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	Τ	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 :

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

- ullet 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

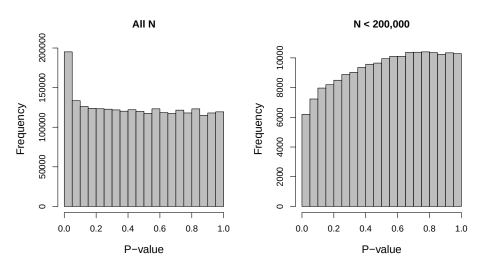
Assume a set of covariates in a column vector \mathbf{X}_i of length c, possibly with c = 1 and extend definitions:

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

Motivating case study: GWAS meta-analysis for BMI

- Looked at \sim 2.5 million SNPs in \sim 340,000 individuals, considering their association with BMI.
- ~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 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:

 $Y_i = 1(P$ -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:

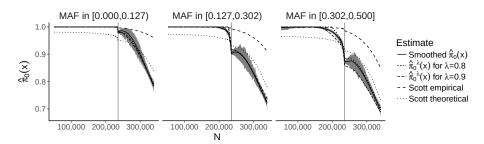
$$\hat{\pi}_0(\mathbf{x}_i) = \frac{\hat{E}[Y_i|\mathbf{X}_i = \mathbf{x}_i]}{1-\lambda}.$$

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

Additional details:

- May need to threshold at 1, given division by 1λ .
- 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 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	ВН
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://bioconductor.org/packages/release/bioc/html/swfdr.html (uses linear, not logistic regression)