

A direct approach to estimating false discovery rates conditional on covariates

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<https://github.com/SiminaB/Presentations>

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Goal

Extend **false discovery rate framework** to easily incorporate **external covariates (meta-data)** in decision of whether to reject a hypothesis when testing many hypotheses at once.

Background

Multiple testing is a ubiquitous issue in modern science:

Need to test relationship between hundreds or thousands of variables/features and one outcome (genomics, metabolomics etc.)

	Fail to reject null	Reject null	Total
Null true	U	V	m_0
Null false	T	S	$m - m_0$
	$m - R$	R	m

R = number of discoveries

V = number of false discoveries

Benjamini and Hochberg, 1995, *JRSSB*.

Background

False discovery rate (FDR) often used as framework to control for multiple testing.

Natural definition:

$$FDR = E \left[\frac{V}{R} \right].$$

Since R can be equal to 0, usually defined as:

$$FDR = E \left[\frac{V}{R} \middle| R > 0 \right] Pr(R > 0).$$

Benjamini and Hochberg, 1995, *JRSSB*.

Background

FDR control and estimation approaches rely on an estimate of the proportion of null hypotheses, π_0 :

$$\pi_0 = P(\text{hypothesis } i \text{ is null}).$$

- Original Benjamini-Hochberg (BH) approach assumes $\pi_0 \equiv 1$
- Storey, 2002, *JRSSB* estimates π_0 as a fixed value and multiplies the BH FDR estimates by $\hat{\pi}_0$.
- We expand this framework to estimate π_0 as a function of external covariates.

Motivating case study

- Genome-wide association (**GWAS**) study looking at associations between millions of genetic loci and BMI (Locke et al, 2015, *Nature*).
- Loci are single nucleotide polymorphisms (SNPs) that usually have 2 possible variants (alleles).
 - ▶ Major allele - more common allele, minor allele - less common allele.
- Each SNP has a different population-level frequency (coded as **MAF = minor allele frequency**).
- Not all SNPs have the same sample size (**N**), since they may be genotyped in different individuals.

Plan to use MAF and N as external covariates!

Building on prior work

Our work builds on the work of Benjamini and Hochberg, Efron, Storey, Scott et al, 2015, *JASA*, who framed the concept of **FDR regression**, extending FDR and π_0 to include covariates.

We focus on estimating π_0 as a function of covariates, then using it as a plug-in estimator to estimate FDR as a function of covariates, à la Storey.

We also use ideas from Ignatiadis et al, 2016, *Nat. Methods* that adjusting for covariates independent of the data - conditional on the null being true - can improve power.

Approach: Extend definitions of π_0 and FDR to include covariates

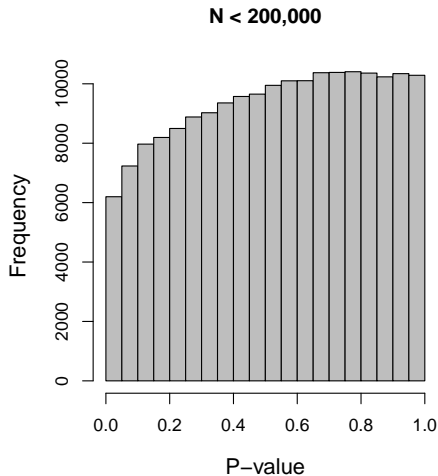
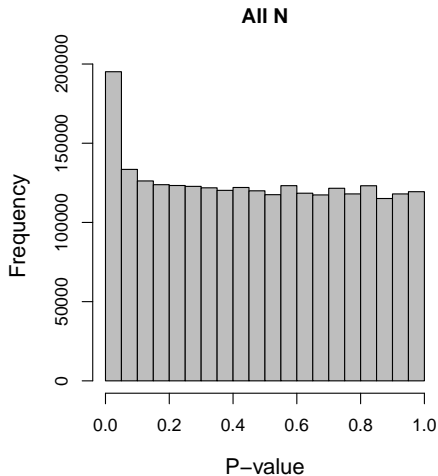
Assume a set of covariates in a column vector \mathbf{X}_i of length c , possibly with $c = 1$ and extend definitions:

$$\begin{aligned}\pi_0(\mathbf{x}_i) &= Pr(\theta_i = 1 | \mathbf{X}_i = \mathbf{x}_i), \\ FDR(\mathbf{x}_i) &= E \left[\frac{V}{R} \middle| R > 0, \mathbf{X}_i = \mathbf{x}_i \right] Pr(R > 0 | \mathbf{X}_i = \mathbf{x}_i).\end{aligned}$$

Motivating case study: GWAS meta-analysis for BMI

- Looked at ~ 2.5 million SNPs in $\sim 340,000$ individuals, considering their association with BMI.
- $\sim 320,000$ were of European descent
 - ▶ Used HapMap CEU population as reference for MAF
- Sample size (N) varies between SNPs (50,002 - 339,224, median = 235,717)

Dependence of p-values on sample sizes



Approach: Estimate $\pi_0(\mathbf{x}_i)$ using logistic regression

Assume that null p-values come from $\text{Uniform}(0,1)$ and the alternative p-values from a distribution with cdf G , so that for a large enough $\lambda \in (0, 1)$, $G(\lambda) \approx 1$.

