An Inferential Perspective on Data Depth

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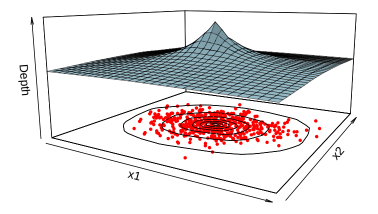
May 18, 2017

- Introduction
- Signed Peripherality Functions
- Nonconvex Penalized Multitask Regression using Depth-based Penalty
- Generalized model discovery using statistical evaluation maps
- Selecting important SNPs from multi-SNP mixed models on Twin Studies GWAS data

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What is depth?

Example: 500 points from $\mathcal{N}_2((0,0)^T, \text{diag}(2,1))$



A scalar measure of how much inside a point is with respect to a data cloud

For any multivariate distribution $F = F_{\mathbf{X}}$, the depth of a point $\mathbf{x} \in \mathbb{R}^p$, say $D(\mathbf{x}, F_{\mathbf{x}})$ is any real-valued function that provides a 'center outward ordering' of **x** with respect to F (Zuo and Serfling, 2000).

Desirable properties (Liu, 1990)

- (P1) Affine invariance: $D(A\mathbf{x} + \mathbf{b}, F_{A\mathbf{X}+\mathbf{b}}) = D(\mathbf{x}, F_{\mathbf{x}})$
- (P2) Maximality at center: $D(\theta, F_X) = \sup_{\mathbf{x} \in \mathbb{R}^p} D(\mathbf{x}, F_X)$ for F_X with center of symmetry θ , the deepest point of F_x .
- (P3) Monotonicity w.r.t. deepest point: $D(\mathbf{x}; F_{\mathbf{X}}) \leq D(\theta + a(\mathbf{x} \theta), F_{\mathbf{X}})$
- (P4) Vanishing at infinity: $D(\mathbf{x}; F_{\mathbf{X}}) \to \mathbf{0}$ as $\|\mathbf{x}\| \to \infty$.

 Halfspace depth (HD) (Tukey, 1975) is the minimum probability of all halfspaces containing a point.

$$\textit{HD}(\boldsymbol{x}, F) = \inf_{\boldsymbol{u} \in \mathbb{R}^p: \boldsymbol{u} \neq \boldsymbol{0}} P(\boldsymbol{u}^T \boldsymbol{X} \geq \boldsymbol{u}^T \boldsymbol{x})$$

Projection depth (PD) (Zuo, 2003) is based on an outlyingness function:

$$O(\mathbf{x}, F) = \sup_{\|\mathbf{u}\|=1} \frac{|\mathbf{u}^T \mathbf{x} - m(\mathbf{u}^T \mathbf{X})|}{s(\mathbf{u}^T \mathbf{X})}; \quad PD(\mathbf{x}, F) = \frac{1}{1 + O(\mathbf{x}, F)}$$

Used extensively for classification, robust estimation of outlyingness,
 L-estimation of location and scale, hypothesis testing;

Although the nonparametric concept of data depth has gained visibility and has seen many applications in recent years, its utility in achieving traditional parametric inferential goals is largely unexplored. In this proposal we develop different approaches to address this.

- Signed Peripherality Functions: robust location and scale inference;
- Nonconvex Penalized Multitask Regression using Depth-based Penalty;
- Generalized Model Discovery using Statistical Evaluation Maps: applications in Indian Monsoon, fMRI and Minnesota Twin Studies data.

- **Signed Peripherality Functions**
- Nonconvex Penalized Multitask Regression using Depth-based

Depth-based Inference May 18, 2017 Spatial signs (Locantore et al., 1999):

$$\mathbf{S}(\mathbf{x}) = \begin{cases} \mathbf{x} \|\mathbf{x}\|^{-1} & \text{if } \mathbf{x} \neq \mathbf{0} \\ \mathbf{0} & \text{if } \mathbf{x} = \mathbf{0} \end{cases}$$

Fix a depth function $D(\mathbf{x}, F) = D(\mathbf{x}, [\mathbf{X}])$. Define the signed peripherality as

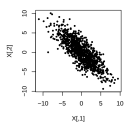
$$\kappa(D(\mathbf{x}, [\mathbf{X}]).\mathbf{S}(\mathbf{x})$$

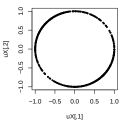
where $\kappa : [0, \infty) \mapsto [0, \infty)$ is a bounded monotone transformation.

Two cases-

- (1) κ is increasing: Robust location inference. Improves on existing estimators for robust location estimates and high-dimensional testing;
- (2) κ is decreasing: Robust scale inference.

- Say **x** follows an elliptic distribution with mean μ , covariance matrix Σ .
- Sign covariance matrix (SCM): $\Sigma_S = \mathbb{E} \mathbf{S} (\mathbf{X} \boldsymbol{\mu}) \mathbf{S} (\mathbf{X} \boldsymbol{\mu})^T$
- SCM has same eigenvectors as Σ. PCA using SCM is robust, but not efficient.





Spatial ranks

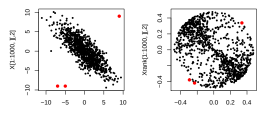
Transform the original observation

$$ilde{\mathbf{x}} = D^-(\mathbf{x}, [\mathbf{X}]) \mathbf{S}(\mathbf{x} - \mathbf{\mu})$$

where $D^-(\mathbf{x}, [\mathbf{X}])$ is the *inverse depth* of \mathbf{x} : any bounded nonnegative-valued monotonically decreasing transformation on the depth function.

