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About

OSCA (OmicS-data-based Complex trait Analysis) is a software tool written in C/C++ for the analysis of complex traits using multi-omics data. It is developed by Futao Zhang, Zhihong Zhu and Jian Yang at Institute for Molecular Bioscience, The University of Queensland. Bug reports or questions: Jian Yang <jian.yang@uq.edu.au>.

Functions currently supported are:

- Estimating the epigenetic (or transcriptomic) relationships between individuals from genome-wide DNA methylation (or gene expression) data.
- Estimating the proportion of phenotypic variance for a complex trait can be "explained" by all DNA methylation (or gene expression) probes.
- Mixed linear model based analysis to test for associations between DNA methylation (or gene expression) probes and complex traits.
- Estimating the joint "effects" of all methylation (transcription) probes on a phenotype (e.g. BMI) in a mixed linear model (analogous to BLUP). These estimated effects can be used to predict the phenotype in a new independent sample.
- eQTL/mQTL analysis with linear regression model. The model can adjust for the relevant covarites. The results are saved in SMR dense BESD format.
- vQTL analysis to test the homogeneity of variance. Three algorithms have been implemented (Bartlett's test, Levene's test and Brown–Forsythe test, Fligner-Killeen test). The results are saved in SMR dense BESD format.

Note: Although this software tool is designed for gene expression and DNA methylation data, it can be applied to any other source of omics data including microbiome, proteome and brain connectome.

Credits

Futao Zhang and Jian Yang developed the methods, software and webpage. Zhihong Zhu contributed to the development of the ORM and OREML methods. Ting Qi contributed to MeCS method. Huanwei Wang contributed to vQTL method. Zhili Zheng provided the template of the website.

Questions and Help Requests

Bug reports or questions to Jian Yang (jian.yang@uq.edu.au) at Institute for Molecular Bioscience, The University of Queensland.

Citations

EWAS method: Zhang, F. et al. (2017) OSCA: a tool for omics-data-based complex trait analysis. In preparation.

Download

Executable Files (version 0.38)

The Linux version is available at: osca_Linux.zip.

The MacOS version is available at: osca_Mac.zip.

The Windows version is available at: osca_Win.zip.

The example files are in example.zip.

The executable files (binary code) are release under MIT lincense.

Update log

- 1. 21 July, 2017: first release.
- 2. 8 January, 2018: added meta analysis, eQTL analysis and vQTL analysis.

Data Management

The DNA methylation (or gene expression) data are stored in a binary format for storage efficiency. We store the data in three separate files .oii (individual information, similar as a PLINK .fam file), .opi (probe information) and .bod (a binary file to store the DNA methylation or gene expression profiles).

To efficiently save the result of eQTL / vQTL analysis, OSCA also supports SMR BESD format.

BOD format

BOD format: an efficient format to store omics data

myeed.oii

```
F01 I01 0 0 NA
F02 I01 0 0 NA
F03 I02 0 0 NA
...
```

Columns are family ID, individual ID, paternal ID, maternal ID and sex (1=male; 2=female; 0=unknown). Missing data are represented by "NA".

myeed.opi

```
1 probe101 924243 Gene01 +
1 probe102 939564 Gene01 -
1 probe103 1130681 Gene01 -
...
```

Columns are chromosome, probe ID (can be the ID of an exon or a transcript for RNA-seq data), physical position, gene ID and gene orientation.

myeed.bod

DNA methylation (or gene expression) data in binary format. Please do not try to open this file with a text editor.

Note: The text below are for advanced users only. We use the first 12 bytes of the .bod file to store the descriptive information for the data. The first 2 bytes are reserved for further extension of the software. The type of data is indicated by the 3rd byte (0 for gene expression data, 1 for DNA methylation data, and 2 for any other type of data). The type of value is indicated by the 4th byte (0 for DNA methylation beta value, 1 for DNA methylation m value and 2 for any other type of value). The number of individuals and number of probes are stored as integers (4 bytes each). For example, for a DNA methylation data set (0x01) of m values (0x01) on 1,342 (0x53e) individuals and 228,694 (0x37d56) CpG sites, the first 12 bytes of the .bod file are:

```
0101 0000 3e05 0000 567d 0300
```

The bytes afterwards are for the DNA methylation or gene expression data.

Make a BOD file

compile data in binary format from text format

```
osca --efile myprofile.txt --methylation-beta --make-bod --out myprofile
```

- --efile reads a DNA methylation (or gene expression) data file in plain text format.
- --methylation-beta indicates methylation beta values in the file.
- --make-bod saves DNA methylation (or gene expression) data in binary format.
- --out saves data (or results) in a file.

myprofile.txt

```
FID IID cg00000658 cg26036652 cg00489772 ...
F01 I01 0.909 0.845 0.41 ...
F01 I02 0.832 0.732 0.503 ...
...
```

This is a file with a header line that contains family ID, individual ID and names of probes. The column of family ID is optional. Please use the flag "--no-fid" for data without family ID.

```
osca ——efile myprofile.txt ——methylation—beta ——make—bod ——no—fid ——out myprofile
```

--no-fid indicates data without family ID.

```
osca --efile myprofile.txt --methylation-m --make-bod --out myprofile
```

--methylation-m indicates DNA methylation m values in the file.

