

**Supplementary Text S2: R codes to run FAIME**

```
#####
# "FAIME" is a new algorithm to predict Functional Analysis of Individual Microarray Expression.
#
# This R code is composed one main function "runFAIME", which is a R wrap for FAIME.      #
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# Last updated on Feb 22th, 2010.                                                       #
# Version 2                                                                           #
# #
# Update recorder:                                                               #
# Added a parameter weightRank=TRUE; if FALSE, no weighted rank will be used instead   #
#####

#####
## Pre-required R package    ##
#####
## To install Bioconductor three packages, include the following in your R code.
## source("http://bioconductor.org/biocLite.R")
## biocLite("Biobase")

library("Biobase")

# Declaration of instants which need to be locally modified by user.
ArrayInput                      # mRNA-expression input file name.
Geneset2gene                     # an 2 column matrix mapping between GO to genes
FDR.T = 0.05                     # Threshold to call a gene-set as significant
genewprobe                        # array of microarray probe IDs with annotated gene symbol

#####
## The key function to implement FAIME algorithm    ##
#####
## parameters (inputs):
## sampleExp: A vector of normalized exp value with probeName
## genemembers: A vector of probeName of the gene members of a geneset of interest
## weightRank: A logical to decide whether weighted rank to be used, default is TRUE.
##      if weithRank="mild", the exp( (rank-N)/N) was applied which controls the weights
## between 0 and 1
## output:
## y: A score assigned to each microRNA

FAIME <- function(sampleExp, genemembers, na.last=TRUE, weightRank,
logCheck=FALSE)
{
  # check if it log transformed, if not, log transformed #
  if (logCheck) {      # by this transformation, negative value will be scaled to NA #
    if (max(sampleExp) > 20) sampleExp <- log2(SampleExp)
  }

  if (any(is.na(names(sampleExp)))) stop("Please input sampleExp with probe IDs")
  allGenes <- names(sampleExp)
  N <- length(allGenes)
  nongenemembers <- allGenes[-which(allGenes %in% genemembers)]
```

```

# Step 1: Calculation of weighted rank of gene expression #
# Rank from the lowest to highest, thus the leading up-regulated genes get the higher
weighted score #
rankedExp <- rank(sampleExp, na.last=na.last)
if(weightRank=="mild") {
    rankscore <- rankedExp*exp((rankedExp-N)/N)
} else {
    if (weightRank) {
        rankscore <- rankedExp*exp(rankedExp/N)
    } else {
        rankscore <- rankedExp
    }
}

# Step 2: Calculate F-score for each individual gene-set per a sample using mRNA
expression of their gene members and that of its none-members #
ST <- sum(rankscore[genemembers])/length(genemembers)
SN <- sum(rankscore[nongenemembers])/length(nongenemembers)
y <- ST - SN
return(y)
}

#####
## The main function to run FAIME ##
#####

## parameters (inputs):
## dat: An expression matrix, row for a probeName and column for a sample
## genewprobe: A vector of gene Symbol for rows of dat, the names of which is probeName
## geneset2gene: An one-to-one mapping matrix with two columns,
##      the 1st column is geneset ID/name, and the 2nd is its gene members
## weightRank: A logical to decide whether weighted rank to be used, default is TRUE.
## if weightRank="mild" the exp( (rank-N)/N) was applied which controls the weights
##      between 0 and 1
## output:
## res: A matrix of transformed microRNA profiling, the score calculated by FAIME

runFAIME <- function(dat, genewprobe, geneset2gene, na.last=TRUE, weightRank=TRUE)
{
    if (class(dat)=="ExpressionSet") dat <- exprs(dat)
    if (is.null(names(genewprobe))) stop("Please input the zz as a named vector for gene
Symbol!")
    allSym <- genewprobe [rownames(dat)]
    seeds <- unique(geneset2gene[,1])
    res <- matrix(nrow=length(seeds), ncol=ncol(dat))
    rownames(res) <- seeds
    colnames(res) <- colnames(dat)
    geneIDs <- rownames(dat)

    for(i in 1:length(seeds))
    {
        genemembers <- geneset2gene[which(geneset2gene [,1]==seeds[i]),2]

```

```
targetP <- unlist(allSym[which(allSym %in% genemembers)] )  
for (j in 1:ncol(dat))  
{  
  oneSampleExp <- dat[,j]  
  names(oneSampleExp) <- geneIDs  
  res[i,j] <- FAIME(oneSampleExp, genemembers=names(targetP),  
    na.last=na.last, weightRank=weightRank)  
}  
}  
length(which(is.na(res[,1]))) # there might be geneset without measured member gene on  
this array #  
return(res)  
}
```