This is the Spatial Rank of x.

• Depth Covariance Matrix (DCM) $\tilde{\Sigma} := \mathbb{V}(\tilde{\mathbf{X}})$. Has more information than spatial signs, so more efficient.



Results using spatial ranks

- Recovery of population eigenvectors and eigenvalues using DCM: asymptotic and robustness properties of estimates;
- Adaptations in Sufficient Dimension Reduction and functional PCA;
- Simulations and real data applications.

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Consider the multitask linear regression model:

$$\mathbf{Y} = \mathbf{XB} + \mathbf{E}$$

where $\mathbf{Y} \in \mathbb{R}^{n \times q}$ is the matrix of responses, and \mathbf{E} is $n \times q$ the noise matrix: each row of which is drawn from $\mathcal{N}_q(\mathbf{0}_q, \mathbf{\Sigma})$ for a $q \times q$ positive definite matrix $\mathbf{\Sigma}$.

We are interested in sparse estimates of the coefficient matrix ${\bf B}$ through solving penalized regression problems of the form

$$\min_{\mathbf{B}} \operatorname{Tr}\{(\mathbf{Y} - \mathbf{X}\mathbf{B})^{T}(\mathbf{Y} - \mathbf{X}\mathbf{B})\} + P_{\lambda}(\mathbf{B}). \tag{1}$$

Our estimator

We incorporate measures of data depth as a row-level penalty function. Specifically, we estimate the coefficient matrix **B** by solving the following constrained optimization problem:

$$\hat{\mathbf{B}} = \underset{\mathbf{B}}{\operatorname{argmin}} \left[\operatorname{Tr} \{ (\mathbf{Y} - \mathbf{X} \mathbf{B})^T (\mathbf{Y} - \mathbf{X} \mathbf{B}) \} + \lambda \sum_{j=1}^p D^-(\mathbf{b}_j, F) \right]$$

where $D^{-}(\mathbf{x}, F)$ is an inverse depth function.

- Regularization penalties can be interpreted as distance from the origin.
- We want to use penalties that are 'distance from a distribution centered at the origin'.
- The distribution F is fixed at the start of the modelling process, and can represent a prior belief on how the different responses are related among themselves.
- Two advantages: (1) penalty attains nonconvex shape by inverting the depth function, (2) has a natural bayesian interpretation.
- For now we assume F to be spherically symmetric plausible from a frequentist perspective.

- As F to be spherically symmetric, D^- becomes a function of the row-norm $r_j = \|\mathbf{b}_j\|_2$: $D^-(\mathbf{b}_j, F) = p_F(r_j)$.
- Use the first order Taylor approximation around a 'close enough' point r_j^* instead of $p_F(r_j)$. This is local linear approximation (Zou and Li, 2008):

$$p_F(r_j) \simeq p_F(r_j^*) + p_F'(r_j^*)(r_j - r_j^*)$$

Thus the modified solution is:

$$\hat{\mathbf{B}}^{(1)} = \underset{\mathbf{B}}{\operatorname{argmin}} \left[\operatorname{Tr}\{(\mathbf{Y} - \mathbf{X}\mathbf{B})^T (\mathbf{Y} - \mathbf{X}\mathbf{B})\} + \lambda \sum_{j=1}^p p_F'(r_j^*) r_j \right]$$

The close enough matrix \mathbf{B}^* to start from can be the least squares estimate. We call this *Local Approximation by Row Norm* (LARN).

1.9	3.3	-2.3	-1.1
0	0	0	0
2.4	4.5	1.5	-3.1
0	0	0	0

0	3.3	-2.3	0
0	0	0	0
2.4	4.5	0	-3.1
0	0	0	0

Pre-thresholding (L) vs. post-thresholding (R) estimators

- Derived theoretical properties: oracle property, near-minimax optimal performance, post-estimation thresholding to recover within row support;
- A block coordinate descent algorithm to compute solution;
- Simulation and data example.

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- In a parametric modelling setup, any candidate model is a subset of the parameter space;
- We have a collection of models, and want to classify them as 'good' or 'bad' based on if they match with a baseline model with respect to a predefined criterion;
- We shall compare sampling distributions of (potentially transformed)
 parameter estimates of a candidate model with that of a baseline model
 using a generic quantity called the e-value.

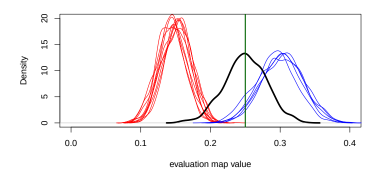
Linear regression: $Y = \mathbf{X}\beta + \epsilon$. Assume that there is a true parameter vector β_0 , with non-zero index set S_0 .

