ELMER: An R/Bioconductor Tool Inferring Regulatory Element Landscapes and Transcription Factor Networks Using Methylomes

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1 Introduction

This document provides an introduction of the *ELMER*, which is designed to use DNA methylation and gene expression data sets from a large number of tissue samples to infer regulatory element landscapes and transcription factor network. It includes functions for identifying probes at distal regulatory regions with differential DNA methylation levels, predicting genes whose expression associates with the differentially methylated probes and discovering the functional regulatory TFs. This package can be easily applied to TCGA public available cancer data sets and to custom DNA methylation and gene expression data sets.

1.1 Installing and loading ELMER

To obtain a copy of ELMER, you will need to install devtools

```
install.packages(devtools)
library(devtools);
devtools::install_github("lijingya/ELMER");
```

2 Download example data

The following steps can be used to download the example data set for ELMER

```
## Loading required package: GenomicRanges
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
##
## The following objects are masked from 'package:parallel':
##
##
      clusterApply, clusterApplyLB, clusterCall, clusterEvalQ, clusterExport,
      cluster {\tt Map}, \ par {\tt Apply}, \ par {\tt Capply}, \ par {\tt Lapply}, \ par {\tt Lapply} {\tt LB}, \ par {\tt Rapply},
##
##
      parSapply, parSapplyLB
##
## The following object is masked from 'package:stats':
##
##
      xtabs
##
## The following objects are masked from 'package:base':
##
##
      anyDuplicated, append, as.data.frame, as.vector, cbind, colnames, do.call,
##
      duplicated, eval, evalq, Filter, Find, get, intersect, is unsorted, lapply,
##
      Map, mapply, match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,
##
      Position, rank, rbind, Reduce, rep.int, rownames, sapply, setdiff, sort,
##
      table, tapply, union, unique, unlist, unsplit
##
## Loading required package: S4Vectors
## Loading required package:
                               stats4
## Loading required package:
                               IRanges
## Loading required package: GenomeInfoDb
```

3 Quick start: running TCGA example

Function, TCGA.pipe, is the easy usage for downloading TCGA data and performing all the analyses in ELMER. For illustration purpose, we skip the downloading step. The user can use the getTCGA function to download TCGA data or use TCGA.pipe by including "download" in the analysis option.

```
TCGA.pipe("LUSC",wd="./ELMER.example",cores=detectCores()/2,permu.size=300,
          analysis = c("distal.enhancer", "diffMeth", "pair", "motif", "TF.search"),
          diff.dir="hypo",rm.chr=paste0("chr",c(1:22,"X","Y")))
## ##################
## Select distal enhancer probes
## ##################
## Warning in (function (probe, distal = TRUE, feature, TSS, TSS.range = list(upstream = 2000, :
Default probes coordinates are for HM450K DNA methylation array
## ##################
## Get differential DNA methylation loci
## ##################
##
##
## ~~~ MEE.data: initializator ~~~
## ##################
## Predict pairs
## ##################
##
##
## ~~~ MEE.data: initializator ~~~
## Identify putative probe-gene pair for hypomethylated probes
## Calculate empirical P value.
##
## ##################
## Motif search
## ##################
##
##
## Identify enriched motif for hypomethylated probes
## 12 motifs are enriched.
## #################
## Search responsible TFs
## ##################
##
##
## ~~~ MEE.data: initializator ~~~
## Identify regulatory TF for enriched motif in hypomethylated probes
```

4 Input data

The whole pipeline analyses in *ELMER* needs at least 4 input files: a matrix of DNA methylation from HM450K platform; a matrix of gene expression for the same samples; a GRanges object containing the information for probes on HM450K such as names and coordinates; a gene annotation which is also a GRanges object. When TCGA data are used, the sample information will be automatically generated by fetch.mee function. However sample information should be provided when using custom data.

