# GTEx summary statistics database

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Last updated: July 7, 2015

## 1 Databases

# 1.1 Multi-tissue summary stats

Summary statistics from the GTEx v6 analysis are stored in MatrixEQTLSumStats.h5 (130GB, in HDF5 format). It contains 38,933 genes (less than previous version! See the list /project/mstephens/datal each gene has several summary statistics matrix of dimension  $N_{cisSNPs} \times N_{tissues}$ . For this release  $N_{tissues} = 44$ . For MatrixEQTL the summary statistics are  $\hat{\beta}$ , T-stat and P-value.

```
MatrixEQTLSumStats.h5

/ENSG00000008735.10 'MatrixEQTL summary statistics of GTEx Release 2015.04.15'
/ENSG00000008735.10/beta (5636, 44)
/ENSG00000008735.10/p-value (5636, 44)
/ENSG00000008735.10/t-stat (5636, 44)
/ENSG00000008735.10/rownames (5636,)
/ENSG00000008735.10/colnames (44,)
...
```

There are two special tables in MatrixEQTLSumStats.h5. One called max, which is the data for the "best" gene-snp pair identified per gene as the snp pair having maximum abs(t-stat) across all snps and all tissues and there is no missing data for any tissue. It contains 16,069 gene-snp pairs. The other called null, which organize gene-snp pair data in the same format as max only that the gene-snp pairs in this table are NOT the best gene-snp pair, but just random samples from all non-best gene-snp pairs. For portability these two tables are also stored separately in MatrixEQTLSumStats.Portable.h5 (58MB, in HDF5 format).

#### 1.2 Meta-database

Unfortunately GTEx names cisSNP by their genomic coordinate in Human Genome Assembly hg19 (b37), not by rsID. To make it possible to search with rsID I created another database snp-gene.db (1.6GB, in SQLite format).

```
snp-gene.db
3_52771872_C_T_b37
                                        ENSG00000016864.12, ENSG00000055955.11, ENSG00000055957.6 ...
3_52774530_C_T_b37
                                        ENSG00000016864.12,ENSG00000055955.11,ENSG00000055957.6 ...
                        rs2159644
3_52774835_AT_A_b37
                        rs35270761.rs397785457 ENSG00000016864.12.ENSG00000055955.11.ENSG00000055957.6 ...
```

### The 3 columns are

- SNP name in GTEx convention
- rsID
- cis-genes, i.e., genes whose TSS is within 100,000bp up/down-stream of the SNP.

This database will be used prior to searching the summary statistics.



# **Note**

- There may be multiple rsID associated with the same coordinate. Indeed such records exists in dbSNP 144, and eventually these rsIDs should merge into one name (as always have happened in history).
- There are 10,297,646 SNPs in total in this dataset. 9,794,339 of them (95.1%) has rsID in dbSNP 144.

# **Queries**

I provide a simple R script SumstatQuery. R to make queries on this dataset. It has:

- A function that lists for given rsID the GTEx SNP ID and all its cis-genes.
- A function that loads sumstats data given gene name and GTEx SNP ID.

To use the script, RSQLite and rhdf5 should be installed, e.g.,

```
install.packages("RSQLite")
source("http://bioconductor.org/biocLite.R")
biocLite("rhdf5")
```

