# **Transcriptome-Wide Association Study Pipeline (TWAS-pipeline)**

#### **INSTALLATIONS**

```
--- TWAS ---
```

The following packages by the developers

Package

s URLs

TWAS https://data.broadinstitute.org/alkesgroup/TWAS/TWAS.2016\_02\_24.tar.bz2

weight https://data.broadinstitute.org/alkesgroup/TWAS/

files

z-score https://data.broadinstitute.org/alkesgroup/TWAS/ETC/CLEAN\_ZSCORES.tar.bz

clean 2 program

are required and be unpacked. In addition, lists of genes in the three populations are made through the following scripts,

```
TWAS=/genetics/bin/TWAS
cd $TWAS
for pop in MET NTR YFS
do
    ls WEIGHTS_$1 | sed 's\/\\g' > $pop.lst
done
```

--- **TWAS-pipeline** ---. The pipeline is installed as follows,

```
git clone https://github.com/jinghuazhao/TWAS-pipeline
```

On our system, TWAS.sh and TWAS\_get\_weights.sh for TWAS and twas.sh, twas2.sh, twas2-collect.sh and twas2-1.sh for TWAS-pipeline have symbolic links under /genetics/bin and available from the \$PATH environment. A Stata equivalent has been developed by Dr lian'an Luan.

To accommodate the suggestion of p value in accordance with the Z-score in the output, pnorm.c is included which can be compiled as follows,

```
gcc pnorm.c -lm -o pnorm
```

and a call pnorm z\_score yields a p value with more decimal places. However, the function is now part of twas-single.sh.

--- **GNU Parallel** ---. Further information is available from here.

#### **RUNNING THE PIPELINE**

Suppose you have a file containing GWAS summary statistics, you can run the pipeline as follows,

```
twas.sh input_file
```

where the input\_file is in tab-delimited format containing SNP\_name, SNP\_pos, Ref\_allele, Alt allele, Beta and SE. The output will be contained in <input file>.imp.

This assumes that ssh can access nodes in a clusters freely and in case this has not been done, a single node mode is more appropriate,

twas-single.sh input\_file

## **TWAS USING GTEX**

This is achieved with gtex.sh and gtex.subs using weights from the GTEx project. File GTEX\_WEIGHTS.bim.gz was created by GTEX\_WEIGHTS.sh in accordance with reference .bim files used by TWAS and GTEX\_WEIGHTS.1st was created to facilitate the imputation.

### **ADDITIONAL TOPICS**

These are available from the wiki page,

- An example on body bone mineral density
- An exposition with GIANT data
- Building reference panel
- Epigenomewide association
- FUSION pipeline

## **ACKNOWLEDGEMENTS**

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#### **REFERENCE**

Gusev A, et al. (2016). Integrative approaches for large-scale transcriptome-wide association studies. Nature Genetics (a copy at Harvard), 48, 245-252