4.1 DNA methylation data

Raw DNA methylation data can be processed by *Methylumi* or *minfi* generating DNA methylation information for each CpG. The DNA methylation level at each CpG is referred to as a beta (β) value, calculated as (M/(M+U)), where M represents the methylated allele intensity and U the unmethylated allele intensity. Beta values range from 0 to 1, reflecting the fraction of methylated alleles at each CpG in the each tumor; beta values close to 0 indicates low levels of DNA methylation and beta values close to 1 indicates high levels of DNA methylation. Generate a matrix with DNA methylation beta values for all the samples (columns) and probe loci (rows) and save matrix as meth.rda

```
load("./ELMER.example/Result/LUSC/LUSC_meth_refined.rda")
Meth[1:10, 1:2]
##
              TCGA-43-3394-11A-01D-1551-05 TCGA-43-3920-11B-01D-1551-05
## cg00045114
                                  0.8190894
                                                                0.8073763
## cg00050294
                                  0.8423084
                                                                0.8241138
## cg00066722
                                  0.9101127
                                                                0.9162212
## cg00093522
                                  0.8751903
                                                                0.8864599
## cg00107046
                                  0.3326016
                                                                0.3445508
## cg00116430
                                  0.6097183
                                                                0.5952469
## cg00152117
                                  0.7074149
                                                                0.6439695
## cg00163018
                                  0.5928909
                                                                0.8250584
## cg00173804
                                  0.9162264
                                                                0.9303684
## cg00223046
                                  0.7826863
                                                                0.7744760
```

4.2 Gene expression data

Gene expression values can be generated from different platforms such as array or RNA-seq, gene level or transcript level gene expression calling. Generate a matrix with gene expression values for all the samples (columns) and genes (rows) and save matrix as RNA.rda

```
load("./ELMER.example/Result/LUSC/LUSC_RNA_refined.rda")
GeneExp[1:10, 1:2]
##
            TCGA-22-5472-01A-01R-1635-07 TCGA-22-5489-01A-01R-1635-07
## ID126767
                                0.0000000
                                                               0.000000
## ID343066
                                0.4303923
                                                               0.000000
## ID26574
                               10.0817831
                                                               10.717673
## ID24
                                6.4462711
                                                                6.386644
## ID23456
                                8.5929182
                                                                9.333097
## ID5825
                               10.5578756
                                                               9.878333
## ID25
                               10.7233258
                                                              11.075515
## ID27
                                8.9761542
                                                                9.569239
## ID29777
                                9.6415206
                                                                9.353424
## ID80325
                                8.9840983
                                                                9.177624
```

4.3 Sample information

Sample information should be stored as a data.frame object containing sample ID, group labels (such as tumor, normal) and other description for each sample. When TCGA data were used, tumor, normal group label will be automatically generated by fetch.mee function by specifying option TCGA=TRUE.

```
mee <- fetch.mee(meth=Meth, exp=GeneExp, TCGA=T)</pre>
## ~~~ MEE.data: initializator ~~~
head(getSample(mee))
                                ID
                                                        meth.ID
                                                                                       exp.ID
## TCGA-43-3394-11 TCGA-43-3394-11 TCGA-43-3394-11A-01D-1551-05 TCGA-43-3394-11A-01R-1758-07
## TCGA-56-8305-01 TCGA-56-8305-01 TCGA-56-8305-01A-11D-2294-05 TCGA-56-8305-01A-11R-2296-07
## TCGA-56-8307-01 TCGA-56-8307-01 TCGA-56-8307-01A-11D-2294-05 TCGA-56-8307-01A-11R-2296-07
## TCGA-56-8308-01 TCGA-56-8308-01 TCGA-56-8308-01A-11D-2294-05 TCGA-56-8308-01A-11R-2296-07
## TCGA-56-8309-01 TCGA-56-8309-01 TCGA-56-8309-01A-11D-2294-05 TCGA-56-8309-01A-11R-2296-07
## TCGA-58-8386-01 TCGA-58-8386-01 TCGA-58-8386-01A-11D-2294-05 TCGA-58-8386-01A-11R-2296-07
##
## TCGA-43-3394-11 Normal
## TCGA-56-8305-01
## TCGA-56-8307-01 Tumor
## TCGA-56-8308-01 Tumor
## TCGA-56-8309-01 Tumor
## TCGA-58-8386-01 Tumor
```