If the profile is of gene expression value.

```
osca --efile myprofile.txt --gene-expression --make-bod --out myprofile
```

--gene-expression indicates gene expression profiles in the file.

For any other type of data:

```
osca --efile myprofile.txt --make-bod --out myprofile
```

A binary file also can be made from a transposed file in text format.

```
osca --tefile mytprofile.txt --methylation-beta --make-bod --no-fid --out myprofile
```

mytprofile.txt

```
FID F01 F01 ...

IID I01 I02 ...

cg00000658 0.909 0.832 ...

cg26036652 0.845 0.732 ...

cg00489772 0.41 0.503 ...
```

The row of family ID is optional. Please use the flag "--no-fid" if there is no family ID in your data.

```
osca — tefile mytprofile.txt — methylation—beta — make—bod — no-fid — out myprofile
```

update probe annotation

Probe information are sometimes not available in the original DNA methylation (or gene expression) data. These information can be updated the command below.

```
osca --befile myprofile --update-opi annotated.opi
```

- --befile reads a DNA methylation (or gene expression) data file in binary format.
- --update-opi reads a fully annotated .opi file.

Manage BOD file(s)

make a subset

To extract a probe

```
osca ——befile myprofile ——probe cg00000658 ——make—bod ——out mysubprofile
```

--probe extracts a specified probe.

To exclude a probe

```
osca ——befile myprofile ——probe—rm cg00000658 ——make—bod ——out mysubprofile
```

--probe-rm excludes a specified probe.

To extract a subset of probes

```
osca --befile myprofile --extract-probe probe.list --make-bod --out mysubprofile
```

--extract-probe extracts a subset of probes.

probe.list

```
cg00000658
cg26036652
...
```

To exclude a subset of probes

```
osca --befile myprofile --exclude-probe probe.list --make-bod --out mysubprofile
```

--exclude-probe excludes a subset of probes.

To extract a subset of individuals

```
osca ——befile myprofile ——keep indi.list ——make—bod ——out mysubprofile
```

--keep extracts a subset of individuals.

indi.list

```
F01 I01
F01 I02
```

To exclude a subset of individuals

```
osca --befile myprofile --remove indi.list --make-bod --out mysubprofile
```

--remove excludes a subset of individuals.

To extract a subset of genes

```
osca --befile myprofile --genes gene.list --make-bod --out mysubprofile
```

--genes extracts a subset of genes.

gene.list

```
MAN1B1
EDEM2
...
```

To extract a gene

```
osca --befile myprofile --gene MAN1B1 --make-bod --out mysubprofile
```

--gene extracts a gene.

To extract a subset of probes based on the order

```
osca --befile myprofile --from-probe probe0 --to-probe probe1 --make-bod --out mysubprofile
```

- --from-probe specifies the start probe.
- --to-probe specifies the end probe.

To extract a subset of probes in a genomic region centred at a specific probe

```
osca --befile myprofile --probe cg00000658 --probe-wind 500 --make-bod --out mysubprofile
```

--probe-wind defines a window \<kb> centred on a specified probe.

To extract a subset of probes on a chromosome

```
osca --befile myprofile --chr 1 --make-bod --out mysubprofile
```

--chr specifies a chromosome to select probes.

To extract a subset of probes based on physical positions

```
osca --befile myprofile --from-probe-kb 100 --to-probe-kb 200 --make-bod --out mysubprofile
```

- --from-probe-kb specifies the start physical position of the probes.
- --to-probe-kb specifies the end physical position of the probes.

merge BOD files

```
osca --befile-flist mybod.flist --make-bod --out myprofile
```

--befile-flist reads a file to get the full paths of the binary files.

mybod.flist

```
path1/my_bod1
path2/my_bod2
...
```

Quality control

To remvoe probes with low variance

```
osca --befile myprofile --std 0.02 --make-bod --out newprofile
```

--std removes the probes with the standard deviation smaller than a specified threshold.

To remove probes with a missing rate threshold

```
osca --befile myprofile --missing-ratio-probe 0.01 --make-bod --out newprofile
```

--missing-ratio-probe specifies a missing proportion threshold to remove probes.

Quality control of methyaltion data

To remove constitutively methylated/unmethylated probes

```
osca --befile myprofile --upper-beta 0.8 --lower-beta 0.2 --make-bod --out newprofile
```

- --upper-beta removes the DNA methylation probes with the mean beta value larger than a specified threshold.
- --lower-beta removes the DNA methylation probes with the mean beta value smaller than a specified threshold.