Define:

$Y_i = 1(\text{P-value from test } i > \lambda) = P(\text{test } i \text{ not rejected at level } \lambda)$.

For a fixed threshold λ :

$$\pi_0(\mathbf{x}_i) \approx \frac{E[Y_i | \mathbf{X}_i = \mathbf{x}_i]}{1 - \lambda}.$$

Use a regression framework to estimate $E[Y_i | \mathbf{X}_i = \mathbf{x}_i]$, then estimate $\pi_0(\mathbf{x}_i)$ by:

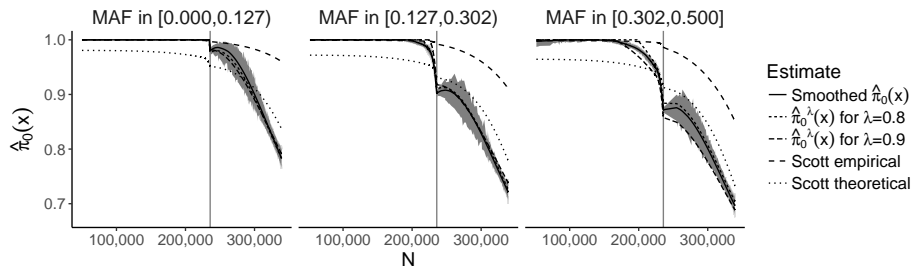
$$\hat{\pi}_0(\mathbf{x}_i) = \frac{\hat{E}[Y_i | \mathbf{X}_i = \mathbf{x}_i]}{1 - \lambda}.$$

Approach: Estimate $\pi_0(\mathbf{x}_i)$ using logistic regression

Additional details:

- May need to threshold at 1, given division by $1 - \lambda$.
- Can fix λ at e.g. 0.8 or 0.9 *or* smooth over a series of thresholds $\lambda \in (0, 1)$.
- Can use a bootstrap approach for confidence intervals.
- For BMI GWAS, split MAFs into 3 categories and used cubic splines with 5 degrees of freedom for N.

Estimates of $\pi_0(\mathbf{x}_i)$ for BMI GWAS



Grey = 90% bootstrap CI

Vertical line = median sample size

Approach: Use plug-in estimate of $\pi_0(\mathbf{x}_i)$ for $\text{FDR}(\mathbf{x}_i)$

- Multiply estimate of FDR obtained via BH approach by our estimate of $\pi_0(\mathbf{x}_i)$
- Note that is the same as approach of Storey, but considers extension to covariates
- Major assumption: Conditional on the null being true, the p-values do not depend on the covariates
i.e. the probability of a feature being from one of the two distributions depends on the covariates but the actual test statistic and p-value does not depend on the covariates further under the null.

Results for estimated $FDR(\mathbf{x}_i)$ for BMI GWAS

Number of SNPs with an estimated $FDR \leq 5\%$ for various approaches.

	BL	Scott T	Scott E	Storey	BH
Number with $\widehat{FDR} \leq 5\%$	13,384	16,697	7,636	12,771	12,500

BL = our approach

Scott T = Scott et al approach with theoretical null

Scott E = Scott et al approach with empirical null

All the discoveries from the BH approach are also present in BL.

Overlap with Storey = 12,740.

Simulations to check FDR control and power

Extensive simulations are presented in our paper/Github page.

Power = TPR (true positive rate) = fraction of truly alternative discoveries out of the total number of truly alternative features.

The good:

- If there is no or low correlation between test statistics, generally shows good control of FDR
- Always leads to an improvement in power over Benjamini-Hochberg, which increases with lower $\pi_0(\mathbf{x}_i)$ (max 6%-11% in absolute terms)
- Improved interpretability compared to Storey's approach
- FDR control is much better compared to Scott FDR regression approach when test statistics are not from a normal distribution

Simulations to check FDR control and power

Extensive simulations are presented in our paper/Github page.

Power = TPR (true positive rate) = fraction of truly alternative discoveries out of the total number of truly alternative features.

The caveats:

- If test statistics are highly correlated, does not appropriately control the FDR
- If $\pi_0(\mathbf{x}_i)$ is high, not much gain in power over Benjamini-Hochberg
- Gain in power over Storey's approach is usually minimal (0-2%)
- Power is better for Scott FDR regression approach when test statistics are from a normal distribution

Conclusions

We developed a direct approach to estimating FDR conditional on covariates.

Why should you use it?

- Improved power compared to BH
- Improved interpretability compared to Storey
- Improved robustness compared to Scott

We hope to extend/apply this approach to a number of other scenarios.

Questions?

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Preprint:

<http://www.biorxiv.org/content/early/2017/07/25/035675>

Code for all analyses/simulations in paper:

<https://github.com/SiminaB/Fdr-regression>

Package which includes FDR regression approach: [https:](https://bioconductor.org/packages/release/bioc/html/swfdr.html)

[//bioconductor.org/packages/release/bioc/html/swfdr.html](https://bioconductor.org/packages/release/bioc/html/swfdr.html)
(uses linear, not logistic regression)