4.4 Probe information

Probe information should be stored as a GRanges object containing the coordinate of each probe on the DNA methylation array and names of each probe. The default probe information is for HM450K.

```
probe <- ReadBed(system.file("extdata","Illumina-methyl-450K-manifest.hg19.bed.xz",</pre>
                             package = "ELMER"))
mee <- fetch.mee(meth=Meth, exp=GeneExp, TCGA=T, probeInfo=probe)</pre>
## ~~~ MEE.data: initializator ~~~
getProbeInfo(mee)
## GRanges object with 1725 ranges and 1 metadata column:
##
                                   ranges strand
            seqnames
                                                                   name
##
               <Rle>
                                  <IRanges> <Rle>
                                                            <character>
        [1]
                chr1 [172674159, 172674159]
                                                             cg00045114
##
        [2]
                chr1 [ 2886818, 2886818]
##
                                                             cg00050294
                chr1 [ 43634520, 43634520]
        [3]
##
                                                      cg00066722
##
        [4]
                chr1 [ 2252019, 2252019]
                                                             cg00093522
##
        [5]
                chr1 [ 16465562, 16465562]
                                                      cg00107046
##
                 . . .
                chr1 [215147891, 215147891]
                                                             cg27589988
##
     [1721]
                                                      ##
     [1722]
                chr1 [ 46632696, 46632696]
                                                      cg27636310
##
     [1723]
                chr1 [ 3472204, 3472204]
                                                             cg27637706
                chr1 [ 3283394,
##
     [1724]
                                  3283394]
                                                      ch.1.131529R
                chr1 [174947362, 174947362]
##
     [1725]
                                                      | ch.1.173213985R
##
     seqinfo: 24 sequences from an unspecified genome; no seqlengths
##
```

4.5 Gene information

Gene information should be stored as a GRanges object containing coordinates of each gene, gene id, gene symbol and gene isoform id. The default gene information is the UCSC gene annotation.

```
load(system.file("extdata","UCSC_gene_hg19.rda",package = "ELMER"))
## In TCGA expression data, geneIDs were used as the rowname for each row. However, numbers
## can't be the rownames, "ID" was added to each gene id functioning as the rowname.
## If your geneID is consistent with the rownames of the gene expression matrix, adding "ID"
## to each geneID can be skipped.
txs$GENEID <- pasteO("ID",txs$GENEID)</pre>
geneInfo <- promoters(txs,upstream = 0, downstream = 0)</pre>
save(geneInfo,file="./ELMER.example/Result/LUSC/geneAnnot.rda")
mee <- fetch.mee(meth=Meth, exp=GeneExp, TCGA=T, geneInfo=txs)</pre>
## ~~~ MEE.data: initializator ~~~
getGeneInfo(mee)
## GRanges object with 13741 ranges and 3 metadata columns:
##
          seqnames
                              ranges strand | tx_name
                                                               GENEID
                                                                               SYMBOL
                               <IRanges> <Rle> | <character> <character> <character>
##
             <Rle>
                                                | uc001hzz.1 ID10000
##
    10000
            chr1 [243651535, 244006584]
                                                                                AKT3
##
    10000
             chr1 [243663021, 244006584]
                                             - | uc001iab.2
                                                                  ID10000
                                                                                AKT3
                                            - | uc021plu.1
- | uc001xmf.3
- | uc010tth.2
             chr1 [243663021, 244006886]
##
    10000
                                                                  ID10000
                                                                                AKT3
    10001 chr14 [ 71050957, 71067384]
##
                                                                  ID10001
                                                                                MED6
##
    10001 chr14 [ 71050957, 71067384]
                                                                  ID10001
                                                                                MED6
             . . .
##
     . . .
                                                     . . .
                                                                   . . .
                                          + | uc003uik.3
           chr7
                    [86781677, 86825648]
                                                                  ID9988
##
     9988
                                                                                DMTF1
     9988 chr7 [86781677, 86825648]
                                                | uc011khb.2
##
                                             +
                                                                   ID9988
                                                                                DMTF1
                                            + | uc003uil.3
##
     9988 chr7 [86781870, 86825648]
                                                                  ID9988
                                                                                DMTF1
##
     9988 chr7 [86792198, 86809018]
                                            + | uc003uim.1
                                                                  ID9988
                                                                                DMTF1
     9988
              chr7 [86792198, 86825648]
                                           +
                                                uc003uin.3
##
                                                                   ID9988
                                                                                DMTF1
##
    seqinfo: 93 sequences (1 circular) from hg19 genome
```