- Get $\hat{\beta}$. Obtain its bootstrap distribution: $[\hat{\beta}]$;
- ② Replace the *j*-th coefficient with 0, name it $\hat{\beta}_{-j}$. Do the same for its bootstrap distribution, say $[\hat{\beta}_{-j}]$. Repeat for all j;
- e-value of *j*-th covariate = mean depth of $\hat{\beta}_{-j}$ with respect to $[\hat{\beta}]$, i.e. $\mathbb{E}D(\hat{\beta}_{-j}, [\hat{\beta}])$;
- Select covariates with e-value less than mean depth of full model:

$$\hat{\mathcal{S}}_0 = \left\{ j : \mathbb{E}D(\hat{\boldsymbol{\beta}}_{-j}, [\hat{\boldsymbol{\beta}}]) < \mathbb{E}D(\hat{\boldsymbol{\beta}}, [\hat{\boldsymbol{\beta}}]) \right\}$$

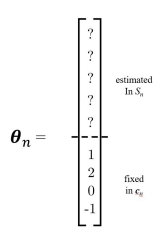
Then $\mathbb{P}(\hat{\mathcal{S}}_0 = \mathcal{S}_0) \to 1$ as $n \to \infty$.

```
DroppedVar
                   Cn
       - x20.2356008
       - \times 3.0.2428004
       - \times 40.2448785
       -x10.2473548
       - x50.2486610
     - \times 20.0.2503475
    <none> 0.2505000
       -x90.2522873
      - x21 0.2538186
      - x22 0.2547132
11
      -x140.2548410
12
      -x170.2554293
13
      - x13 0.2559990
14
      - \times 10.0.2564211
1.5
      - x24 0.2566334
16
      - x19 0.2568725
17
      - \times 250.2573902
       - x8 0.2578656
18
19
      - x16 0.2588032
      -x120.2590218
21
       - \times 6.0.2595048
22
      - \times 23.0.2598039
2.3
      - \times 150.2605307
24
      - x11 0.2606763
25
      - x18 0.2610460
2.6
       -x70.2613168
```



- Start with full model;
- ② Drop an essential predictor ⇒ Model becomes wrong ⇒ e-value shifts to left;
- Orop a non-essential predictor ⇒ Model still correct but nested within full model ⇒ e-value shifts to right.

The framework



Given traingular sequences of observable data and unknown parameters:

$$\{(\mathcal{B}_n, \boldsymbol{\theta}_n), n \in \mathbb{N}\}$$

$$\mathcal{B}_n = \{B_{n1}, \dots, B_{nk_n}\}, \quad \boldsymbol{\theta}_n \in \boldsymbol{\Theta}_n \subseteq \mathbb{R}^{p_n}$$

a candidate model \mathcal{M}_n is specified by:

- (a) The set of indices $S_n \subseteq \{1, 2, ..., p_n\}$ where parameter values are estimated from the data;
- **(b)** An ordered vector of known constants $\mathbf{c}_n = (c_{nj} : j \neq S_n)$ for parameters not indexed by S_n .

Denote the model space of \mathcal{M}_n by Θ_{mn} .

Among all such possible models, we designate one of them as the *preferred* model: say $\mathcal{M}_{*n} = (\mathcal{S}_{*n}, \mathbf{c}_{*n})$.

This is the baseline model we shall compare all models with.

Example

For variable selection, the model with all covariates is the preferred model.

For hypothesis testing, the null model is preferred model.

Designate an element of the preferred model space Θ_{*n} as the *preferred* parameter vector θ_{0n} . This is generally informative of the data generating process.

For any candidate model \mathcal{M}_n , we consider estimating functionals $\Psi_{sni}(.)$ that admit an unique minimizer $\theta_{mn} \in \Theta_{mn}$:

$$\Psi_{sn}(\theta) = \mathbb{E}\sum_{i=1}^{k_n} \Psi_{sni}(\theta, B_{ni}); \quad \theta_{mn} = \operatorname*{argmin}_{\theta \in \Theta_{mn}} \Psi_{sn}(\theta)$$

The estimate corresponding to model \mathcal{M}_n is obtained by minimizing its sample version:

$$\hat{\boldsymbol{\theta}}_{mn} = \operatorname*{argmin}_{\boldsymbol{\theta} \in \boldsymbol{\Theta}_{mn}} \sum_{i=1}^{k_n} \Psi_{sni}(\boldsymbol{\theta}, B_{ni})$$

We assume there exist $a_{sn} \uparrow \infty$ such that $[a_{sn}(\hat{\theta}_{sn} - \theta_{sn})]$ 'converges' to a distribution as $n \to \infty$. Here θ_{sn} is θ_{mn} is \mathcal{S}_n indices, and same for $\hat{\theta}_{sn}$.

Now we map parameters from the parameter space to a common multivariate space using a known smooth function:

$$\mathbf{G}_{mn}: \mathbf{\Theta}_n \mapsto \mathbb{R}^{d_n}; \quad d_n = o(\min_{\mathcal{S}} \{a_{\mathcal{S}n}, a_{*n}\})$$

This is for easy comparison of different kinds of modelling methods.

Example

Want to compare the following models built on data

$$\{(y_i, X_{1i}, X_{2i}) : i = 1, \ldots, n\}$$
:

- (1) Linear regression $Y_i = X_{1i}\beta_1 + X_{2i}\beta_2 + \epsilon_i, \epsilon_i \sim N(0, \sigma^2), \sigma > 0$;
- (2) Semiparametric regression $Y_i = X_{1i}\beta_1 + g(X_{2i}) + \epsilon_i$ for unknown g;
- (3) Single index model $Y_i = h(X_{1i}\beta_1 + X_{2i}\beta_2) + \epsilon_i$ for unknown h;

Comparing all model in terms of prediction errors has a clearer interpretation.

