#### Introduction

This tutorial is for running the FASE pipeline. The pipeline is based on differential alternative splicing events and predicts the transcript structure and their concentration along with survival analysis. In fact this is the first kind of pipeline that takes advantage of differential alternative splicing events for finding novel transcripts that are neglected due to low expression in transcript level statistical analysis.

#### Installation

Most of the R libraries required by FASE will be installed by default along with FASE installation, which includes limma, edgeR, Rsubread, parallel, etc. Besides, following tools must be installed on Linux system before starting the analysis with FASE if pre-processing functions (see part 1 of the tutorial) are being used. However, if read counts have already been summarized, these tools are not required:

- 1. <u>fastq-dump</u> is required for converting SRA files to fastq files. It is required in order to pre-process sra files automatically.
- 2. bowtie2 is required to build a genome index, for read mapping by tophat2.
- 3. tophat2 is required for mapping reads to reference genome.
- 4. samtools is required to sort and index the bam files.

# Sample case study

1. <u>SRA files</u>: In this tutorial, SRA files from NCBI SRA BioProject ID: PRJNA342407 have been used. In this study, one set of C. elegans (4 biological replicates for each condition) were grown on OP50: control (WT) and treated (eat-2(ad1116)(-)) and another set were grown on RNAi-seeded plates: control and treated (hrpu-1 or smg-2). For this tutorial, two samples each from control (WT) and eat-2 (-) should be downloaded using accession IDs: SRR5606854 (WT Day 1-04), SRR5606849 (WT Day 1\_03), SRR5606847 (eat-2(-) Day 1\_03) and SRR5606848 (eat-2(-) Day 1\_04). Alternative splicing, differential gene expression and differential junction expression would be compared between the WT and eat-2 (-) C. elegans. All four files are paired-end.

For a comprehensive analysis, the SRA files should be renamed as WT-R1, WT-R2, eat2-R1 and eat2-R2, respectively. These files should be saved in an empty folder.

- 2. <u>GTF file</u>: GTF file is required for adding annotation to genes, junctions and alternative splicing events. GTF for C. elegans (WBcel235) can be downloaded from Ensembl (or using link: https://asia.ensembl.org/Caenorhabditis elegans/Info/Index).
- 3. <u>Reference genome</u>: It is required by tophat2 for mapping the reads. Reference genome for C. elegans (WBcel235) can be downloaded from Ensembl (or using link: https://asia.ensembl.org/Caenorhabditis\_elegans/Info/Index).

GTF and reference genome should be saved together in a folder, separate from SRA files.

FASE needs to pre-processes RNA-Seq data before analysing alternative splicing, differential gene expression and differential junction expression. Therefore the tutorial is divided in two parts: Pre-processing and downstream analysis. Pre-processing can be performed by any of the three methods, depending on user preference: (i) using automatic pre-processing function: ppAuto, (ii) using three sequential wrapper functions: ppRawData, ppSumEIG and ppFASE, or (iii) using manual tools and functions. For this purpose, the pre-processing part is sub-divided in three sections.

## Open a linux terminal in the directory containing reference genome # Step:1. Running bowtie to build genome index on linux terminal. This will generate six files containing genome index in the same directory as reference genome. This step needs to be performed only once for a reference genome.

bowtie2-build Caenorhabditis\_elegans.WBcel235.dna.toplevel.fa Caenorhabditis\_elegans.WBcel235.dna.toplevel # done

## Open an R session in the directory containing SRA files # Step:2. Installing and loading FASE.

install.packages('FASE')
library(FASE)

# done

# Step:3. Declaring genomeBI with the path of reference genome index. Please note that reference genome filename should not contain '.fa' extension while declaring genomeBI. genomeBI <- 'path/Caenorhabditis\_elegans.WBcel235.dna.toplevel' # done

## Step:4. Parsing the downloaded GTF file for possible junctions which is required to find intron retention events using intronGTF parser function. An intron parsed GTF file will be generated by default with '\_corrected' appended to the GTF filename, in the same directory as downloaded GTF file. This GTF file should be used in FASE. Note: intronGTF parser needs to be run only once for a GTF file.

gtf <- intronGTFparser('path/Caenorhabditis\_elegans.WBcel235.99.gtf')
# done</pre>