4.6 MEE.data

The above 5 components will generate a MEE.data object as the main input for mulitple functions in ELMER.

```
mee <- fetch.mee(meth=Meth, exp=GeneExp, TCGA=T, probeInfo=probe, geneInfo=txs)

## ~~~ MEE.data: initializator ~~~

mee

## *** Class MEE.data, method show ***

## a meth

## num [1:1725, 1:234] 0.819 0.842 0.91 0.875 0.333 ...

## - attr(*, "dimnames")=List of 2

## ..$ : chr [1:1725] "cg00045114" "cg00050294" "cg00066722" "cg00093522" ...

## ..$ : chr [1:234] "TCGA-43-3394-11A-01D-1551-05" "TCGA-56-8305-01A-11D-2294-05" "TCGA-56-8307-01A-11D

## NULL

## * exp

## num [1:3894, 1:234] 0 0.214 10.048 5.007 8.63 ...

## - attr(*, "dimnames")=List of 2

## ..$ : chr [1:3894] "ID126767" "ID343066" "ID26574" "ID24" ...</pre>
```

```
## ..$: chr [1:234] "TCGA-43-3394-11A-01R-1758-07" "TCGA-56-8305-01A-11R-2296-07" "TCGA-56-8307-01A-11R
## NULL
## * sample
## 'data.frame': 234 obs. of 4 variables:
## $ ID : chr "TCGA-43-3394-11" "TCGA-56-8305-01" "TCGA-56-8307-01" "TCGA-56-8308-01" ...
## $ meth.ID: chr "TCGA-43-3394-11A-01D-1551-05" "TCGA-56-8305-01A-11D-2294-05" "TCGA-56-8307-01A-11D-22
## $ exp.ID : chr "TCGA-43-3394-11A-01R-1758-07" "TCGA-56-8305-01A-11R-2296-07" "TCGA-56-8307-01A-11R-22
               : chr "Normal" "Tumor" "Tumor" "Tumor" ...
## $ TN
## NULL
## * probeInfo
## GRanges object with 1725 ranges and 1 metadata column:

        seqnames
        ranges strand
        |
        name

        <Rle>
        <IRanges><Rle>
        <character>

##
                                                                             cg00045114
cg00050294
cg00066722
cg00093522
cg00107046
           [1] chr1 [172674159, 172674159] *
##
           [2] chr1 [ 2886818, 2886818]
##
                                                                *
          [3] chr1 [ 43634520, 43634520]
[4] chr1 [ 2252019, 2252019]
                                                                *
##
##
                                                                * |
          [5] chr1 [ 16465562, 16465562]
##
                                                                * |
##
## [1721] chr1 [215147891, 215147891] * |
## [1722] chr1 [ 46632696, 46632696] * |
## [1723] chr1 [ 3472204, 3472204] * |
                                                                             cg27589988
                                                                              cg27636310
                                                                               cg27637706
## [1724] chr1 [ 3283394, 3283394] * | ch.1.131529R
## [1725] chr1 [174947362, 174947362] * | ch.1.173213985R
##
      seqinfo: 24 sequences from an unspecified genome; no seqlengths
## * geneInfo
## GRanges object with 13741 ranges and 3 metadata columns:
## seqnames ranges strand | tx_name GENEID SYMBOL
                                          <IRanges> <Rle> | <character> <character> <character>
##
                 <Rle>
## 10000 chr1 [243651535, 244006584] - | uc001hzz.1 ID10000 AKT3
## 10000 chr1 [243663021, 244006584] - | uc001iab.2 ID10000 AKT3
## 10000 chr1 [243663021, 244006886] - | uc021plu.1 ID10000 AKT3
## 10001 chr14 [71050957, 71067384] - | uc001xmf.3 ID10001 MED6
## 10001 chr14 [71050957, 71067384] - | uc010tth.2 ID10001 MED6
## ... ... ... ... ... ... ... ... ...
     9988 chr7 [86781677, 86825648] + | uc003uik.3 ID9988

9988 chr7 [86781677, 86825648] + | uc011khb.2 ID9988

9988 chr7 [86781870, 86825648] + | uc003uil.3 ID9988

9988 chr7 [86792198, 86809018] + | uc003uim.1 ID9988

9988 chr7 [86792198, 86825648] + | uc003uim.3 ID9988
                                                                                                              DMTF1
##
##
                                                                                                               DMTF1
##
                                                                                                                DMTF1
##
                                                                                                                DMTF1
##
                                                                                                               DMTF1
##
      seqinfo: 93 sequences (1 circular) from hg19 genome
## ***** End Print (MEE.data) *****
```

5 Illustration of ELMER analysis steps

A subset of chromosome 1 data from TCGA LUSC were used as illustruation.