For $\mathbf{g} \in \mathbb{R}^{d_n}$ and $\mathcal{G}'_n \subseteq \mathbb{R}^{d_n}$ define the following:

$$d(\mathbf{g}, \mathcal{G}'_n) := \inf_{\mathbf{g}' \in \mathcal{G}'_n} \|\mathbf{g} - \mathbf{g}'\|$$

Then

(a) For two sequences of models, say $\{\mathcal{M}_{1n}\}$ and $\{\mathcal{M}_{2n}\}$, we say $\{\mathcal{M}_{1n}\}$ is nested within $\{\mathcal{M}_{2n}\}$ if, for all sequences $\{\mathbf{g}_{1n}:\mathbf{g}_{1n}\in\mathcal{G}_{1n}\}$ we have

$$\lim_{n\to\infty} d(\boldsymbol{g}_{1n},\mathcal{G}_{2n}) = 0$$

with $\mathcal{G}_{1n} := \{\mathbf{G}_{m1n}(\boldsymbol{\theta}(\mathcal{M}_{1n})) : \boldsymbol{\theta}(\mathcal{M}_{1n}) \in \boldsymbol{\Theta}_{m1n}\}$ etc.

- **(b)** A sequence of models $\{\mathcal{M}_n\}$ is called *adequate* if the model \mathcal{M}_{0n} corresponding to the singleton set $\Theta_{0n} = \{\theta_{0n}\}$, is nested within \mathcal{M}_n .
- (c) A model that is not adequate is an inadequate model.

Covers obvious cases:

$$(*,*,*,0)$$
 – adequate model

$$(*,*,*,*)$$
 – full model

$$(*,*,0,*)$$
 - inadequate model

Covers limiting cases:, e.g. $(*,*,*,\delta_n)$, $\delta_n = o(1)$ will be an adequate model in our framework.

Such data generating models, e.g.

$$Y_{ni} = X_{1i}\beta_{01} + X_{2i}\delta_n + \epsilon; \quad \beta_{01} \in \mathbb{R}, \delta_n = o(1)$$

for linear regression, frequently arise from prior choices in bayesian variable selection techniques.

We now introduce an *evaluation function*:

$$E_n: \mathbb{R}^{d_n} imes ilde{\mathbb{R}}^{d_n} \mapsto [0, \infty)$$

to quantify the relative position of $\hat{\mathbf{G}}_{mn} \equiv \mathbf{G}_{mn}(\hat{\boldsymbol{\theta}}_{mn})$ with respect to the preferred model estimate distribution. Denote this by

$$E_n(\hat{\mathbf{G}}_{mn}, [\hat{\mathbf{G}}_{*n}])$$

e-value is simply a functional of the distribution of this random evaluation function. Denote this by $e_n(\mathcal{M}_n)$.

We shall now elaborate on the following choice of the e-value:

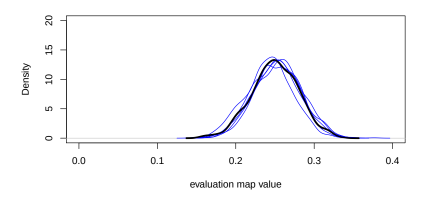
$$e_n(\mathcal{M}_n) = \mathbb{E} E_n(\hat{\mathbf{G}}_{mn}, [\hat{\mathbf{G}}_{*n}])$$

Results

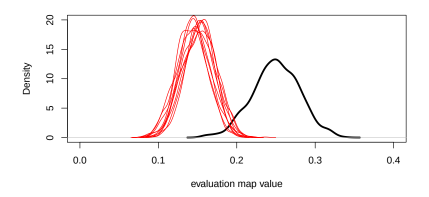
Theorem

Under some regularity conditions, as $n \to \infty$:

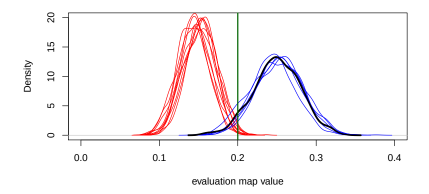
- For the preferred model $e_n(\mathcal{M}_{*n}) \to e_* < \infty$;
- ② For any adequate model, $|e_n(\mathcal{M}_n) e_n(\mathcal{M}_{*n})| \to 0$;
- **3** For any inadequate model, $e_n(\mathcal{M}_n) \to 0$.



For large enough n, e-values for all adequate models will be close to that of the preferred model.



But e-values of all inadequate models will be very small.



Thus we can choose an appropriate threshold ϵ_n such that any model with e-value below that threshold is inadequate, but e-value above the threshold implies the model is adequate.

Bootstrap estimation of e-values

We shall use bootstrap to generate multiple copies of the preferred model estimate and thus approximate $[\hat{\mathbf{G}}_{*n}]$.

Under standard regularity conditions (Chatterjee and Bose, 2005), we shall calculate the bootstrap estimate $\hat{\theta}_{rmn}$ by solving

$$\hat{oldsymbol{ heta}}_{rmn} = \mathop{\mathsf{argmin}}_{oldsymbol{ heta} \in \Theta_{mn}} \sum_{i=1}^{k_n} \mathbb{W}_{\mathit{rsni}} \Psi_{\mathit{sni}}(oldsymbol{ heta}, B_i)$$

where \mathbb{W}_{rsni} are i.i.d. weights chosen independently from the data satisfying:

$$\begin{array}{ll} \mathbb{E}\mathbb{W}_{rsn1} & = 1 \\ \mathbb{V}\mathbb{W}_{rsn1} & = \tau_{sn}^2\uparrow\infty \\ \tau_{sn}^2 & = o(a_{sn}^2), \\ \mathbb{E}\mathbb{W}_{rsn1}\mathbb{W}_{rsn2} & = O(k_n^{-1}), \\ \mathbb{E}\mathbb{W}_{rsn1}^2\mathbb{W}_{rsn2}^2 & \to 1, \\ \mathbb{E}\mathbb{W}_{rsn1}^4 & < \infty. \end{array}$$

This is the *Generalized bootstrap*.