## Step:5. Declaring number of cores to be used throughout the analysis. tophat2, samtools, featureCounts, EPrnaseq and iPrnaseq allow parallel processing.

p <- 1 # done

## Step:6. Running ppAuto function. ppAuto performs: (i) SRA conversion to fastq using fastq-dump, (ii) read mapping using tophat2, (iii) sorting and indexing bam files using samtools, (iv) generation of junction read count matrix, (v) summarization of read counts for exons, introns and genes, (vi) generation of readMembershipMatrix,

intronMembershipMatrix and Gcount matrix (required detection of alternative splicing events).

# After completion of ppAuto command, following files would be generated in the folder containing SRA files: eight fastq files (with \_1 and \_2 appended to each SRA filename), srlist.Rdata that contains list of samples, four directories (WT-R1\_tophat\_out, WT-R2\_tophat\_out, eat2-R1\_tophat\_out and eat2-R2\_tophat\_out) containing tophat output, JunctionCounts.Rdata containing read counts for mapped junctions, counts\_genes.Rdata containing read counts for all mapped genes, counts\_exons.Rdata containing read counts for all mapped exons, counts\_introns.Rdata contining read counts of all mapped introns, RMM.data containing read membership matrix (association of exons with exons, introns and junctions (skipping and flanking)), iMM.Rdata containing intron membership matrix (association of introns with exons, introns and junctions (skipping and flanking)) and Gcount.Rdata containing per gene meta-feature read counts.

ppAuto(p = p, genomeBI = genomeBI, gtf = gtf, pairedend = T, files = 'sra') # pre-processing done

## Open a linux terminal in the directory containing reference genome # Step:1. Running bowtie to build genome index on linux terminal. This will generate six files containing genome index in the same directory as reference genome. This step needs to be performed only once for a reference genome.

bowtie2-build Caenorhabditis\_elegans.WBcel235.dna.toplevel.fa Caenorhabditis\_elegans.WBcel235.dna.toplevel # done

## Open an R session in the directory containing SRA files # Step:2. Installing and loading FASE. install.packages('FASE') library(FASE) # done

# Step:3. Declaring genomeBI with the path of reference genome index. Please note that reference genome filename should not contain '.fa' extension while declaring genomeBI. genomeBI <- 'path/Caenorhabditis\_elegans.WBcel235.dna.toplevel' # done

## Step:4. Parsing the downloaded GTF file for possible junctions which is required to find intron retention events using intronGTF parser function. An intron parsed GTF file will be generated by default with '\_corrected' appended to the GTF filename, in the same directory as downloaded GTF file. This GTF file should be used in FASE. Note: intronGTF parser needs to be run only once for a GTF file.

gtf <- intronGTFparser('path/Caenorhabditis\_elegans.WBcel235.99.gtf')
# done</pre>

## Step:5. Declaring number of cores to be used throughout the analysis. tophat2, samtools, featureCounts, EPrnaseq and iPrnaseq allow parallel processing.

p <- 1 # done

## Step:6. Calling ppRawData function. ppRawData requires genome build index and intron parsed GTF file to: (i) convert SRA files to fastq files using fastq-dump, (ii) map RNA-Seq reads to the reference genome using tophat2, (iii) sort and index bam files using samtools, and (iv) generate junction count matrix. After completion of ppRawData command, following files would be generated in the folder containing SRA files: eight fastq files (with \_1 and \_2 appended to each SRA filename), srlist.Rdata that contains list of samples, four directories (WT-R1\_tophat\_out, WT-R2\_tophat\_out, eat2-R1\_tophat\_out and eat2-R2\_tophat\_out) containing tophat output and JunctionCounts.Rdata containing read counts for mapped junctions

ppRawData(pairedend = T, p = p, genomeBI = genomeBI, gtf = gtf)
# done

# loading srlist for ppSumEIG. srlist is a list of samples.

# load('srlist.Rdata')

## Step:7. Calling ppSumEIG function. ppSumEIG is a wrapper function for featureCounts. It performs read count summarization for genes, exons and introns. For this, it requires srlist generated by ppRawData as srlist.Rdata, intron parsed GTF file and sorted and indexed bam files (passed automatically within the function). After completion, ppSumEIG saves three files in current directory: counts\_genes.Rdata, counts\_exons.Rdata and counts\_introns.Rdata.

