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.

--- **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.lst 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

The work is possible with an EWAS project within the MRC Epidemiology Unit, for which colleagues and collaborators have contributed.

REFERENCES

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