Theorem

Suppose

$$\hat{e}_n(\mathcal{M}_n) = \mathbb{E}_r \mathcal{E}_n(\hat{\mathbf{G}}_{rmn}, [\hat{\mathbf{G}}_{r_1*n}])$$

based on two independent sets of bootstrap samples. Then for the above bootstrap scheme, as $n \to \infty$:

- For any adequate model, $|\hat{\mathbf{e}}_n(\mathcal{M}_n) \hat{\mathbf{e}}_n(\mathcal{M}_{*n})| \stackrel{P_n}{\rightarrow} o_P(1)$;
- ② For any inadequate model, $\hat{e}_n(\mathcal{M}_n) \stackrel{P_n}{\to} o_P(1)$.

where P_n is probability conditional on the data.

Fast bootstrap approximation of coefficient estimates

The bootstrap sample is related to the actual estimate through score vectors and hessian matrices:

$$\hat{\boldsymbol{\theta}}_{rsn} = \hat{\boldsymbol{\theta}}_{sn} - \frac{\tau_{sn}}{a_{sn}} \left[\sum_{i=1}^{k_n} \Psi_{sni}''(\hat{\boldsymbol{\theta}}_{sn}, B_i) \right]^{-1/2} \sum_{i=1}^{k_n} W_{rsni} \Psi_{sni}'(\hat{\boldsymbol{\theta}}_{sn}, B_i) + \mathbf{R}_n$$

with
$$\mathbb{E}_r \|\mathbf{R}_n\|^2 = o_P(1)$$
 and $W_{rsni} := (\mathbb{W}_{rsni} - 1)/\tau_{sn}$.

This means we can compute $\hat{\theta}_{rsn}$ just by generating Monte-Carlo samples and reusing other model objects. This makes the bootstrap procedure very fast.

Depth-based Inference May 18, 2017 Start with the preferred model estimate $\hat{\theta}_{*n} = (\hat{\theta}_{*n1}, \dots, \hat{\theta}_{*np_n})^T$. For any model \mathcal{M}_n , define $\hat{\theta}_{mn}$ as:

$$\hat{\boldsymbol{\theta}}_{mnj} = \left\{ egin{array}{ll} ext{Estimated } \hat{\boldsymbol{\theta}}_{*nj} & ext{ for } j \in \mathcal{S}_n; \\ ext{Known } c_{nj} & ext{ for } j
otin \mathcal{S}_n. \end{array}
ight.$$

We shall work with these plugin estimates. This saves time while still ensuring consistency of all model estimates.

Same for bootstrap versions:

$$\hat{\boldsymbol{\theta}}_{rmnj} = \left\{ egin{array}{ll} ext{Estimated } \hat{\boldsymbol{\theta}}_{r*nj} & ext{for } j \in \mathcal{S}_n; \\ ext{Known } c_{nj} & ext{for } j \notin \mathcal{S}_n. \end{array}
ight.$$

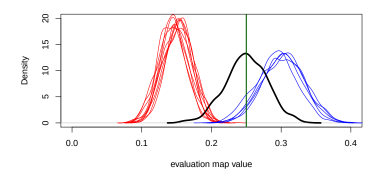
Now take fixed $p \equiv p_n$, and drop subscripts in \mathcal{M}_n , \mathcal{M}_{*n} etc. Take $E_n = D$, a depth function, and the full model as preferred model. Then

Theorem

For two nested models: \mathcal{M}_1 nested within \mathcal{M}_2 , we have $e_n(\mathcal{M}_1) > e_n(\mathcal{M}_2)$ for large enough n.

Notice now that all adequate models are by definiton nested within the preferred model. So $e_n(\mathcal{M}_*) < e_n(\mathcal{M})$ for any adequate model. Thus for large enough n, we can take threshold $\epsilon_n = e_n(\mathcal{M}_*)$ now.

- (1,2,3,0) preferred parameter vector
- (*,*,*,0) adequate model
- (*,*,*,*) full model



- Start with full model;
- ② Drop an essential predictor ⇒ Model becomes inadequate ⇒ e-value shifts to left:
- Orop a non-essential predictor ⇒ Model still adequate but nested within full model ⇒ e-value shifts to right.

Numerical studies and data applications

$$egin{aligned} \mathbf{Y}_i &= \mathbf{X}_i oldsymbol{eta} + oldsymbol{\epsilon} \in \mathbb{R}^{n_i} \ oldsymbol{\epsilon} &\sim \mathcal{N}(\mathbf{0}, \mathbf{V}_i); \quad \mathbf{V}_i = \sigma^2 \mathbf{I}_{n_i} + \mathbf{Z}_i \Delta \mathbf{Z}_i^{\mathsf{T}} \end{aligned}$$

- m subjects, n_i observations per subject, $n = m \times n_i$ total observations;
- $p = 9, \beta = (1, 1, 0, 0, 0, 0, 0, 0, 0)^T$;
- Elements of X₁,..., X_m chosen from Unif(-2,2), random effect design matrix Z_i is first 4 columns of X_i;

•

$$\Delta = \left(egin{array}{cccc} 9 & & & & \ 4.8 & 4 & & & \ 0.6 & 1 & 1 & & \ 0 & 0 & 0 & 0 \end{array}
ight)$$

- Two settings: (i) m = 30, $n_i = 5$, (ii) m = 60, $n_i = 10$;
- We use i.i.d. draws of Gamma(1,1) as bootstrap weights $W_i + 1$.

