# A practical evaluation of recent methods in high-dimensional inference

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May 7, 2015

#### Problem and motivation

- $x \in \mathbb{R}^p, y \in \mathbb{R}$  have a joint distribution P where  $y|x \sim N(x^T\beta, \sigma^2)$
- Observe  $X = (x_1, ..., x_n)^T$ ,  $Y = (y_1, ..., y_n)$  iid
- Problem: test  $H_i$ :  $\beta_0 = i$  for i = 1, ..., p
- Motivation: x are SNPs (mutations), y is phenotype

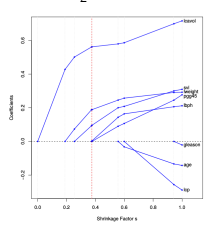
## Methods

	Control	<i>p</i> > <i>n</i>
Classical inference (Pearson 1930)	Marginal	No
Debiased lasso (Javanmard et al. 2014)	Marginal	Yes
Knockoffs (Barber et al. 2014)	FDR	?
Covariance test (Lockhart et al. 2014)	??	Yes
+ FDR control (G'Sell et al. 2013)	FDR	Yes

# The LASSO path

All three methods share an association with LASSO:

$$\hat{\beta}_{\lambda} = \operatorname{argmin}_{\beta} \frac{1}{2} ||X\beta - Y||^2 + \lambda ||\beta||_1$$

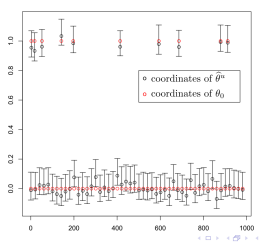


(Image credit: ??)



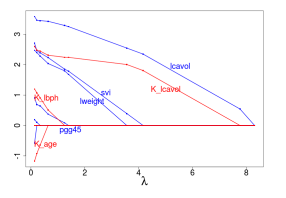
## Debiased regularized M-estimators

- (2014) Javanmard and Montanari
- Standard assumptions + sparsity condition on  $\beta$  + large n and p asymptotics



## Knockoff filter

- (2014) Barber and Candés
- Finite sample  $Y \sim N(X\beta, \sigma^2 I)$ ,  $n \leq p$ , control FDR
- Extension to p > n, FWER control, etc. forthcoming...



lweight 22.5652 lcavol 20.5199 svi 4.4871 lbph 1.1865 age 0.0829 gleason 0.0387 lcp -0.2359 pgg45 -3.3742

#### Covariance test

- (2014) Lockhart, Taylor, Tibshirani (x 2)
- Standard assumptions  $Y \sim N(X\beta, \sigma^2 I) + \text{large } p$  asymptotics
- See also non-asymptotic exact test (Lee, Sun x 2, Taylor 2015)
- What kind of Type I error does it control?

Step	Predictor entered	Forward stepwise	Lasso
1	lcavol	0.000	0.000
2	lweight	0.000	0.052
3	svi	0.041	0.174
4	lbph	0.045	0.929
5	pgg45	0.226	0.353
6	age	0.191	0.650
7	lcp	0.065	0.051
8	gleason	0.883	0.978

#### FDR control for covariance test

- G'Sell, Wager, Chouldechova, Tibshirani (2013)
- Two methods to control FDR for convariance test... but under a different definition of Type I error
- Type I error is defined according to the incremental null (next slide)

• For a subset E of the variables, define  $\beta^E = (X_E^T X_E)^{-1} X_E^T X \beta$ 

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- Selective inference. Condition on a randomly selected subset E, test hypotheses  $H_i: \beta_i^E = 0$  for all  $i \in E$

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• Incremental null. (Informal explanation). I consider adding variables one at a time; I make a mistake whenever I add a variable to the model which doesn't improve the fit of the model (and is redundant). If I reject multiple variables, the false rejections are the variables which were already redundant at the time I added them. However, if a variable is initially useful and only becomes redundant as more variables are added, it is not considered a mistake.

- For a subset E of the variables, define  $\beta^E = (X_E^T X_E)^{-1} X_E^T X \beta$
- The full model null. Test multiple hypotheses  $H_i$ :  $\beta_i = 0$
- In this talk, we define type I errors according to full model null...