# 2.1 Example query

The demo script below shows an example to extract information for gene ENSG00000171960.6 and SNP rs6600419, and examples to extract the best/null gene-snp pairs.

```
R
source("/project/mstephens/gtex/scripts/SumstatQuery.R")
# Load data for given gene
dat <- GetSS("ENSG00000171960.6", "/project/mstephens/gtex/analysis/april2015/query/MatrixEQTLSumStats.h5")
print(names(dat))
## [1] "beta"
                               "p-value" "t-stat" "z-score"
dim(dat$"beta")
## [1] 6344 44
dat$"beta"[1:4,1:4]
                                        Adipose_Subcutaneous Adipose_Visceral_Omentum Adrenal_Gland Artery_Aorta
## 1 \212\161 C T b37
                                                                                                           NaN NaN 0.06785597
                                                       0.04608095
## 1_42124246_A_AC_b37
                                                        -0.03087348
                                                                                                     -0.03222961 -0.1286249 -0.13490575
## 1_42124311_G_A_b37
                                                                                                                               NaN -0.21858459
                                                         0.12400229
                                                                                                      0.08011222
                                                       0.04630340
## 1_42124614_C_T_b37
                                                                                                                                           NaN 0.06785597
                                                                                                                 NaN
# Output information for the SNP of interest
dat$"p-value"["1_43124701_A_G_b37", ]
                                  Adipose\_Subcutaneous
                                                                                                Adipose_Visceral_Omentum
 ##
                                                 1.632499e-01
                                                                                                                    3.462833e-02
  ##
                                                Adrenal\_Gland
                                                                                                                      Artery_Aorta
  ##
                                                  8.168959e-01
                                                                                                                      5.600338e-01
                                                                                                                    Artery_Tibial
  ##
                                             Artery Coronary
  ##
                                                 7.659/26e-01
                                                                                                                      7.397466e-01
  {\it \#\# Brain\_Anterior\_cingulate\_cortex\_BA24}
                                                                                          Brain\_Caudate\_basal\_ganglia
  ## ...
dat$"z-score"["1_43124701_A_G_b37", ]
                                                                                                Adipose_Visceral_Omentum
                                  Adipose_Subcutaneous
                                                                                                                       -2.11267803
                                                   -1.39422418
                                                Adrenal\_Gland
  ##
                                                                                                                      Artery_Aorta
  ##
                                                   -0.23153929
                                                                                                                         0.58279135
                                             Artery_Coronary
                                                                                                                     Artery\_Tibial
                                                    0.29768626
  ##
                                                                                                                         0.33218889
  ## Brain_Anterior_cingulate_cortex_BA24
                                                                                       Brain_Caudate_basal_ganglia
###
# Load the best gene-snp data
mdat <- GetSS("max", "/project/mstephens/gtex/analysis/april2015/query/MatrixEQTLSumStats.h5")</pre>
dim(mdat$"t-stat")
mdat$"p-value"[1:4,1:4]
mdat$"t-stat"["ENSG00000000419.8_20_49461813_G_C_b37",]
## Is this gene-snp pair most significant in spleen?
dat <- GetSS("ENSG00000000419.8", "/project/mstephens/gtex/analysis/april2015/query/MatrixEQTLSumStats.h5")
idx.to.show <- matxMax(abs(dat$"t-stat"))
rownames(dat$"t-stat")[idx.to.show[1]]
colnames(dat$"t-stat")[idx.to.show[2]]
###
# Load the "null" gene-snp data
ndat <- GetSS("null", "/project/mstephens/gtex/analysis/april2015/query/MatrixEQTLSumStats.h5")</pre>
dim(ndat$"t-stat")
# Look up GTEx SNP ID
ShowSNP("rs6600419", "/project/mstephens/gtex/analysis/april2015/query/snp-gene.db")
## GTEx SNP ID: 1_43124701_A_G_b37
\textit{\#\# cisGenes: ENSG0000065978.13, ENSG00000164007.6, ENSG00000171960.6, ENSG000002364.1, ENSG00000234917.1, ENSG00000236180.2}
# Look up rs ID given GTEx SNP ID
Show SNP ("1\_43124701\_A\_G\_b37", "/project/mstephens/gtex/analysis/april2015/query/snp-gene.db") = (12.43124701\_A\_G\_b37", "/project/mstephens/gtex/analysis/april2015/query/snp-gene.db") = (12.43124701\_A_G\_b37", "/project/mstephens/gtex/april2015/query/snp-gene/gtex/april2015/query/snp-gene/gtex/april2015/query/snp-gene/gtex/april2015/query/snp-gene/gte
## rsID(s): rs6600419
\textit{\#\# cisGenes: ENSG00000065978.13, ENSG00000164007.6, ENSG00000171960.6, ENSG00000230254.1, ENSG00000234917.1, ENSG00000236180.2}
##
# Create matched training/testing sets
##
```

```
N1 <- 8000
N2 <- 16069
strong.train <- SubsetMatLists(mdat, seq(1, N1))</pre>
strong.test <- SubsetMatLists(mdat, seq(N1 + 1, N2))</pre>
strong.train.genes <- as.character(lapply(strsplit(rownames(strong.train$beta), "_"), function(x) x[1]))
strong.test.genes <- as.character(lapply(strsplit(rownames(strong.test$beta), "_"), function(x) x[1]))
null.genes <- as.character(lapply(strsplit(rownames(ndat$beta), "_"), function(x) x[1]))</pre>
null.train <- SubsetMatLists(ndat, which(null.genes %in% strong.train.genes))</pre>
null.test <- SubsetMatLists(ndat, which(null.genes %in% strong.test.genes))</pre>
```

For MatrixEQTLSumStats.h5, for given rsID you first search the GTEx SNP ID via ShowSNP, then load the cisGene of interest via GetSS and extract information using the GTEx SNP ID as the row name key.



If R complains the given row name does not exist, try to flip the DNA strand in the GTEx SNP ID and search again. For example, instead of searching for 1\_43124701\_A\_G\_b37 you search for 1\_43124701\_T\_C\_b37.

The best gene-snp data can be loaded to memory, via GetSS("max", "MatrixEQTLSumStats.h5"), or the portable version GetSS("max", "MatrixEQTLSumStats.Portable.h5")

Same for the "null" gene-snp data, via GetSS("null", "MatrixEQTLSumStats.h5"), or the portable version GetSS("null", "MatrixEQTLSumStats.Portable.h5")