"A simple testing procedure for multiseq"

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1 Overview

- After making changes in a software package multiseq, we run a set of tests to check if the changes in multiseq affect output as we planned (e.g., changes shouldn't affect other outputs).
- We simulate four sets of data typical data and data with no reads with/without library read depth.

2 Simulation of data set

2.1 Collect information to simulate data

I'll copy phenotype data (from which we will sample data) and signals for group 1 and group 2. See compareWaveQTLandmultiseq_dsQTL_final.org for detailed description of how to obtain this information.

```
cd ~/multiseq/
mkdir data
```

```
cd data/
mkdir simulation
cd simulation/
mkdir dsQTL
cp ~/multiscale_analysis/analysis/simulation/sample_size/simulation_QTLfinal_v2
    /data/pheno.dat dsQTL
cp ~/multiscale_analysis/analysis/simulation/sample_size/simulation_QTLfinal_v2
    /data/alt.sig0 dsQTL
cp ~/multiscale_analysis/analysis/simulation/sample_size/simulation_QTLfinal_v2
    /data/alt.sig1 dsQTL
```

I'll copy library read depth for this DNase-seq data (you can download from WaveQTL repo) to this directory.

```
cp ~/WaveQTL/data/dsQTL/library.read.depth.dat ~/multiseq/data/simulation/dsQTL
/
```

Information have been save in: #+besin_src bash cd ~/multiseq/data/simulation/dsQTL alt.sig0 alt.sig1 library.read.depth.dat pheno.dat #+end src

2.2 Simulate data

The function 'simulate.data' is a modification of the script in multiscale_analysis repo. This script uses functions in 'my.utils.R' in multiscale_analysis repo.

```
setwd("~/multiseq/data/simulation/dsQTL/")
source("~/multiscale_analysis/src/R/my.utils.R")
##' 'simulate.data' simulate data sets by resampling reads from real data (real .read.counts) for given signals (sig0, sig1).
##'
##' @param seed seed number to set up
##' @param numGroupO number of samples for GroupO
##' @param numGroup1 number of samples for Group1
##' @param real.read.counts a vector of size T (e.g., 1024); t-th element contains number of reads at t-th position; from which we will resample reads for simulation;
##' @param sigO a vector of size T (e.g., 1024); t-th element contains probability of sampling read at t-th position in real.read.counts for GroupO
```

```
##' Oparam sig1 a vector of size T (e.g., 1024); t-th element contains
   probability of sampling read at t-th position in real.read.counts for
##' Oparam real.library.read.depth a vector of size M (>1); from whch we will
   sample library read depth
##' @param over.dispersion parameter used in sample.from.Binomial.with.
   Overdispersion; see 'my.utils.R' for details.
##' @return a list of data, group, library.read.depth; data contains simulated
   data; matrix of (numGroup0+numGroup1) by T; group contains group indicator
   for simulated data; a vector of size (numGroup0+numGroup1); library.read.
   depth contains simulated library read depth; a vector of size (numGroup0+
   numGroup1)
simulate.data <- function(seed = 1, numGroup0, numGroup1, sig0, sig1, real.read</pre>
   .counts, real.library.read.depth = NULL, over.dispersion=NULL){
 genoD = c(rep(0, numGroup0), rep(1, numGroup1))
 numSam = length(genoD)
 numBPs = length(sig0)
 ## phenotype data
 phenoD = matrix(data=NA, nr= length(genoD), nc = numBPs)
 ## let's sample!!!
 set.seed(seed)
 ## upper and lower bound!
 trunc.fun = function(x){
   x = max(0, x)
   return(min(1,x))
 p.sig0 = sapply(sig0, trunc.fun)
 p.sig1 = sapply(sig1, trunc.fun)
 ## geno = 0
 wh0 = which(genoD == 0)
 if(length(wh0) > 0){
   phenoD[wh0,] = sample.from.Binomial.with.Overdispersion(num.sam = length(
       wh0), total.count = real.read.counts, mu.sig = p.sig0, over.dispersion =
        over.dispersion)
 }
```

```
## geno = 1
 wh1 = which(genoD == 1)
 if(length(wh1) > 0){
   phenoD[wh1,] = sample.from.Binomial.with.Overdispersion(num.sam = length(
       wh1), total.count = real.read.counts, mu.sig = p.sig1, over.dispersion =
        over.dispersion)
 if(is.null(real.library.read.depth)){
   library.read.depth=NULL
 }else{
   library.read.depth = sample(real.library.read.depth, numSam, replace = TRUE
 return(list(data = phenoD, group = genoD, library.read.depth = library.read.
     depth))
}
seed = 1
numGroup0 = 10
numGroup1 = 10
sig0 = scan("~/multiseq/data/simulation/dsQTL/alt.sig0", what=double())
sig1 = scan("~/multiseq/data/simulation/dsQTL/alt.sig1", what=double())
real.DNase.dat = read.table("~/multiseq/data/simulation/dsQTL/pheno.dat", as.is
    = TRUE)
real.read.counts = ceiling(as.numeric(apply(real.DNase.dat, 2, sum)))
real.library.read.depth = scan("~/multiseq/data/simulation/dsQTL/library.read.
   depth.dat", what=double())
## data with sample size 20 with library read depth
res = simulate.data(seed = seed, numGroup0 = numGroup0, numGroup1 = numGroup1,
   sig0 = sig0, sig1= sig1, real.read.counts = real.read.counts, real.library.
   read.depth = real.library.read.depth, over.dispersion=NULL)
str(res)
## List of 3
## $ data : int [1:20, 1:1024] 0 0 0 0 0 0 0 0 0 0 ...
## $ group : num [1:20] 0 0 0 0 0 0 0 0 0 ...
## $ library.read.depth: num [1:20] 44257311 37331440 30843655 36823292
   50966695 ...
apply(res$data,1,sum)
## [1] 65 60 63 62 81 75 57 65 57 66 39 40 49 46 44 48 53 43 37 51
```

```
## data with sample size 20 with library read depth, but all samples in group 2
    have zero read count.
res = simulate.data(seed = seed, numGroup0 = numGroup0, numGroup1 = numGroup1,
   sig0 = sig0, sig1= rep(0, length(sig1)), real.read.counts = real.read.
   counts, real.library.read.depth = real.library.read.depth, over.dispersion=
   NULL)
apply(res$data,1, sum)
## [1] 65 60 63 62 81 75 57 65 57 66 0 0 0 0 0 0 0 0 0
## data with sample size 20 without library read depth
res = simulate.data(seed = seed, numGroup0 = numGroup0, numGroup1 = numGroup1,
   sig0 = sig0, sig1= sig1, real.read.counts = real.read.counts, over.
   dispersion=NULL)
apply(res$data,1,sum)
## [1] 65 60 63 62 81 75 57 65 57 66 39 40 49 46 44 48 53 43 37 51
res$library.read.depth
## NULL
## data with sample size 20 without library read depth, but all samples in
   group 2 have zero read count.
res = simulate.data(seed = seed, numGroup0 = numGroup0, numGroup1 = numGroup1,
   sig0 = sig0, sig1= rep(0, length(sig1)), real.read.counts = real.read.
   counts, over.dispersion=NULL)
apply(res$data,1,sum)
## [1] 65 60 63 62 81 75 57 65 57 66 0 0 0 0 0 0 0 0 0
res$library.read.depth
##NULL
```