## Methods

## But what's actually used in practice?

	Control	p > n
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+ FDR control (G'Sell et al. 2013)	FDR	Yes
Marginal screening	???	Yes

#### Statistical Validation

- These procedures are derived under strong assumptions (linearity, gasusianity, homoscedasticty)
- How well do they work in real data where these assumptions are violated?
- We could validate inference procedures in real data if only we knew the 'true'  $\beta$ , (re)defined as

$$\beta = \mathbf{E}[\mathbf{x}\mathbf{x}^T]^{-1}\mathbf{E}[\mathbf{y}\mathbf{x}]$$

### Statistical Validation

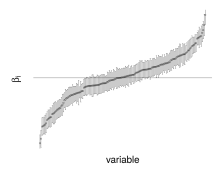
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• Possibility: take a dataset with large p and  $humongous\ n$ , so we can get an extremely precise estimate of  $\beta$  using OLS. Then test the high-dimensional inference procedures on subsamples of size  $n_0 \le p < n$  of the data

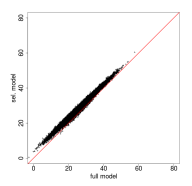
- Data with p = 163 survey questions from an online personality test, n = 49086 (after processing)
- Predict self-reported age of respondent, y, from their responses
- Is n large enough for us to confidently say which  $\beta_i = 0$  (for use as ground truth?)

Coefficient estimates  $\pm$  3 sd

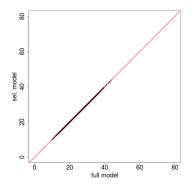


Consider declaring all variables whose intervals cross 0 to be null. Then  $p_1=105$  (out of 163)

- If n were large enough, then for the selected model S we should have  $\hat{y} = \sum_{i=1}^{p} X \hat{\beta}_i$  close to  $\hat{y}_S = \sum_{i \in S} X_i \hat{\beta}_i$
- But...



- Here n is not large enough for p = 163
- If we reduce the dimensionality to 15 by subsampling columns, it looks more convincing that we selected the correct 10 variables



#### Dillemma

- It is by no means *impossible* to get large enough data to estimate high-dimensional  $\beta$ , with say, p > 100
- But if were *easy* to get such large *n* data... we wouldn't need these new inference techniques in the first place!

# Why not use simulations?

- Simulations can be used to test robustness of the procedure
- In simulations, we can add all the nonlinearities, nongaussianity, etc.
   that we want

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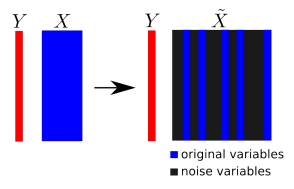
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- **Disadvantage:** Are these simulations relevant? How can we tell the simulated models are realistic?

#### Idea

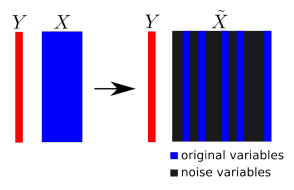
I give you real data mixed in with noise variables



• Can you identify the original columns from the noise columns?

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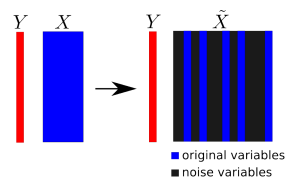
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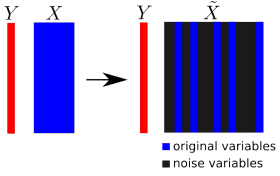
#### Idea

I give you real data *mixed in* with noise variables



- Can you identify the original columns from the noise columns?
- I can test your procedure this way, because I know the ground truth!
- Caveat: this test is unrealistically 'easy' (due to lack of correlations)

- Synthetic negative controls (SNCs) are artificial columns which are correlated to X, yet still have zero (population) regression coefficients
- Suppose I give you real data + SNCs, then you apply high-dimensional inference. If you reject any SNCs, we know these are errors!
- This gives us some measure of performance on "real" data (maybe?)



