Write-Up ADDIS Re-run and Verification

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Operations in January and February

Re-analysis of GTEx Using MRPC-ADDIS

DETAILED DESCRIPTION:

Implementation of the ADDIS improvement to the MRPC algorithm: This involved adapting the existing scripts developed by M. Badsha to rerun the GTEx data using the ADDIS version of the FDR control. The distribution of model types (M0, M1, ... M4, Other) relative to each tissue was retained from the analyses for both MRPC-LOND and MRPC-ADDIS for comparison. Additionally, a host of programs were developed to identify the specific classification of each trio analyzed in each tissue. These programs were further adapted to identify the number of trios for each tissue classified as cis or trans mediated (M1 type 1 or M1 type 2). Such information was again compared with results from MRPC LOND with the intention of understanding the differences in inferred networks between the two FDR control methods.

In General, MRPC-ADDIS loosens the rejection threshold such that more edges/directions are inferred. There was typically a reshuffling of Model types with some specific trends such as most M1's whose graph classification was changed was converted to M2 or M4. M0's were converted to either M1, M2, M3, or M4 (with a large number in each tissue transferring to M3). The path by which M1 was converted to M2 or M4 was generally a direction flip between the cis and trans leading to a dependence structure of the cis gene dependent on both the trans gene and variant. The other method was the inference of an direct edge between the variant and both genes and a general edge between the cis and trans gene.

MAIN POINTS AND FILE LOCATIONS:

- Re-run of GTEx data with MRPC using the ADDIS FDR control
- → Done using a slightly modified version of M. Badsha's scripts.
- \rightarrow All associated output data files are stored at:

/mnt/ceph/jarredk/AddisReRunFiles

 \rightarrow The script used to Re-run **MRPC** using **ADDIS** is located at:

 $/mnt/ceph/jarredk/ADDIS_verify/ADDIS_ReRun.R$

- Summarizing of differences between LOND and ADDIS trio classifications:
- \rightarrow quantified the distribution of trio classifications for each tissue as counts of (M0, M1, ... M4, Other) \rightarrow table can be accessed through the cluster at:

/mnt/ceph/jarredk/ADDIS verify/Trios analysis GTEx v8 allPCs V3 ordered updated.xlsx

• followed up on FDR methods by investigating specific trios and their classification differences between FDR methods:

GENERAL STEPS:

- 1 read in SNP, PC, and final output files generated from ADDIS_rerun() (trio->model information)
- 2 catalog and store each trios LOND and ADDIS model classification
- 3 count/summarize trios between methods or within tissues (generally both)
- 4 rerun MRPC on trio using both **LOND** and **ADDIS** methods and compare graph and correlation matrices
- \rightarrow R-functions developed to assist in tracking trios are in:

$$/mnt/ceph/jarredk/ADDIS_verify/ADDIS_Post_Analysis_processing.R$$

 \bullet followed up and compared mediation model trios of Type-1 (cis mediated) and Type-2 (trans-mediated) for both **ADDIS** and **LOND**

GENERAL STEPS:

- 1 read final output files generated from ADDIS_rerun() for both FDR methods (trio->model information)
- 2 use MRPC to re-classify each trio identified as M1 to obtain causal graph matrix for next step
- 3 match trio to appropriate M1-type (1 or 2) by comparing causal graph matrix to type-1 and type-2 expected matrix
- 4 store M1-type classification information in a tiered list and save
- \rightarrow function to classify M1-type is called *ADDIS.M1.Check()* and is part of the R-script:

$$/mnt/ceph/jarredk/ADDIS_verify/ADDIS_Post_Analysis_processing.R$$

 \rightarrow tables summarizing the counts and relative proportions of cis/trans mediated M1 models can be found at:

$$/mnt/ceph/jarredk/ADDIS_verify/$$

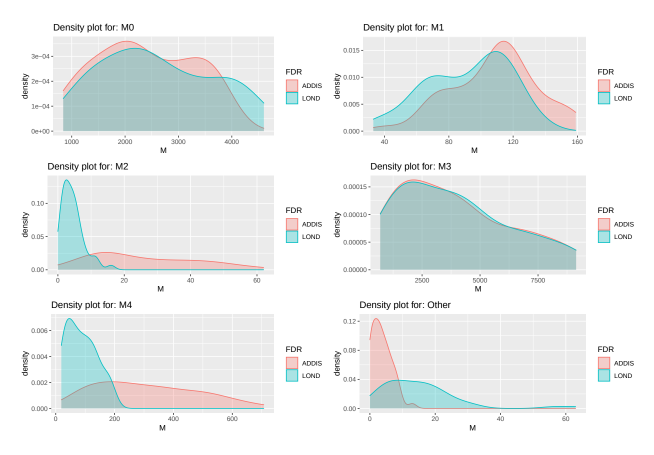


Figure 1: Density plots showing the distribution of each graph type under both ADDIS and LOND

Table 1: table of the average difference in count and percentage between ADDIS and LOND and their standard errors for the differences

	M0	M1	M2	M3	M4	Other
mean.change.ct	-235.06250	16.47917	21.41667	-13.83333	222.45833	-11.45833
SD.change.ct	187.55248	12.39078	13.72191	127.75214	121.25373	10.03814
$\rm mean.change\%$	-0.01742	0.00116	0.00166	-0.00107	0.01653	-0.00085
SD.change%	0.02104	0.00200	0.00186	0.01165	0.01706	0.00125

Table 2: Table of confidence intervals for the average count of each classified graph for all tissues under the LOND FDR control

	lower.limit	mean	upper.limit
$\overline{\text{M0}}$	2604.155	2613.533	2622.910
M1	91.490	91.718	91.946
M2	4.442	4.471	4.499
M3	3914.535	3934.987	3955.439
M4	88.627	89.077	89.526
Other	14.995	15.102	15.209

Table 3: Table of confidence intervals for the average count of each classified graph for all tissues under the ADDIS FDR control $\,$

lower.limit mean upp	
	er.limit
M0 2368.250 2376.159 2	384.068
M1 107.943 108.171	108.400
M2 25.807 25.952	26.097
M3 3890.639 3911.791 3	932.943
M4 310.763 312.288	313.813
Other 3.636 3.662	3.689