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Simulation results

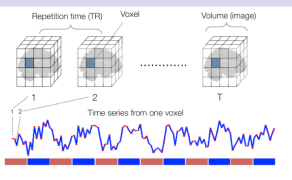
Method	Tuning	FPR%	FNR%	Model size	FPR%	FNR%	Model size
		$n_i = 5, m = 30$			$n_i = 10, m = 60$		
e-value based	$\tau_n/\sqrt{n}=1$	57.4	0.0	5.24	43.8	0.0	4.03
	2	30.4	0.0	3.32	12.3	0.0	2.42
	3	15.6	0.0	2.54	3.2	0.0	2.10
	4	7.3	0.0	2.24	1.0	0.0	2.03
	5	3.0	0.0	2.09	0.7	0.0	2.02
	6	1.7	0.0	2.05	0.3	0.0	2.01
	7	1.0	0.0	2.03	0.0	0.0	2.00
	8	0.7	0.0	2.02	0.0	0.0	2.00
	9	0.0	0.0	2.00	0.0	0.0	2.00
	10	0.0	0.0	2.00	0.0	0.0	2.00
Peng and Lu (2012)	BIC	21.5	9.9	2.26	1.5	1.9	2.10
	AIC	17	11.0	2.43	1.5	3.3	2.20
	GCV	20.5	6	2.30	1.5	3	2.18
	$\sqrt{\log n/n}$	21	15.6	2.67	1.5	4.1	2.26

Comparison between our method and that proposed by Peng and Lu (2012) through average false positive percentage, false negative percentage and model size

Method	τ_n/\sqrt{n}	Setting 1	Setting 2
e-value based	1	2	16
	2	36	67
	3	60	91
	4	80	97
	5	91	98
	6	95	99
	7	97	100
	7	98	100
	8	100	100
	10	100	100
Bondell et al. (2010)		73	83
Peng and Lu (2012)	49	86	
Fan and Li (2012)		90	100

Comparison of our method and three sparsity-based methods of mixed effect model selection through accuracy of selecting correct fixed effects

Data analysis: spatial dependency in fMRI visual task data



Dimensions of 3D voxel array $64 \times 64 \times 33$

- Data collected from 19 subjects, each of which were shown two types of pictures at certain time points within a test period (Wakeman and Henson, 2015).
- Each subject went through 9 runs. In each run 210 images of their brain were recorded in 2-second intervals.
- We use the data from a single run on subject 1, and perform a voxelwise analysis to find out the effect of time lags and activation of neighboring voxels on the activation of each voxel.

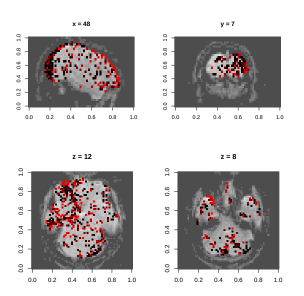
$$y_i(t) = x_{ia}(t)\beta_{ia} + x_{ib}(t)\beta_{ib} + \sum_{l=1}^{q} t^{l-1}\gamma_{il} + \sum_{n \in N_i} y_n(t)\delta_{i,n} + \epsilon_i(t)$$
$$= \tilde{\mathbf{x}}_i(t)^T \boldsymbol{\theta}_i + \epsilon_i(t)$$

- Want to quantify effect of immediate neighbors: N_i the set of neighbors of i-th voxel;
- Edge or corner voxels excluded: so 26 neighbors for a voxel, thus total 30 predictors.
- We estimate the set of non-zero coefficients in θ_i using our method. Suppose this set is R_i , and its subsets corresponding to neighbor and non-neighbor (i.e. stimuli and drift) terms are S_i and T_i , respectively.

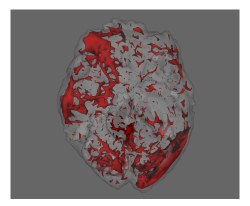
To quantify the effect of neighbors we now calculate the corresponding *F*-statistic:

$$F_i = \frac{(\sum_{n \in S_i} \tilde{x}_{i,n} \hat{\theta}_{i,n})^2}{(y_i(t) - \sum_{n \in T_i} \tilde{x}_{i,n} \hat{\theta}_{i,n})^2} \frac{|n - T_i|}{|S_i|}$$

and obtain its *p*-value, i.e. $P(F_i \ge F_{|S_i|,|n-T_i|})$.



Plot of significant p-values at $\alpha = 0.05$ at specified cross-sections



A smoothed surface obtained from the p-values clearly shows high spatial dependence in right optic nerve, auditory nerves, auditory cortex and left visual cortex areas as well as Cerebellum

- Introduction
- Signed Peripherality Functions
- Nonconvex Penalized Multitask Regression using Depth-based Penalty
- Generalized model discovery using statistical evaluation maps
- Selecting important SNPs from multi-SNP mixed models on Twin Studies GWAS data

Genome-Wide Association Studies (GWAS) based on families are used in behavioral genetics to control for environmental variation, thus requiring smaller sample size to detect Single Nucleotide Polymorphisms (SNP) responsible behind traits like alcoholism and drug addiction, and also to quantify gene-environment interaction.

