Schizophrenia

May, 2015

August 25, 2016

evtTADA

Extended

Schizophreni data

- extTADA pipeline.
- Apply extTADA to schizophrenia data.

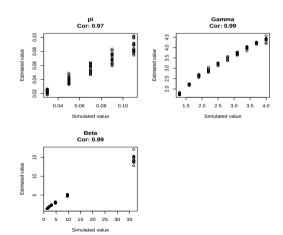
Steps

- De novo mutations: the same as original TADA (Or using binomial distribution).
- Inherited/Case-control:
 - 1 Only use a non heterogeneous population (obtain by using LM/GLM).
 - 2 Use an approximate model in the estimation process.
- Estimate all parameters using a MCMC method from Equation 1 (known risk genes are not necessary).

$$P(x|\phi_1,\phi_0) = \prod_{i=1}^{m} \left[\pi \prod_{j=1}^{K_1} f_{1_{DN_j}} \prod_{h=1}^{K_2} f_{1_{CC_h}} + (1-\pi) \prod_{j=1}^{K_1} f_{0_{DN_j}} \prod_{h=1}^{K_2} f_{0_{CC_h}} \right]$$
(1)

CC correlation

Use original TADA model to simulate => use approximate CC model estimate parameters.



Schizophrenia data Sample sizes from different studies.

Source	De novo	${\sf Non/Transmitted}$	Case	Control
Fromer et al. (2014)	617	617		
Girard et al. (2011)	14			
Gulsuner et al. (2013)	105			
McCarthy et al. (2014)	57			
Xu et al. (2012)	231			
Giulio et al. (2016)			4954	6239
Total	1024	617	4954	6239

TADA

Schizophrenia data

NoFXaC

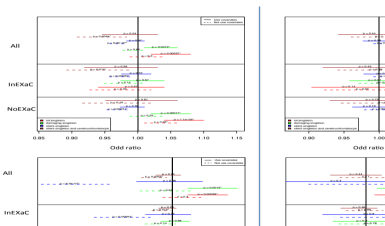
tot singleton
damaging singleton
sitent singleton
sitent singleton

-0.3

-0.2

-0.4

SCZ Case control



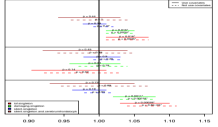
p = 1.1e-05*

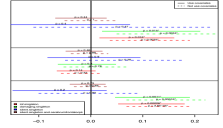
0.1

0.2

p = 0.2

-0.1





extTADA

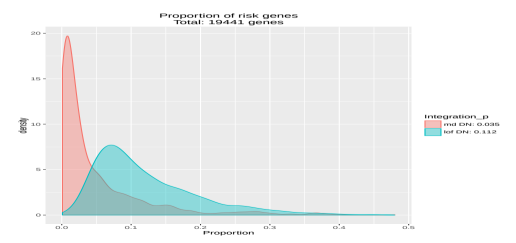
Extended

Schizophrenia data

Mutation	dnControl	dnCase	IOR	uOR	odds.ratio	pvalue
lof	43	111	1.34	2.87	1.94	0.000302
missense	334	612	1.45	2.15	1.77	7.04e-09
silent	134	227	0.994	1.62	1.27	0.0552
damaging missense (md)	31	100	1.6	3.83	2.44	1.21e-05
silentCerebellumocPk.narrowPeak	14	30	0.788	3.18	1.55	0.216
silentCerebrumfrontalocPk.narrowPeak (nr)	14	50	1.42	5.19	2.63	0.00109
silentFrontalcortexocPk.narrowPeak	21	55	1.13	3.37	1.92	0.0122

Estimate parameters for only SCZ CC

md DN: missense damaging de novo

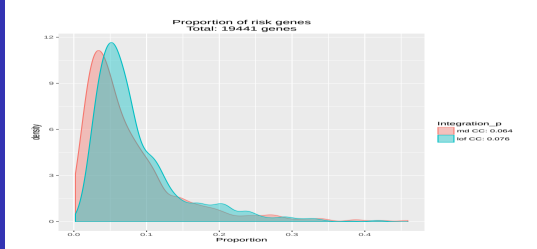


Estimate parameters for only SCZ CC

extTADA

Extend-

Schizophrenia data

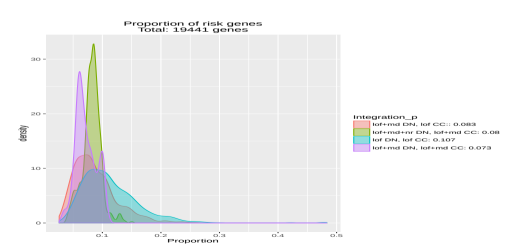


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Extende

Schizophrenia data

Estimate simultaneously parameters for SCZ DN and CC



data

How many risk genes

Previous studies (The unseen species problem):

- Xu et al (2012): **868 genes** based on 231 parent-proband trios and 34 unaffected trios.
- Fromer et al (2014): 4000 to 12000 genes based on 623 schizophrenia trios (use LoF and NS mutations).

Current study (extTADA)

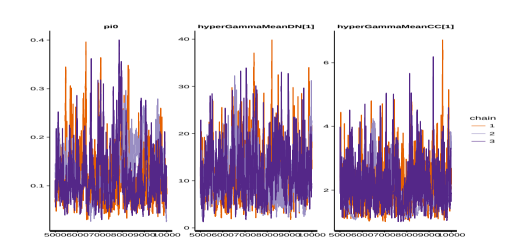
• \sim 2000 genes.

MCMC results for one DN + one CC

extTADA

Extende TADA

Schizophrenia data



Schizophrenia data

Top genes from extTADA

FDR
0.0545668538593245
0.0621595450057018
0.124379091145705
0.166929375517358
0.209294278939901
0.239767227415426
0.262757710600909
0.280827362530608
0.296109953864339

Schizophrenia data

- Choose top 100, 500, 1000 genes from extTADA results.
- Intersect with known gene sets.

Gene set	nGene	Top 100	Top 500	Top 1000
SNPsINdel.denovo.aut	2789	4.00E-05	1.00E-05	1.00E-05
SNPsINdel.denovo.scz	737	1.00E-05	1.00E-05	1.00E-05
celf4	2461	4.60E-04	1.00E-05	1.00E-05
constrained	936	1.00E-05	1.00E-05	1.00E-05
denovo.aut	2789	1.00E-05	1.00E-05	1.00E-05
denovo.scz	737	1.00E-05	1.00E-05	1.00E-05
fmrp	1194	1.00E-05	1.00E-05	1.00E-05
listLoFtolerantEXAC	2969	1.00E-05	1.00E-05	1.00E-05
listMcRae2016	74	1.00E-05	1.00E-05	1.00E-05
newListFromSupTable2	45	1.00E-05	1.00E-05	1.00E-05
nmdarc	74	5.00E-04	1.23E-01	1.38E-02
pLI09	3231	1.00E-05	1.00E-05	1.00E-05
rbfox13	3226	6.20E-04	1.00E-05	1.00E-05
rbfox2	2892	6.30E-04	1.00E-05	1.00E-05
synaptome	1812	2.01E-03	1.90E-04	6.00E-05

p-values were calculated by sampling 100,000 times.