5.1 Selection of probes within biofeatures

Function, get.feature.probe, is used to select probes that are located within biofeatures such as H3K27ac ChIP-seq peaks. As default, the get.feature.probe function will automatically select distal enhancer probes on HM450K which are at least 2kb away from the TSS annotated by GENCODE V15 and UCSC-gene and locate within the putative comprehensive enhancers from REMC, ENCODE and FANTOM5.

```
#get distal enhancer probes that are 2kb away from TSS and overlap with REMC and FANTOM5
#enhancers on chromosome 1
Probe <- get.feature.probe(probe=probe, rm.chr=paste0("chr",c(2:22,"X","Y")))
save(Probe,file="./ELMER.example/Result/LUSC/probeInfo_feature.rda")</pre>
```

5.2 Identifying differentially methylated probes

Function, get.diff.meth, will be used to identify differentially methylated probes among the ones within biofeatures, which are selected in the above step.

```
## fetch.mee can take path as input.
mee <- fetch.mee(meth="./ELMER.example/Result/LUSC/LUSC_meth_refined.rda",</pre>
                  exp="./ELMER.example/Result/LUSC/LUSC_RNA_refined.rda", TCGA=T,
                  probeInfo="./ELMER.example/Result/LUSC/probeInfo_feature.rda",
                  geneInfo="./ELMER.example/Result/LUSC/geneAnnot.rda")
## ~~~ MEE.data: initializator ~~~
sig.diff <- get.diff.meth(mee, cores=detectCores()/2, dir.out ="./ELMER.example/Result/LUSC",</pre>
                           diff.dir="hypo", pvalue = 0.01)
sig.diff$hypo[1:10,]
                      ## significantly hypomethylated probes
                    probe
                                pvalue tumorMinNormal
                                                            adjust.p
## cg00045114 cg00045114 7.307478e-13
                                        -0.3499588 2.419463e-12
## cg00050294 cg00050294 4.440329e-09
                                            -0.5000920 1.057951e-08
## cg00093522 cg00093522 1.143338e-23
                                           -0.3596944 1.001147e-22
## cg00163018 cg00163018 2.240026e-21
                                            -0.3838195 1.558083e-20
## cg00173804 cg00173804 2.767232e-18
                                            -0.3744557 1.455328e-17
## cg00223046 cg00223046 8.142751e-11
                                            -0.3313773 2.317862e-10
## cg00255699 cg00255699 6.386989e-13
## cg00292636 cg00292636 6.368018e-34
## cg00329272 cg00329272 6.864176e-39
                                           -0.4308994 2.126941e-12
                                            -0.4833375 2.112468e-32
                                            -0.4355332 5.638431e-37
## cg00340127 cg00340127 3.056522e-18
                                            -0.5857140 1.602584e-17
# get.diff.meth automatically save output files.
# qetMethdiff.hypo.probes.csv contains statistics for all the probes.
# getMethdiff.hypo.probes.significant.csv contains only the significant probes.
dir(path = "./ELMER.example/Result/LUSC", pattern = "getMethdiff")
                                                    "getMethdiff.hypo.probes.significant.csv"
## [1] "getMethdiff.hypo.probes.csv"
```

5.3 Identifying putative probe-gene pairs

Function, get.pair function, will be used to identify putative target genes for selected probes. This step is the most time consuming step since it requires a large amount calculations for permutation. The greater the permutation time is, the longer it will take. It is recommended to use multiple cores for this step. Default permutation time is 1000 which may