- Given random vector  $x \in \mathbb{R}^p$ , let e be noise in  $\mathbb{R}^p$  independent of x.
- Let  $\Gamma$  be a fixed  $p \times q$  matrix. Define synthetic negative controls  $z \in \mathbb{R}^q$  by by

$$z = x'\Gamma + e$$

and let  $\tilde{x} = (x, z)$ , so that

$$\tilde{x}_1 = x_1, \ldots, \tilde{x}_p = x_p$$

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Then

$$\forall i \in \{1, \dots, p\} : \beta_i = \tilde{\beta}_i$$
$$\forall i \in \{p+1, \dots, p+q\} : \tilde{\beta}_i = 0$$



- Recall that  $\hat{\beta}_i$  is the *univariate regression* coefficient of Y on  $X_{i|-i}$ , where  $X_{i|-i}$  is the *residual of*  $X_i$  after  $X_i$  is regressed on the other columns..
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ullet The orthogonal projection  $P_X^\perp$  of  $ilde X_{p+1}$  is

$$P_X^{\perp} \tilde{X} = P_X^{\perp} X \Gamma_i + P_X^{\perp} E_i = 0 + E_i$$

since  $P_X^{\perp}X = 0$ ; meanwhile since  $E_i \perp X$ ,  $P_X^{\perp}E_i = E_i$ .



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• Since  $E_i \perp y$ , we have  $Cor(P_X^{\perp} \tilde{X}_{p+i}, y) = 0$ , hence  $\tilde{\beta}_{p+i} = 0$ 



• Population version:  $\beta_i = 0$  if the projection of  $X_i$  on the null space of the other covariates is uncorrelated with Y

- Since  $E_i \perp y$ , we have  $\operatorname{Cor}(P_X^{\perp} \tilde{X}_{p+i}, y) = 0$ , hence  $\tilde{\beta}_{p+i} = 0$
- And since  $\beta_j = 0$  for all the added variables  $j = p + 1, \dots p + q$ , it follows that  $\tilde{\beta}_i$  is unchanged for  $i = 1, \dots, p$ .

## Using SNCs to evaluate procedures

- Take low-dimensional real data mixed with SNCs (synthetic negative controls), apply inference procedure
- Proxy for Type I error: Rejected SNCs
- Proxy for Power: Rejected original variables

# A step-by-step tutorial (in R)

1. Take the prostate data

2. Construct 20 synthetic negative controls

```
> GAMMA <- matrix(rnorm(8 * 20), 8, 20)
> E <- matrix(rnorm(97 * 20), 97, 20)
> sncs <- as.matrix(x) %*% GAMMA + 2 * E
> sncs <- data.frame(sncs)
> colnames(sncs)
  [1] "X1" "X2" "X3" "X4" "X5" "X6" ...
[19] "X19" "X20"
```

- 3. Create combined design matrix
- $> x2 \leftarrow cbind(x, sncs)$

4. Try marginal screening

```
5. Try covariance test
> library(covTest)
> covTest(lars(as.matrix(x2), y), as.matrix(x2), y)
$results
Predictor_Number Drop_in_covariance P-value
                               69.0292
                                         0.0000
                 1
                 5
                                1.5390 0.2219
                                6.8094 0.0020
                11
                                0.8559 0.4294
(Numbers 1, 5, 2 are original, 11 is a SNC)
```

```
6. Try debiased lasso (code at
http://web.stanford.edu/ montanar/sslasso/)
> res <- SSLasso(as.matrix(x2), y)
[1] "10% done"
...
[1] "90% done"
> rej <- (res$up < 0) | (res$low > 0)
> names(x2)[rej]
[1] "lcavol" "lweight" "svi"
```

### Disclaimer!

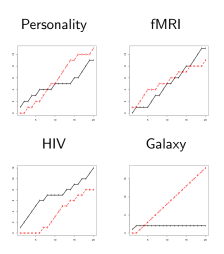
- I am not proposing SNCs as a methodology for inference
- There is a danger of inferring that Type I error has been controlled from lack of rejection of SNCs. There are no formal guarantees of this!
- One should interpret results from experiments with SNCs in the same way one interprets simulation results with purely synthetic data

# More Experiments!

Data	n	$p_1$	Linear?	Gaussian?	Constant $\sigma^2$ ?
Personality	3000	163	No	No	No
fMRI	1750	53	No	OK	No
HIV	842	207	No	Yes?	OK?
Galaxy	323	4	No	OK	No

- We add  $n/2 p_1$  synthetic negative controls
- X is scaled,  $\Gamma$  is a gaussian matrix, Var(E) is chosen to yield 'interesting' results
- Personality data is subsampled