To QC with detection p-values

```
osca --befile myprofile --detection-pval-file dpval.txt --dpval-mth 0 --dpval-thresh 0.05 --make-bod --out newprofile
```

- --detection-pval-file reads a file that contains DNA methylation detection p-values.
- --dpval-mth specifies a method to do quality control with the detection p-values, 0 for removing the probes with one or more detection p-values violating a threshold (default as 0.05), and 1 for dropping the samples violating a proportion threshold (default as 1%) and simultaneously dropping the probes violating a proportion threshold (default as 1%).
- --dpval-thresh specifies a threshold of detection p-value.

```
osca --befile myprofile --detection-pval-file dpval.txt --dpval-mth 1 --ratio-probe 0.01 --ratio-sample 0.01 --dpval-thresh 0.05 --make-bod --out newprofile
```

- --ratio-probe specifies a proportion threshold to remove probes.
- --ratio-samplespecifies a proportion threshold to remove individuals.

the file "dpval.txt" is in the same format with a transposed profile file. Option "--no-fid" is also valid here.

adjust the covariates for each probe

```
osca —befile myprofile —covar my.cov —qcovar my.qcov —adj-probe —make—bod —out newprofile
```

--adj-probe adjusts the covariates for each probe.

other options

Transform methylation beta value to methylation m value and vice versa

```
osca --befile myprofile --m2beta --make-bod --out newprofile
```

--m2beta calculates the methylation beta value from the methylation m value.

```
osca --befile myprofile --beta2m --make-bod --out newprofile
```

--beta2m calculates the methylation m value from the methylation beta value.

Query a BOD file

make a text format file

```
osca --befile myprofile --make-efile --out myprofile.txt
```

--make-efile saves the DNA methylation (or gene expression) data in text format.

--make-tefile saves the DNA methylation (or gene expression) data in transposed text format.

NOTE: The options in Manage BOD file(s) can also be applied.

osca --befile myprofile --make-tefile --out mytprofile.txt

Calculate the variance and the mean of probes

```
osca --befile myprofile --get-variance --get-mean --out newprofile
```

- --get-variance calculates the variance of each probe.
- --get-mean calculates the mean of each probe.

newprofile.var.txt

```
cg00000957 0.000812127
cg00001349 0.00560701
```

newprofile.mean.txt

```
cg00000957 0.901574
cg00001349 0.860279
...
```

BESD format

For the detail information please refer to SMR BESD format. To efficiently save the eQTL / v QTL result, we extended the BESD format.

Note: The text below are for advanced users only.

We use the first 4 bytes of the .besd file to store the descriptive information for the data.

BESD dense format 1 (OSCA and SMR supported): the descriptive information is 0x00000000 and the data onwards is a vector of beta values followed by a vector of SE values of each probe across all the snps. The beta and se values are saved as float-precision (4B).

BESD dense format 2 (OSCA supported): the descriptive information is 0x00000004 and the data onwards is the beta value followed by the SE value of each SNP across all the probes. The beta and se values are saved as float-precision (4B).

BESD sparse format 1 (OSCA and SMR supported): the descriptive information is 0x40400000. The beta values and the SE values as saved in the CSC format

BESD sparse format 2 (OSCA supported): the descriptive information is 0x00000003. The beta values and the SE values as saved in the CSC format with removing the redundant information.

Query a BESD file

This feasuer is memory efficent even to query with huge dense BESD file.

Command line options for SNPs

To query the eQTL resutls for a single SNP, we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --snp rs123 --out myquery
```

- --query saves in text format a subset of the eQTL summary dataset based on the specified eQTL p-value threshold. The default value is 5.0e-8.
- --snp specifies a single SNP.

myquery.txt

```
SNP Chr BP A1 A2 Freq Probe Probe_Chr Probe_bp Gene Orientation b se p
rs01 1 1001 A G 0.23 cg01 1 1101 gene1 + -0.033 0.006 3.8e-08
rs01 1 1001 A G 0.06 cg02 1 1201 gene2 - 0.043 0.007 8.1e-10
.....
```

To query the eQTL resutls excluding a single SNP, we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --snp-rm rs123 --out myquery
```

--snp-rm specifies a single SNP to exclude.

To query the eQTL resutls extracting a subset of SNPs , we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --extract-snp mysnp.list --out myquery
```

--extract-snp extracts a subset of SNPs for analysis.

To query the eQTL resutls excluding a subset of SNPs , we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --exclude-snp mysnp.list --out myquery
```

--exclude-snp excludes a subset of SNPs from analysis.

To query eQTL resutls for a range of SNPs in a genomic region

```
smr --beqtl-summary myeqtl --query 5.0e-8 --from-snp rs123 --to-snp rs456 --out myquery
```

- --from-snp specifies the start SNP.
- --to-snp specifies the end SNP.

NOTE: All SNPs should be on the same chromosome.

To query eQTL results for all SNP on a chromosome

```
smr --beqtl-summary myeqtl --query 5.0e-8 --snp-chr 1
```

--snp-chr specifies a chromosome to select SNPs.

NOTE: The probes in the result could be on the other chromosomes if there are *trans-eQTLs*.

To query SNPs based on physical positions

```
smr --beqtl-summary myeqtl --query 5.0e-8 --snp-chr 1 --from-snp-kb 100 --to-snp-kb 200 --out myquery
```

- --from-snp-kb specifies the start physical position of the region.
- --to-snp-kb specifies the end physical position of the region.

NOTE: You will need to specify a chromosome (using the '--snp-chr' option) when using this option.