need 12 hrs by 4 cores. However 10,000 permutations is recommended which may cost 2 days, if high confidence results are desired.

```
### identify target gene for significantly hypomethylated probes.
Sig.probes <- read.csv("./ELMER.example/Result/LUSC/getMethdiff.hypo.probes.significant.csv",
                      stringsAsFactors=F)[,1]
head(Sig.probes) # significantly hypomethylated probes
## [1] "cg00045114" "cg00050294" "cg00093522" "cg00163018" "cg00173804" "cg00223046"
## Collect nearby 20 gene for Sig.probes
nearGenes <-GetNearGenes(TRange=getProbeInfo(mee,probe=Sig.probes),</pre>
                        geneAnnot=getGeneInfo(mee),cores=detectCores()/2)
## Identify significant probe-gene pairs
Hypo.pair <-get.pair(mee=mee,probes=Sig.probes,nearGenes=nearGenes,
                    permu.dir="./ELMER.example/Result/LUSC/permu",permu.size=300,Pe = 0.01,
                    dir.out="./ELMER.example/Result/LUSC",cores=detectCores()/2,label= "hypo")
## Calculate empirical P value.
head(Hypo.pair) ## significant probe-gene pairs
##
                           Probe GeneID Symbol Distance Sides
                                                                        Raw.p
                                                                                      Pe
## cg20701183.ID8543 cg20701183 ID8543 LM04 2563 L1 7.453984e-14 0.003322259
## cg19403323.ID255928 cg19403323 ID255928 SYT14 87477 R1 1.671937e-12 0.003322259
## cg12213388.ID84451 cg12213388 ID84451 KIAA1804 993548 L4 2.527644e-12 0.003322259
## cg26607897.ID55811 cg26607897 ID55811 ADCY10 292476 R3 4.593610e-12 0.003322259
## cg10574861.ID8543 cg10574861 ID8543 LM04
                                                    4715 L1 4.770162e-12 0.003322259
## cg26607897.ID23432 cg26607897 ID23432 GPR161
                                                    563308 R6 8.048248e-12 0.003322259
# get.pair automatically save output files.
#getPair.hypo.all.pairs.statistic.csv contains statistics for all the probe-gene pairs.
#getPair.hypo.pairs.significant.csv contains only the significant probes.
dir(path = "./ELMER.example/Result/LUSC", pattern = "getPair")
## [1] "getPair.hypo.all.pairs.statistic.csv"
## [2] "getPair.hypo.pairs.significant.csv"
## [3] "getPair.hypo.pairs.significant.withmotif.csv"
```

5.4 Motif enrichment analysis on the selected probes

Function, get.enriched.motif, will be used to calculate enrichment of the motifs from factorbook and JASPER for the selected probes. Odds Ratio is used to assess the enrichment levels and 95% confidence interval of Odds Ratio is calculated.

```
min.incidence = 10,lower.OR = 1.1)
### 6 motifs are enriched.
names(enriched.motif) # enriched motifs
## [1] "AP1" "BARHL2" "IRF" "PRDM1" "TCF7L2" "TP53"
# get.enriched.motif automatically save output files.
# getMotif.hypo.enriched.motifs.rda contains enriched motifs and the probes with the motif.
# getMotif.hypo.motif.enrichment.csv contains summary of enriched motifs.
dir(path = "./ELMER.example/Result/LUSC", pattern = "getMotif")
## [1] "getMotif.hypo.enriched.motifs.rda" "getMotif.hypo.motif.enrichment.csv"
# motif enrichment figure will be automatically generated.
dir(path = "./ELMER.example/Result/LUSC", pattern = "motif.enrichment.pdf")
## [1] "hypo.motif.enrichment.pdf"
```

5.5 Identifying regulatory TF

Function, get.TFs, will use the anti-correlation of a particular TF and the level of demethylation at its binding sites to predict the regulatory TF.

```
### identify regulatory TF for the enriched motifs
load("./ELMER.example/Result/LUSC/getMotif.hypo.enriched.motifs.rda")
TF <- get.TFs(mee=mee, enriched.motif=enriched.motif,dir.out="./ELMER.example/Result/LUSC",
              cores=detectCores()/2, label= "hypo")
# get. TFs automatically save output files.
# getTF.hypo.TFs.with.motif.pvalue.rda contains statistics for all TF with average
# DNA methylation at sites with the enriched motif.
# getTF.hypo.significant.TFs.with.motif.summary.csv contains only the significant probes.
dir(path = "./ELMER.example/Result/LUSC", pattern = "getTF")
## [1] "getTF.hypo.significant.TFs.with.motif.summary.csv"
## [2] "getTF.hypo.TFs.with.motif.pvalue.rda"
# TF ranking plot based on statistics will be automatically generated.
dir(path = "./ELMER.example/Result/LUSC/TFrankPlot", pattern = "pdf")
## [1] "AP1.TFrankPlot.pdf"
                                 "BARHL2.TFrankPlot.pdf" "FOX.TFrankPlot.pdf"
## [4] "IRF.TFrankPlot.pdf"
                                 "MYC_USF.TFrankPlot.pdf" "NFKB1.TFrankPlot.pdf"
## [7] "PRDM1.TFrankPlot.pdf"
                                 "SOX2.TFrankPlot.pdf"
                                                          "SPI1.TFrankPlot.pdf"
## [10] "TCF7L2.TFrankPlot.pdf"
                                 "TP53.TFrankPlot.pdf"
                                                          "UA7.TFrankPlot.pdf"
## [13] "UA9.TFrankPlot.pdf"
```