Two challenges:

- SNPs highly correlated, weak signals of individual SNPs;
- Need to use mixed models to account for within-family dependence.

$$\begin{aligned} \mathbf{Y}_i &= \alpha + \mathbf{G}_i \boldsymbol{\beta}_g + \mathbf{C}_i \boldsymbol{\beta}_c + \boldsymbol{\epsilon}_i \\ \boldsymbol{\epsilon}_i &\sim \mathcal{N}_{n_i}(\mathbf{0}, \mathbf{V}_i); \quad \mathbf{V}_i = \sigma_a^2 \mathbf{\Phi}_i + \sigma_c^2 \mathbf{1} \mathbf{1}^T + \sigma_e^2 \mathbf{I}_{n_i} \end{aligned}$$

- Total m families, with the i-th pedigree containing n_i individuals;
- $\mathbf{Y}_i = (y_{i1}, \dots, y_{in_i})^T$ are the quantitative trait values for individuals in *i*-th pedigree, $\mathbf{G}_i \in \mathbb{R}^{n_i \times p_s}$ containing their genotypes for a bunch of SNPs, $\mathbf{C}_i \in \mathbb{R}^{n_i \times p}$ contain the data on individual-specific covariates;
- Three variance components correspond to polygenic effect due to other SNPs, shared environment effect and individual-specific effects. This is called the ACE model.

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 $\Phi_{Adopted} = \mathbf{I}_4$

$$\mathbf{\Phi}_{MZ} = \begin{bmatrix} 1 & 0 & 1/2 & 1/2 \\ 0 & 1 & 1/2 & 1/2 \\ 1/2 & 1/2 & 1 & 1 \\ 1/2 & 1/2 & 1 & 1 \end{bmatrix},$$

$$\mathbf{\Phi}_{DZ} = \begin{bmatrix} 1 & 0 & 1/2 & 1/2 \\ 0 & 1 & 1/2 & 1/2 \\ 1/2 & 1/2 & 1 & 1/2 \\ 1/2 & 1/2 & 1/2 & 1 \end{bmatrix},$$

 Φ_i depends on the type of the *i*-th family:

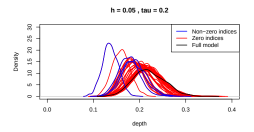
MZ = family with identical or monozygous twins, DZ = family with identical or dizygous twins.

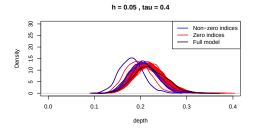
Objective

Want to detect the non-zero entries of β_a in the above model.

State-of-the-art is to perform single-SNP analysis and then correct for multiple testing. This loses power. We want to use the *e*-values method to improve that.

Problem: weak signal of individual SNPs





Using means as *e*-values makes the procedure very conservative.

But it may still be possible to use a tail quantile to distinguish between distributions. We now take as e-value the q-th quantile of the evaluation map distribution, for some fixed $q \in (0,1)$. Under slightly stronger assumptions than before, all results go through for a general evaluation map in the population and bootstrap worlds.

- Estimate the full model coefficient, say $\hat{eta}_g \equiv \hat{eta}$ (by R package regress)
- **②** Obtain its bootstrap distribution: $[\hat{\beta}]$;
- **3** Replace the *j*-th coefficient with 0, name it $\hat{\beta}_{-j}$. Do the same for its bootstrap distribution, say $[\hat{\beta}_{-j}]$. Repeat for all j;
- e-value of j-th covariate = tail probability of the q-th quantile of $[E(\hat{\beta}_{-j}, [\hat{\beta}])]$ with respect to $[E(\hat{\beta}, [\hat{\beta}])]$;
- Select j-th covariate if its e-value is less than tq, for some 0 < t < 1.

- 250 pedigrees, each of size 4: consisting of parents and MZ twins;
- $\alpha = 0$, no environmental covariates:
- 50 SNPs in correlated blocks of 6.4.6.4 and 30: MAF of SNPs in the blocks 0.2, 0.4, 0.4, 0.25 and 0.25;
- $\sigma_a^2 = 4, \sigma_c^2 = 1, \sigma_a^2 = 1;$
- First SNP of first 4 blocks are causal: each having heritability (a measure of magnitude of non-zero effect) h/6%;
- Full setup replicated 1000 times.
- Methods compared: mBIC2 - Variant of BIC that control false discovery rate at 0.05; RFGLS - Fast method of fitting single-SNP ACE models. Do Benjamini-Hochberg correction on *p*-values to control FDR at 0.05.