To guery based on a flanking region of a SNP

```
smr ——beqtl—summary myeqtl ——query 5.0e—8 ——snp rs123 ——snp—wind 50 ——out myquery
```

--snp-wind defines a window centred on a specified SNP.

Command line options for probes

To query based on a single probe

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe cg123 --out myquery
```

--probe specifies a single probe.

To query excluding a single probe

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe-rm cg123 --out myquery
```

--probe-rm specifies a single probe to exclude.

To query the eQTL resutls extracting a subset of probes , we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --extract-probe myprobe.list --out myquery
```

--extract-probe extracts a subset of probes for analysis.

To query the eQTL resutls excluding a subset of probes , we could use this command

```
smr --beqtl-summary myeqtl --query 5.0e-8 --exclude-probe myprobe.list --out myquery
```

--exclude-probe excludes a subset of probes from analysis.

To query based on a range of probes

```
smr --beqtl-summary myeqtl --query 5.0e-8 --from-probe cg123 --to-probe cg456 --out myquery
```

- --from-probe specifies the start probe.
- --to-probe specifies the end probe.

NOTE: All probes should be on the same chromosome.

To query based on a chromosome

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe-chr 1
```

--probe-chr specifies a chromosome to select probes.

NOTE: The SNPs in the result could be on the other chromosomes if there are trans-eQTLs.

To query based on physical positions of the probes

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe-chr 1 --from-probe-kb 1000 --to-probe-kb 2000 --out myquery
```

- --from-probe-kb specifies the start physical position of the probes.
- --to-probe-kb specifies the end physical position of the probes.

NOTE: You will need to specify a chromosome (using the '--probe-chr' option) when using this option.

To query based on a flanking region of a probe

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe cg123 --probe-wind 1000 --out myquery
```

--probe-wind defines a window centred on a specified probe.

To query based on a gene

```
smr --beqtl-summary myeqtl --query 5.0e-8 --gene gene1 --out myquery
```

--gene specifies a single gene to select probes.

Command line option for cis-region

```
smr --beqtl-summary myeqtl --query 5.0e-8 --probe cg123 --cis-wind 2000 --out myquery
```

File-list options

To query based on a list of SNPs

```
smr --beqtl-summary myeqtl --extract-snp snp.list --query 5.0e-8 --out myquery
```

To query based on a list of probes

```
smr --beqtl-summary myeqtl --extract-probe probe.list --query 5.0e-8 --out myquery
```

To gurey based on a list of genes

```
smr ——beqtl—summary myeqtl ——genes gene.list ——query 5.0e—8 ——out myquery
```

--genes extracts a subset of probes which tag the genes in the list.

gene.list

```
gene1
gene2
gene3
...
```

Manage a BESD file

make a subset of data

All the data management options in Query a BESD file can be here.

for example, To extract a subset of SNPs and/or probes

```
smr --beqtl-summary myeqtl --extract-snp mysnp.list --extract-probe myprobe.list --make-besd --out mybesd
```

tansform to SMR compatible format

```
smr --beqtl-summary myeqtl --make-besd --to-smr --out mybesd
```

--to-smr tranforms BESD file to SMR compatible format.

shrink a BESD file

To remove probes without any value across all the SNPs and SNPs without any value across all the probes.

```
smr --beqtl-summary myeqtl --make-besd --besd-shrink --out mybesd
```

--besd-shrink removes probes and SNPs that have no value.

ORM

estimate ORM

```
osca ——befile myprofile ——make—orm ——out myorm
```

```
osca --befile myprofile --make-orm-bin --out myorm
```

--make-orm or --make-orm-bin estimates the omics relationship matrix (ORM) between pairs of individuals from a set of probes and save the lower triangle elements of ORM to binary files. This format is compatible with the GRM in the GCTA software.

```
osca --befile myprofile --make-orm-gz --out myorm
```

--make-orm-gz estimates the ORM and save the lower triangle elements of ORM to compressed plain text files.

```
osca --befile myprofile --make-orm --orm-alg 1 --out myorm
```

--orm-alg specifies the algorithm to estimate the ORM. 1 for standardized data of each probe, 2 for centred data of each probe and 3 for standardized data of each individual. The default option is 1.

Note that although we describe the options above using DNA methylation and gene expression data, all the options can be applied to any other source of omics data mentioned above.

Principal Component Analysis

```
osca --orm myorm --pca 20 --out mypca
```

- --orm reads the ORM binary files.
- --pca conducts principal component analysis and saves the first n (default as 20) PCs.

mypca.eigenval

```
122.014
95.3064
70.8055
...
```

mypca.eigenvec

```
R06C01 R06C01 0.012637 -0.00328532 0.0263842 -0.00595246
R05C02 R05C02 0.0106692 -0.0184545 -0.0159826 0.013951
R04C02 R04C02 0.0024727 -0.0118185 -0.0256503 -0.0106235
...
```

Users can also manipulate the ORM in the analysis.

osca --orm myorm --remove indi.list --pca 20 --out mypca

```
osca --orm myorm --keep indi.list --pca 20 --out mypca
```

```
osca --orm myorm --orm-cutoff 0.05 --pca 20 --out mypca
```

--orm-cutoff removes one of a pair of individuals with estimated omics relationships larger than the specified cut-off value.

```
osca ——merge-orm myorm.flist ——pca 20 ——out mypca
```

--merge-orm reads multiple ORMs in binary format.