6 Generating figures

6.1 Scatter plots

6.1.1 Scatter Plot of 20 nearby genes

Generate scatter plots for one probes' nearby 20 gene expression vs DNA methylation at this probe. Figure 1

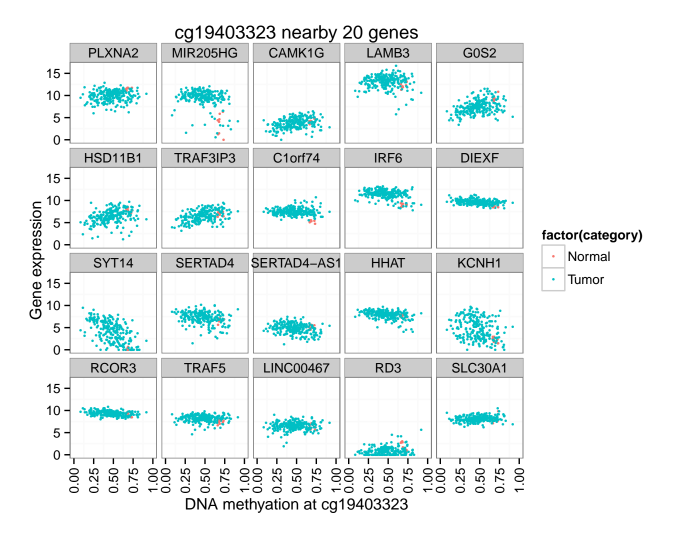


Figure 1: Each scatter plot shows the methylation level of an example probe cg19403323 in all LUSC samples plotted against the expression of one of 20 adjacent genes.

6.1.2 Scatter Plot of One Pair

```
Generate a scatter plot for one probe-gene pair. Figure 2
```

6.1.3 TF expression vs. average DNA methylation

Generate scatter plot for TF expression vs average DNA methylation of the sites with certain motif. Figure 3

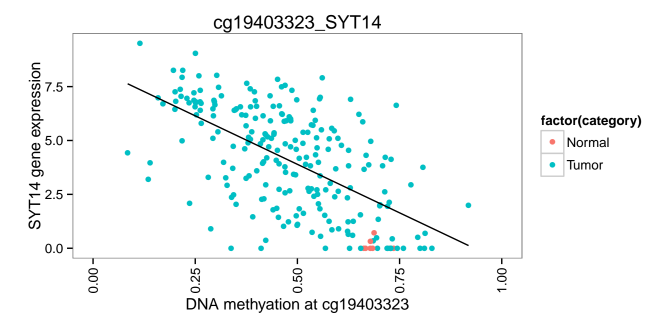


Figure 2: Scatter plot shows the methylation level of an example probe cg19403323 in all LUSC samples plotted against the expression of the putative target gene SYT14.

6.2 Schematic plot

Schematic plot shows a breif view of linkages between genes and probes. To make a schematic plot, "Pair" object should be generated first.

6.2.1 Nearby Genes

Generate schematic plot for one probe with 20 nearby genes and label the gene significantly linked with the probe in red. Figure 4

```
grid::grid.newpage()
schematic.plot(pair=pair, byProbe="cg19403323", save=FALSE)
## cg19403323
```

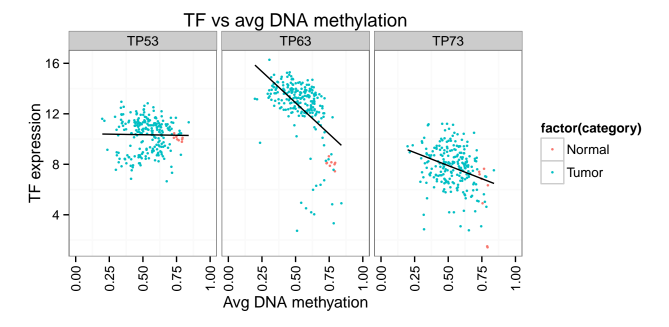


Figure 3: Each scatter plot shows the average methylation level of sites with the TP53 motif in all LUSC samples plotted against the expression of the transcription factor TP53, TP63, TP73 respectively.