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Simulation results

6x	mBIC2	RFGLS	quantile e-values				
Heritability		+BH	q	t = 0.8	t = 0.7	t = 0.6	t = 0.5
			0.9	0.95/0.97	0.95/0.97	0.95/0.98	0.94/0.98
h = 10	0.79/0.99	0.95/0.92	0.5	0.96/0.97	0.96/0.98	0.95/0.98	0.94/0.98
			0.2	0.96/0.94	0.96/0.97	0.95/0.97	0.95/0.98
			0.9	0.72/0.95	0.7/0.96	0.69/0.96	0.66/0.97
h = 5	0.41/0.99	0.62/0.97	0.5	0.78/0.94	0.75/0.94	0.72/0.95	0.71/0.96
			0.2	0.83/0.91	0.78/0.94	0.75/0.95	0.73/0.95
			0.9	0.26/0.97	0.24/0.97	0.23/0.98	0.21/0.98
h = 2	0.11/0.99	0.14/0.99	0.5	0.34/0.95	0.28/0.96	0.27/0.97	0.26/0.97
			0.2	0.46/0.91	0.34/0.95	0.3/0.96	0.27/0.96
			0.9	0.12/0.98	0.1/0.98	0.09/0.99	0.08/0.99
h = 1	0.05/0.99	0.04/0.99	0.5	0.16/0.96	0.13/0.97	0.12/0.97	0.11/0.98
			0.2	0.25/0.93	0.16/0.96	0.13/0.97	0.13/0.97
			0.9	-/0.99	-/0.99	-/0.99	-/0.99
h = 0	-/0.99	-/0.99	0.5	-/0.98	-/0.98	-/0.99	-/0.99
			0.2	-/0.94	-/0.98	-/0.98	-/0.99

Average true positive/ true negative detection proportions over 1000 replications

Simulation results

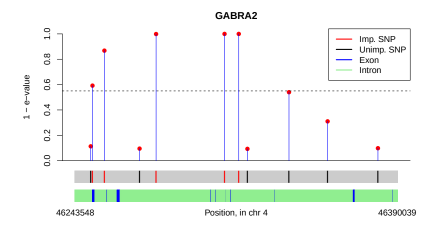
6x	mBIC2	RFGLS	quantile e-values				
Heritability		+BH	q	t = 0.8	t = 0.7	t = 0.6	t = 0.5
			0.9	0.96/0.97	0.96/0.97	0.95/0.98	0.94/0.98
h = 10	0.84/0.99	0.96/0.99	0.5	0.96/0.97	0.96/0.97	0.95/0.98	0.95/0.98
			0.2	0.97/0.95	0.96/0.97	0.96/0.97	0.95/0.98
			0.9	0.73/0.95	0.71/0.95	0.7/0.96	0.67/0.97
h = 5	0.48/0.99	0.64/0.99	0.5	0.79/0.93	0.76/0.94	0.73/0.95	0.72/0.95
			0.2	0.85/0.91	0.79/0.93	0.76/0.94	0.74/0.95
			0.9	0.29/0.96	0.27/0.97	0.25/0.98	0.23/0.98
h = 2	0.16/0.99	0.16/0.99	0.5	0.37/0.95	0.31/0.96	0.3/0.96	0.29/0.97
			0.2	0.53/0.91	0.38/0.95	0.33/0.95	0.3/0.96
			0.9	0.15/0.97	0.13/0.98	0.12/0.98	0.10/0.99
h = 1	0.08/0.99	0.05/0.99	0.5	0.2/0.96	0.17/0.97	0.15/0.97	0.13/0.98
			0.2	0.35/0.93	0.21/0.96	0.17/0.97	0.16/0.97
			0.9	-/0.97	-/0.98	-/0.98	-/0.99
h = 0	-/0.98	-/0.99	0.5	-/0.95	-/0.97	-/0.97	-/0.98
			0.2	-/0.90	-/0.95	-/0.97	-/0.97

Average true positive/ true negative detection proportions over 1000 replications: true positive = can detect any SNP in the same block

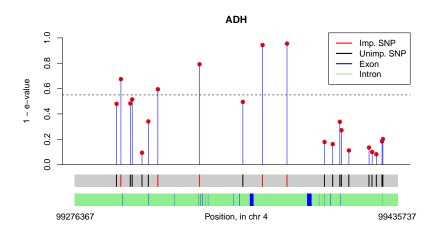
- Analyze data on families with MZ twins: 682 families;
- Response variable: amount of alcohol consumption;
- Look at models specific to well-studied genes for alcoholism: GABRA2, ADH1B, ADH1C, SLC6A3, SLC6A4, OPRM1, CYP2E1, DRD2, ALDH2, and COMT;
- Group together ADH genes as individual genes have very small number of SNPs. Also do SLC6A4+DRD2 together as they are known to interact with each other.
- We take q = 0.9, t = 0.5 to increase specificity, i.e. detection of true negatives.

SNP GABRA2 11/5 ADH 21/5 SLC6A3 18/4		
GABRA2 11/5 ADH 21/5 SLC6A3 18/4	Gene	Total/detected
ADH 21/5 SLC6A3 18/4		SNP
SLC6A3 18/4	GABRA2	11/5
	ADH	21/5
	SLC6A3	18/4
SLC6A4 5/0	SLC6A4	5/0
OPRM1 46/29	OPRM1	46/29
CYP2E1 9/5	CYP2E1	9/5
DRD2 17/0	DRD2	17/0
ALDH2 5/5	ALDH2	5/5
COMT 15/9	COMT	15/9
SLC6A4 + DRD2 22/0	SLC6A4 + DRD2	22/0

Table of analyzed genes and detected SNPs in them



Detects rs1808851 and rs279856, which have very high correlation with the well-known rs279858. This was missed by a previous analysis (Irons, 2012).



Detects rs13103626 and rs10516430 at positions 99317251 and 99337881 in chr 4: close to the well-known rs1229984 at position 99318162.

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Conclusion and future work

In this thesis, I have proposed several methods of using depths or depth-like quantities in traditional statistical inference, e.g. PCA, sparse regression, model selection.

Future works include:

- Formulation of depth in a general Hilbert space;
- Exploring different uses of the e-values framework: for example in multiple testing;
- Extension of the signed rank and depth-based penalization framework.

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