OREML

```
osca --reml --pheno my.phen --out myreml
```

- --reml performs REML (restricted maximum likelihood) analysis. This option is usually followed by the option --orm (one ORM) or --merge-orm (multiple ORMs) to estimate the variance explained by the probes that were used to estimate the omics relationship matrix.
- --pheno reads phenotype data from a plain text file. Missing value should be represented by "NA".

my.phen

```
R06C01 R06C01 32.6332
R05C02 R05C02 23.9411
R04C02 R04C02 29.7441
```

```
osca --reml --orm myorm --pheno my.phen --mpheno 1 --out myreml

--mpheno reads a list of comma-delimited trait numbers if the phenotype file contains more than one trait, e.g. "1,3" tells OSCA to take the first and the third trait
```

NOTE: current version only supports single trait analysis in one run.

for analysis. OSCA always takes the first trait for analysis unless this option is specified.

```
osca --reml --orm myorm --pheno my.phen --out myreml
osca --reml --orm myorm --pheno my.phen --keep indi.list --out myreml
```

```
osca --reml --orm myorm --pheno my.phen --orm-cutoff 0.05 --out myreml
```

With multiple ORMs

```
osca — reml — merge-orm myorm.flist — pheno my.phen — reml—alg 0 — out myreml
```

--reml-alg specifies the algorithm to do REML iterations, 0 for average information (AI), 1 for Fisher-scoring and 2 for EM. The default option is 0, i.e. AI-REML, if this option is not specified.

```
osca --reml --merge-orm myorm.flist --pheno my.phen --reml-maxit 100 --out myreml
```

--reml-maxit specifies the maximum number of iterations. The default number is 100 if this option is not specified.

```
osca --reml --orm myorm --pheno my.phen --covar my.covar --out myreml
```

--covar reads discrete covariates from a plain text file.

my.covar

```
R06C01 R06C01 F 0
R05C02 R05C02 F 1
R04C02 R04C02 M 1
...

osca --reml --orm myorm --pheno my.phen --qcovar my.qcovar --out myreml
```

--qcovar reads quantitative covariates from a plain text file.

my.qcovar

```
R06C01 R06C01 25
R05C02 R05C02 16
R04C02 R04C02 30
...
```

osca --reml --orm myorm --pheno my.phen --reml-est-fix --out myreml

--reml-est-fix displays the estimates of fixed effects on the screen.

```
osca --reml --orm myorm --pheno my.phen --reml-no-lrt --out myreml
```

--reml-no-lrt turns off the LRT.

EWAS

Mixed Linear Model Association

```
osca ——mlma ——befile myprofile ——pheno my.phen ——out my
```

If you have already computed the ORM

```
osca --mlma --befile myprofile --pheno my.phen --orm myorm --out my
```

--mlma initiates an MLM based association analysis including the target probe (the probe to be tested for association) in the ORM. The results will be saved in a plain text file with .mlma as the filename extension.

my.mlma

```
Chr Probe bp Gene Orientation b se p
1 cg00003287 201346149 TNNT2 - -0.156 0.597 0.794
1 cg00008647 207082900 IL24 + 0.032 0.354 0.926
1 cg00009292 50882082 DMRTA2 - 0.120 1.182 0.919
...
```

This is a text file with headers. Columns are chromosome, probe, probe BP, gene, orientation, effect size, standard error and p-value.

```
osca --mlma-loco --befile myprofile --pheno my.phen --out my
```

--mlma-loco initiates an MLM based association analysis with the chromosome where the target probe is located excluded from the ORM. The results will be saved in a plain text file with .loco.mlma as the filename extension.

```
osca --mlma --lxpo 0.1 --befile myprofile --pheno my.phen --out my
```

```
osca --mlma-loco --lxpo 0.1 --befile myprofile --pheno my.phen --out my
```

--lxpo specifies a percentage of probes to exclude from calculating the ORM. This option should accompany --mlma or --mlma or

Linear Regression

```
osca --befile myprofile --pheno my.phen --linear --out my
```

```
osca --befile myprofile --pheno my.phen --qcovar my.qcovar --covar my.covar --linear --out my
```

--linear saves linear regression statistics to a plain text file.

my.linear

```
probeChr
          ProbeID Probe_bp
                            BETA
                                    SE P NMISS
                                         9.11e-01
   cg00003287 201346149
                          -0.0594 0.531
                                                     1337
                                                    1337
   cg00008647 207082900 0.5003 0.263
                                         5.72e-02
   cq00009292 50882082
1
                          1.0512 1.026
                                         3.06e-01
                                                    1337
```

This is a text file with headers. Columns are chromosome, probe, probe BP, effect size, standard error, p-value and number of non-missing individuals.