6.2.2 Nearby Probes

Generate schematic plot for one gene with the probes which the gene is significantly linked to. Figure 5

```
grid::grid.newpage()
schematic.plot(pair=pair, byGene="ID255928", save=FALSE)
```

6.3 Motif enrichment plot

Motif enrichment plot shows the enrichment levels for the selected motifs. Figure6

6.4 TF ranking plot

TF ranking plot shows statistic $-log10(P \ value)$ assessing the anti-correlation level of TFs expression level with average DNA methylation level at sites with a given motif. Figure 7

2.15Mb

cg19403323

chr1:209605478-211752100

SYT14 SERTAD4-AS1

Figure 4: The schematic plot shows probe colored in blue and the location of nearby 20 genes. The genes significantly linked to the probe were shown in red.

```
sessionInfo()
## R version 3.1.3 (2015-03-09)
## Platform: x86_64-apple-darwin14.1.0 (64-bit)
## Running under: OS X 10.10.2 (Yosemite)
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
## [1] stats4
                parallel stats
                                    graphics grDevices utils
                                                                 datasets methods
## [9] base
##
## other attached packages:
## [1] ELMER_0.99.4
                           GenomicRanges_1.18.4 GenomeInfoDb_1.2.4 IRanges_2.0.1
## [5] S4Vectors_0.4.0
                           BiocGenerics_0.12.1
##
## loaded via a namespace (and not attached):
## [1] BiocStyle_1.4.1 codetools_0.2-11 colorspace_1.2-6 digest_0.6.8
                                                                          evaluate_0.5.5
## [6] formatR_1.0
                        ggplot2_1.0.0
                                         grid_3.1.3
                                                         gtable_0.1.2
                                                                          highr_0.4
## [11] knitr_1.9
                        labeling_0.3
                                         MASS_7.3-39
                                                         munsell_0.4.2
                                                                          plyr_1.8.1
## [16] proto_0.3-10
                        Rcpp_0.11.5
                                         reshape_0.8.5
                                                         reshape2_1.4.1
                                                                          scales_0.2.4
## [21] snow_0.3-13
                        stringr_0.6.2
                                         tools_3.1.3 XVector_0.6.0
```

chr1:209537469-210111538

0.57Mb

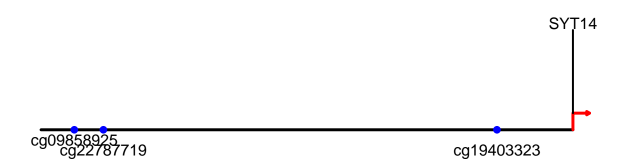


Figure 5: The schematic plot shows the gene colored in red and all blue colored probes, which are significantly linked to the expression of this gene.

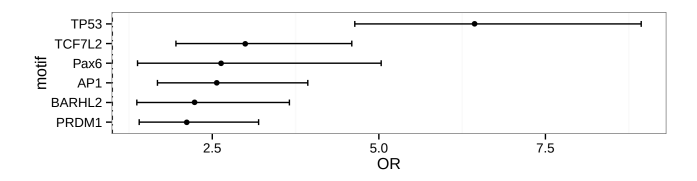


Figure 6: The plot shows the Odds Ratio (x axis) for the selected motifs with OR above 1.3 and lower boundary of OR above 1.3. The range shows the 95% confidence interval for each Odds Ratio.

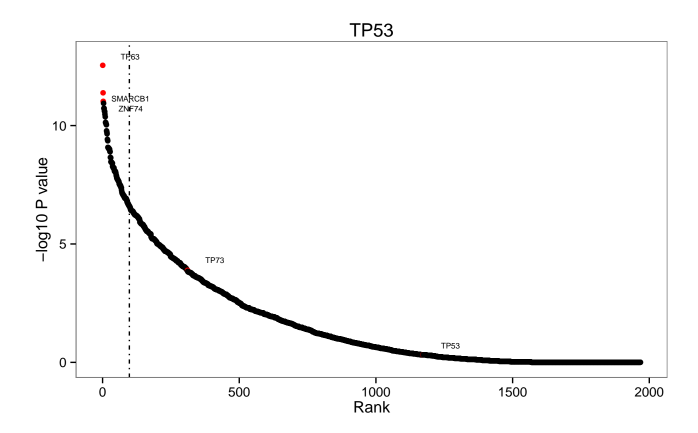


Figure 7: Shown are TF ranking plots based on the score ($-\log(P \text{ value})$) of association between TF expression and DNA methylation of the TP53 motif in the LUSC cancer type . The dashed line indicates the boundary of the top 5% association score. The top 3 associated TFs and the TF family members (dots in red) that are associated with that specific motif are labeled in the plot.