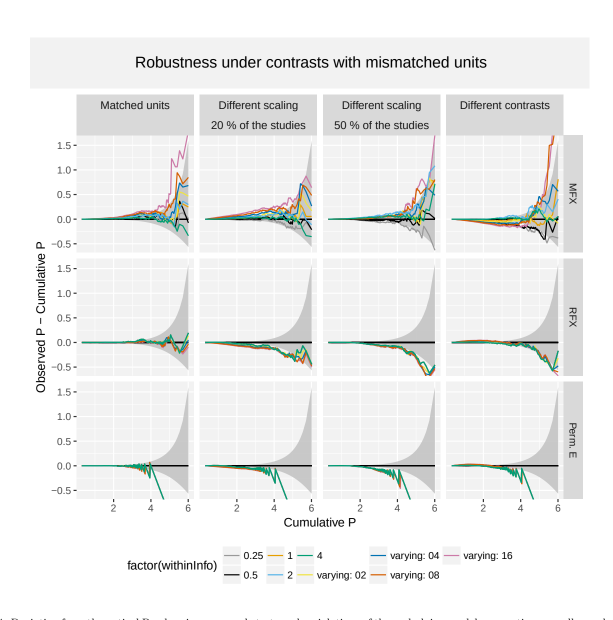


Figure S4: Deviation from theoretical P-values in one-sample tests under violations of the underlying model assumptions: small sample sizes (A), heteroscedasticity (B) and heterogeneity (C). Deviation from theoretical Z in two-sample tests with unit mismatch, under ideal circumstances for each statistical approach ($\tau^2 = 1$ and k = 5, 25, 50 with matched ("nominal") or mismatched ("different scaling target", "different scaling algorithm", "different contrast vector scaling") units.