EWAS simulation

The phenotypes are simulated based on a set of real DNA methylation (or gene expression) data and a simple model $y = sum(x_ib_i) + \epsilon$, where y is a vector of phenotypes, x_i is a vector of raw DNA methylation (or gene expression) profile or standardized profile of the i-th "causal" probe, b_i is the effect of the i-th causal probe and $\epsilon \sim N(0, var(sum(x_ib_i))(1/h^2-1))$ is a vector of residual effect.

```
osca --simu-qt --simu-hsq 0.1 --befile myprofile --simu-causal-loci mycausal.list --out mypheno
```

- --simu-qt smulates a quantitative trait.
- --simu-hsq specifies the proportion of variance in phenotype explained by the causal probes. The default value is 0.1.
- --simu-causal-loci reads a list of probes as causal probes. If the effect sizes are not specified in the file, they will be generated from a standard normal distribution.

mycausal.list

```
cg04584301 1.55182
cg04839274 -0.106226
cg16648571 0.0257417
```

This is a text file with no headers. Columns are probe ID and effect size.

```
osca --simu-qt --simu-hsq 0.1 --simu-eff-mod 0 --befile myprofile --simu-causal-loci mycausal.list --out mypheno
```

--simu-eff-mod specifies whether or not to standardize the causal probe, 0 for standardized profile and 1 for raw profile. The default value is 0.

```
osca --simu-cc 100 300 --simu-hsq 0.1 --simu-k 0.1 --befile myprofile --simu-causal-loci mycausal.list --out mypheno
```

- --simu-cc simulates a case-control trait and specifies the number of cases and the number of controls.
- --simu-k specifies the disease prevalence. The default value is 0.1 if this option is not specified.

Prediction Analysis

```
osca --reml --orm myorm --pheno my.phen --reml-pred-rand --out myblp
```

--reml-pred-rand predicts the random effects by the BLUP (best linear unbiased prediction) method. This option is to estimate the aggregated effect of all the probes (used to compute the ORM) to the phenotype of an individual. The aggregated omics effects of all the individuals will be saved in a plain text file *.indi.blp.

myblp.indi.blp

This is a text file with no headers. Columns are family ID, individual ID, an intermediate variable, the aggregated omics effect, another intermediate variable and the residual effect.

```
osca --befile myprofile --blup-probe myblp.indi.blp --out myblp
```

--blup-probe calculates the BLUP solutions for the probe effects.

myblp.probe.blp

```
cg04584301 -0.000654646
cg04839274 0.000602484
cg16648571 -4.70356e-05
...
```

This is a text file with no headers. Columns are probe ID and BLUP of the probe effect.

```
osca --befile myprofile --score myblp.probe.blp --out myscore

osca --befile myprofile --score myblp.probe.blp 1 2 --out myscore
```

--score reads score files for probes and generates predicted omics profiles for individuals. (Note that this option largely follows the --score option in PLINK.) It allows users to specify the column numbers for probe ID and score (the default values are 1 and 2 as shown in the example above).

myscore.profile

```
FID IID PHENO CNT SCORE

131000028 422572 -9 20000 -7.269198e-07

131000031 243421 -9 20000 9.322096e-06

131000179 338728 -9 20000 -1.250443e-05
...
```

This is a text file with headers. Columns are family ID, individual ID, phenotype, Number of non-missing probes and score.

osca --befile myprofile --score myblp.probe.blp --score-has-header --out myscore

--score-has-header indicates probe score file has headers.

eQTL/mQTL Analysis

basic option

```
osca --eqtl --bfile mydata --befile myprofile --out myeqtl
```

- --eqtl turns on eQTL analysis.
- --bfile reads individual-level SNP genotype data (in PLINK binary format) from a reference sample for LD estimation, i.e. .bed, .bim, and .fam files.

compute with multi-tasks

```
osca --eqtl --bfile mydata --befile myprofile --task-total 1000 --task-id 1 --thread-num 10 --out myeqtl
```

- --task-total specifies the total task number to patition computation by probes.
- --task-id specifies the current task ID which ranges form 1 to the total task number.
- --thread-num specifies the number of OpenMP threads for parallel computing. The default value is 1.

include covariates

```
osca --eqtl --bfile mydata --befile myprofile --covar mycovar --qcovar myqcovar --task-total 1000 --task-id 1 --thread-num 10 --out myeqtl
```

include data management options

```
osca — eqtl — bfile mydata — befile myprofile — extract-snp mysnp.list — maf 0.01 — task-total 1000 — task-id 1 — thread-num 10 — out myeqtl
```

--maf removes SNPs based on a minor allele frequency (MAF) threshold in the individual-level SNP genotype data.

save result in BESD dense format 2

The options above suffer from repeated recoding the genotype. This could be aggravated as the probe number increase. This option could alleviate the situation a little bit.

```
osca --eqtl --bfile mydata --befile myprofile --besd-snp-major --task-total 1000 --task-id 1 --thread-num 10 --out myeqtl
```

--besd-snp-major conducts eQTL analysis SNP by SNP across all the probes and saves the result in BESD dense format 2.