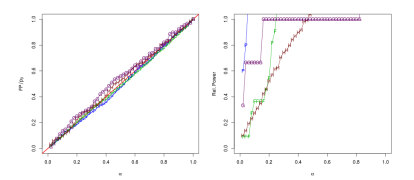
# Marginal Screening



Legend: 0 = False positives, 1 = True positives

## **Ordinary Least Squares**

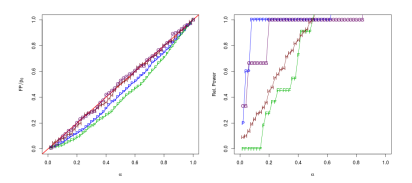
"Rel. power" = TP/(max number of TPs at  $\alpha = 0.5$  for any method)



Legend: P = Personality, F = fMRI, H = HIV, G = Galaxy

#### Debiased Lasso

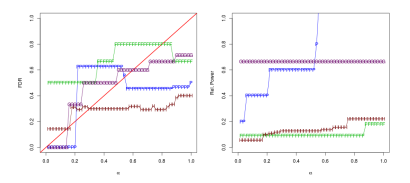
Can you spot the difference from the previous slide?



Legend: P = Personality, F = fMRI, H = HIV, G = Galaxy

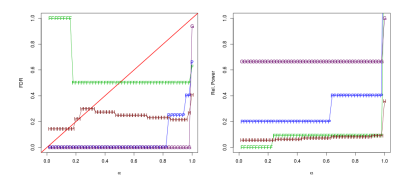
### Covariance Test

Forward Stop: reject first  $\hat{k}$ , where  $-\frac{1}{\hat{k}} \sum_{i=1}^{\hat{k}} \log(1-p_i) \leq \alpha$ 



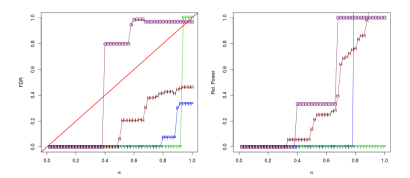
### Covariance Test

Strong Stop: reject first  $\hat{k}$ , where  $\frac{m}{\hat{k}}e^{\sum_{j=\hat{k}}^{p}\log(p_{j})/j}\leq \alpha$ 



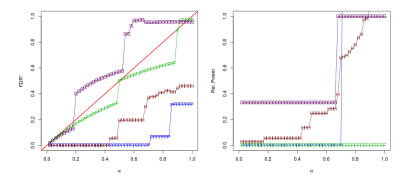
### Knockoffs

#### Using Knockoff+ threshhold



### Knockoffs

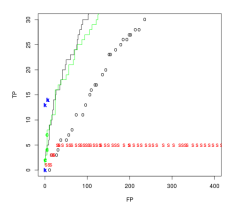
Note: 
$$FDR^* = \mathbf{E}[FP/(FP + TP + 1/\alpha)]$$



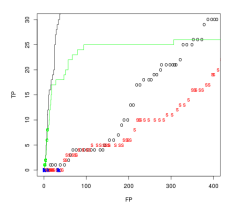
## Variable Ranking Criteria

- Forget about Type I error for a second...
- Use procedures to rank variables by p-value
- Easy to compare procedures with different Type I criteria and also non-inference variable selection
- (Optional) score by Area Under Curve (AUC), etc.

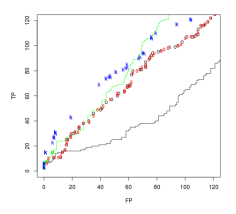
# Variable Ranking: Personality



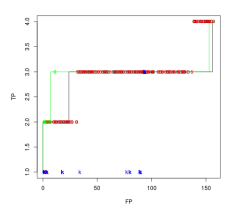
# Variable Ranking: fMRI



# Variable Ranking: HIV



# Variable Ranking: Galaxy

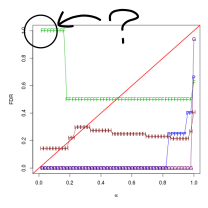


## Commentary

- We should not conclude too much from four experiments with rather arbitrary generation parameters...
- Debiased lasso similiar to OLS but more conservative, less powerful
- Knockoffs vs covariance test:
  - Knockoffs may control FDR more robustly than Covariance test (especially at small  $\alpha$ )
  - Knockoffs and covariance are similar in power overall but have different case-by-case behavior
- Knockoffs tend to be conservative, but have good variable ranking in some cases (Personality, fMRI)
- Marginal screening remains annoyingly effective...

### Another look at covTest

### Strong stop type I error

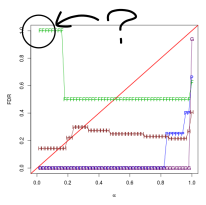


• Why did this negative control get rejected in the fMRI data at such a low  $\alpha$ ?

(thanks to Stefan!)

### Another look at covTest

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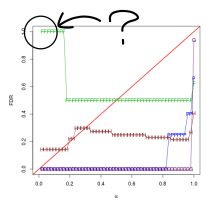


- According to the incremental null, the negative control was not a mistake... it is the best single predictor by far!
- The particular negative control took an average of the original columns

(thanks to Stefan!)

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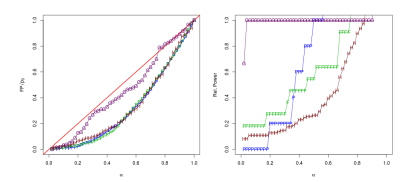
• It is true that  $\beta_i = 0$  for the rejected variable... but should we really disregard such strong "proxy" variables?

## Low sample size

Data	n	р	$p_1$	Linear?	Gaussian?	Constant $\sigma^2$ ?
Personality	100	1500	163	No	No	No
fMRI	100	875	53	No	OK	No
HIV	100	421	207	No	Yes?	OK?
Galaxy	100	161	4	No	OK	No

- Reduce the sample size to 100, so that p >> n
- Same number of negative controls, but larger added noise (easier)
- Covariance test requires estimate  $\hat{\sigma}$ : "cheat" by using OLS estimate from *original* data

### Debiased Lasso

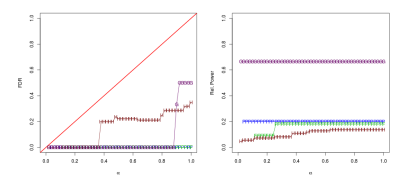


Legend: P = Personality, F = fMRI, H = HIV, G = Galaxy



### Covariance Test

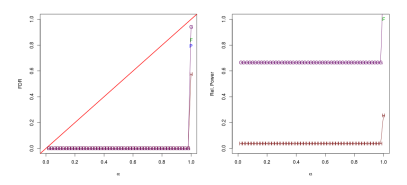
Forward Stop: reject first  $\hat{k}$ , where  $-\frac{1}{\hat{k}} \sum_{i=1}^{\hat{k}} \log(1-p_i) \leq \alpha$ 



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Strong Stop: reject first  $\hat{k}$ , where  $\frac{m}{\hat{k}}e^{\sum_{j=\hat{k}}^{p}\log(p_{j})/j} \leq \alpha$ 



## Commentary

- Debiased lasso and Covariance test + forward stop continue to control Type I error while finding true positives
- Covariance test + strong stop appears too conservative

## Are these experiments actually that realistic!?

- Here we are implicitly assuming that real data always consists of a few "active variables" and many null variables
- If that's true, it seems reasonable to model the distribution of the inactive variables conditional on knowing a superset of the active variables

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- But how do we know that  $\beta$  is really sparse?
- Even more importantly, how do we know that inferring coefficients of the *full model*  $\beta$  are a meaningful objective? E.g. should we consider the covariance test to have made a mistake in the fMRI data?

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- Even more importantly, how do we know that inferring coefficients of the *full model*  $\beta$  are a meaningful objective? E.g. should we consider the covariance test to have made a mistake in the fMRI data?
- How can we decide between the full model null, the incremental null, or an entirely different framework altogether?
- Feedback from the practitioner is the only way we can tell if we have the right formulation for any particular application

### Questions to consider

- Why is OLS more powerful than lasso in some of these experiments even when  $\beta$  is sparse? Look at covariance conditions in the theory of LASSO...
- Why do knockoffs or lasso beat marginal screening/OLS in the HIV data? Was it due to how we generated the SNCs or is due to something special about the data itself?
- Suppose we wanted to validate selective inference or the incremental null. How can we do this with synthetic negative controls (other than pure noise?)

# Closing thoughts

"Both the client and the statistician... must base their thinking on a recognition that their assumptions will always require review and reappraisal..."

John Tukey

# Acknowledgements

Thanks to Will Fithian and Stefan Wager for useful discussions.

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