fast eQTL analysis

The options above fit the covariates in the model as $y=xb+C\beta+e$. It is equivalent to $y=y-C\beta^{\hat{}}$, $y=x-C\beta^{\hat{}}$, $y=x-C\beta$

```
osca --eqtl --bfile mydata --befile myprofile --fast-eqtl --task-total 1000 --task-id 1 --thread-num 10 --out myeqtl
```

--fast-eqtl runs fast eQTL analysis.

vQTL Analysis

basic option for variance QTL analysis

```
osca —vqtl —bfile mydata —befile myprofile —out myvqtl
```

--vqtl turns on vQTL analysis.

compute with multi-tasks

```
osca —-vqtl —-bfile mydata —-befile myprofile —-task-total 1000 —-task-id 1 —-thread-num 10 —-out myvqtl
```

specify the algorithm to analyse the homogeneity of variance

We provide 4 methods to run vQTL analysis. 0 for Bartlett's test; 1 for Levene's test with mean; 2 for Levene's test with median; 3 for Fligner-Killeen test.

```
osca — eqtl — bfile mydata — befile myprofile — vqtl—mtd 0 — task—total 1000 — task—id 1 — thread—num 10 — out myvqtl
```

--vqtl-mtd specifies the algorithm to analyse the homogeneity of variance. The default option is 0.

include data management options

```
osca --vqtl --bfile mydata --befile myprofile --extract-snp mysnp.list --maf 0.01 --task-total 1000 --task-id 1 --thread-num 10 --out myvqtl
```

save result in BESD dense format 2

```
osca --vqtl --bfile mydata --befile myprofile --besd-snp-major --task-total 1000 --task-id 1 --thread-num 10 --out myvqtl
```

vQTL analysis in cis-regions

Conduct the vQTL analysis only in cis-regions. The result will be saved in BESD sparse format 2.

```
osca --vqtl --bfile mydata --befile myprofile --cis-wind 2000 --task-total 1000 --task-id 1 --thread-num 10 --out myeqtl
```

--cis-wind defines a window centred around the probe to select SNPs for the vQTL analysis. The default value is 2000Kb.

Meta Analysis

Meta (meta analysis in unrelated samples)

meta for GWAS summary data

```
osca ——gwas—flist mygwas.flist ——meta ——out mymeta
```

--meta implements the conventional inverse-variance-weighted meta-analysis assuming all the cohorts are independent. Pleae refer to de Bakker PI et al. 2008 Hum Mol Genet for the details.

--gwas-flist reads a file to get file paths of the GWAS summary data.

mygwas.flist

```
Height.01.COJO
Height.02.COJO
Height.03.COJO
...
```

This file has no headers.

The input format of the GWAS summary data follows that for GCTA-COJO analysis (http://cnsgenomics.com/software/gcta/#COJO).

Height.01.COJO

```
SNP A1 A2 freq b se p n
rs1001 A G 0.8493 0.0024 0.0055 0.6653 129850
rs1002 C G 0.03606 0.0034 0.0115 0.7659 129799
rs1003 A C 0.5128 0.045 0.038 0.2319 129830
.....
```

Columns are SNP, the effect (coded) allele, the other allele, frequency of the effect allele, effect size, standard error, p-value and sample size. The headers are not keywords and will be omitted by the program. Important: "A1" needs to be the effect allele with "A2" being the other allele and "freq" needs to be the frequency of "A1". NOTE: For a case-control study, the effect size should be log(odds ratio) with its corresponding standard error.

meta for eQTL summary data

```
osca --besd-flist mybesd.flist --meta --out mymeta
```

--besd-flist reads a file to get the full paths of the BESD files.

mybesd.flist

```
path1/my besd1
path2/my besd2
path3/my_besd3
```

This file has no headers. The eQTL summary data should be in BESD formta.

MeCS (meta analysis in correlated sampels)

MeCS for GWAS summary data

```
osca --gwas-flist mygwas.flist --mecs --out mymecs
```

--mecs implements the MeCS analysis.

specify a method to estimate the cohort correlation matrix.

```
osca --gwas-flist mygwas.flist --mecs --mecs-mth 0 --out mymecs
```

--mecs-mth specifies a method to estimate the cohort correlation matrix. 0 for computing the Pearson Correlaton Coefficient using the beta values and 1 for estimating the cohort correlation using the beta values of pairwised common SNPs. The default value is 0.

use z-score to estimate the cohort correlation matrix.

```
osca --qwas-flist mygwas.flist --mecs --mecs-mth 0 --pcc-z --out mymecs
```

--pcc-z indicates using z-score to compute the Pearson Correlaton Coefficiet matrix. The default value is false.

specify a p-value threshold to exclude the significant SNPs from calculating the cohort correlation matrix.

```
osca --gwas-flist mygwas.flist --mecs --mecs-mth 0 --pmecs 0.01 --out mymecs
```

--pmecs reads a p-value threshold to exclude the significant SNPs from calculating the cohort correlation matrix. The default value is 0.01.

input the cohort correlation matrix

```
osca --qwas-flist mygwas.flist --mecs --cor-mat cor.mat --out mymecs
```

-cor-mat reads a p-value threshold to exclude the significant SNPs from calculation the cohort correlation matrix. The default value is 0.01.

mybesd.flist

```
1.000000000 0.03911566 0.001821693 0.021464142 0.017527909
0.039115655 1.00000000 0.010180606 0.016671469 0.017503133
0.001821693 0.01018061 1.000000000 0.001058645 0.004646767
0.021464142 0.01667147 0.001058645 1.000000000 0.034315463
0.017527909 0.01750313 0.004646767 0.034315463 1.000000000
```

This file is a symmetric matrix without header.

MeCS for eQTL summary data

MeCS is a method that only requires summary-level cis-eQTL data to perform a meta-analysis of cis-eQTLs from multiple cohorts (or tissues) with sample overlaps. It estimates the proportion of sample overlap from null SNPs in the cis-regions and meta-analysed the eQTL effects using a generalized least squares approach. The method can be applied to data from genetic studies of molecular phenotypes (e.g. DNA methylation and histone modification).

NOTE: Only the information in the cis-region would be used.

```
osca --besd-flist mybesd.flist --mecs --out mymecs
```

Options Reference

--blup-probe --chr

--covar

```
--befile
                         reads a DNA methylation (or gene expression) data file in binary format
--befile-flist
```

reads a file to get the full paths of the binary files --beta2m

calculates the methylation m value from the methylation beta value

calculates the BLUP solutions for the probe effects

specifies a chromosome to select probes

reads discrete covariates from a plain text file

reads a file that contains DNA methylation detection p-values --detection-pval-file --dpval-mth specifies a method to do quality control with the detection p-values

--dpval-thresh specifies a threshold of detection p-value

reads a DNA methylation (or gene expression) data file in plain text format --efile

--exclude-probe excludes a subset of probes --extract-probe extracts a subset of probes --from-probe specifies the start probe

--from-probe-kb specifies the start physical position of the probes

--gene extracts a gene

--genes extracts a subset of genes

--gene-expression indicates gene expression profiles in the file --get-mean calculates the mean of each probe --get-variance calculates the variance of each probe --keep extracts a subset of individuals

--linear saves linear regression statistics to a plain text file

--lower-beta removes the DNA methylation probes with the mean beta value smaller than a specified threshold

specifies a percentage of probes to exclude from calculating the ORM --lxpo calculates the methylation beta value from the methylation m value --m2beta saves DNA methylation (or gene expression) data in binary format --make-bod --make-efile saves the DNA methylation (or gene expression) data in text format

estimates the omics relationship matrix (ORM) and save the lower triangle elements of ORM to binary files --make-orm --make-orm-bin estimates the omics relationship matrix (ORM) and save the lower triangle elements of ORM to binary files

estimates the omics relationship matrix (ORM) and save the lower triangle elements of ORM to compressed plain text files --make-orm-gz --make-tefile saves the DNA methylation (or gene expression) data in transposed text format

--merge-orm reads multiple ORMs in binary format --methylation-m indicates methylation m values in the file indicates methylation beta values in the file --methylation-beta

initiates an MLM based association analysis including the target probe (the probe to be tested for association) in the ORM --mlma --mlma-loco initiates an MLM based association analysis with the chromosome where the target probe is located excluded from the ORM

--mpheno reads a list of comma-delimited trait numbers if the phenotype file contains more than one trait --missing-ratio-probe specifies a missing proportion threshold to remove probes

--no-fid indicates data without family ID --orm reads the ORM binary files

--orm-alg specifies the algorithm to estimate the ORM

--orm-cutoff removes one of a pair of individuals with estimated omics relationships larger than the specified cut-off value

saves data (or results) in a file --out

conducts principal component analysis and saves the first n (default as 20) PCs --pca

reads phenotype data from a plain text file --pheno extracts a specified probe --probe

--probe-wind defines a window centred on a specified probe

--probe-rm excludes a specified probe

--qcovar reads quantitative covariates from a plain text file --ratio-probe specifies a proportion threshold to remove probes --ratio-sample specifies a proportion threshold to remove individuals --reml performs REML (restricted maximum likelihood) analysis specifies the algorithm to do REML iterations --reml-alg --reml-est-fix displays the estimates of fixed effects on the screen --reml-maxit specifies the maximum number of iterations

--reml-no-lrt

--reml-pred-rand predicts the random effects by the BLUP (best linear unbiased prediction) method

--remove excludes a subset of individuals

--score reads score files for probes and generates predicted omics profiles for individuals

--score-has-header indicates probe score file has headers --simu-causal-loci reads a list of probes as causal probes

simulates a case-control trait and specifies the number of cases and the number of controls --simu-cc

--simu-eff-mod specifies whether or not to standardize the causal probe

--simu-hsq specifies the proportion of variance in phenotype explained by the causal probes specifies the disease prevalence. The default value is 0.1 if this option is not specified --simu-k --simu-qt simulates a quantitative trait

--std removes the probes with the standard deviation smaller than a specified threshold

--to-probe specifies the end probe

--to-probe-kb specifies the end physical position of the probes

--update-opi reads a fully annotated .opi file

--upper-beta removes the DNA methylation probes with the mean beta value larger than a specified threshold