ppSumEIG(p = p, gtf = gtf, pairedend = T, srlist = srlist)
# done

# loading summarized read counts for exons, introns and junctions for ppFASE function.

load('counts\_exons.Rdata')
load('counts\_introns.Rdata')
load('JunctionCounts.Rdata')

## Step:8. Running ppFASE. ppFASE generates three matrices required for alternative splicing analyses: read membership matrix (RMM) which contains association of exons with exons, introns and junctions (flanking and skipping) (saved as RMM.Rdata by ppFASE), intron membership matrix which contains association of introns with exons, introns and junctions (flanking and skipping) (saved as iMM.Rdata by ppFASE) and Gcount which is a list of genewise read count summarization of meta-features times samples (saved as Gcount.Rdata by ppFASE). ppFASE requires summarized intron read counts (generated by ppSumEIG and saved as counts\_introns.Rdata), exon read counts (generated by ppSumEIG and saved as counts\_exons.Rdata), junction matrix (generated by ppRawData and saved as JunctionCounts.Rdata) and intron parsed GTF file.

ppFASE(intronCount = intronCount, exonCount = exonCount, JunctionMatrix = JunctionMatrix, gtf = gtf)

# pre-processing done

# 

## Open a linux terminal in the directory containing reference genome

# Step:1. Running bowtie to build genome index on linux terminal. This will generate six files containing genome index in the same directory as reference genome. This step needs to be performed only once for a reference genome.

bowtie2-build Caenorhabditis\_elegans.WBcel235.dna.toplevel.fa Caenorhabditis\_elegans.WBcel235.dna.toplevel # done

## Open an R session in the directory containing SRA files
## Step:2. Generating and saving srlist (list of samples)
srlist <- dir()
save(srlist, file = 'srlist.Rdata')</pre>

## Step:3. converting sra files to fasta files

## Open a linux terminal in the directory containing SRA files. Run fastq-dump to convert SRA files to fastq files. It will generate eight fastq files (two fastq files for each SRA file as the reads are paired-end).

fastq-dump -I -v --split-files WT-R1 WT-R2 eat2-R1 eat2-R2

## Step:4. Running tophat

## Open a linux terminal (or same terminal used for fastq-dump) in the directory containing fastq files.

tophat -o ./WT-R1\_tophat\_out -p 5 -N 6 -r -44 --min-intron-length 50 --max-intron-length 5000 --mate-std-dev 30 --read-edit-dist 6

path/Caenorhabditis\_elegans.WBcel235.dna.toplevel ./WT-R1\_1.fastq ./WT-R1\_2.fastq tophat -o ./WT-R2\_tophat\_out -p 5 -N 6 -r -44 --min-intron-length 50 --max-intron-length 5000 --mate-std-dev 30 --read-edit-dist 6

path/Caenorhabditis\_elegans.WBcel235.dna.toplevel ./WT-R2\_1.fastq ./WT-R2\_2.fastq tophat -o ./eat2-R1\_tophat\_out -p 5 - N 6 - r - 44 --min-intron-length 50 - max-intron-length 5000 - mate-std-dev 30 - read-edit-dist 6

path/Caenorhabditis\_elegans.WBcel235.dna.toplevel ./eat2-R1\_1.fastq ./eat2-R1\_2.fastq tophat -o ./eat2-R2\_tophat\_out -p 5 -N 6 -r -44 --min-intron-length 50 --max-intron-length 5000 --mate-std-dev 30 --read-edit-dist 6

path/Caenorhabditis\_elegans.WBcel235.dna.toplevel ./eat2-R2\_1.fastq ./eat2-R2\_2.fastq #done

## Step:5. Sorting bam files.

## Open a linux terminal in the directory containing the tophat output. Run samtools sort for sorting the bam files generated by tophat2. It will generate a file, accepted\_hits\_sorted.bam, for each tophat output. The unsorted bam files, accepted\_hits.bam, can be deleted after running this command. @ option in samtools sort denotes number of cores.

samtools sort accepted\_hits.bam -o accepted\_hits\_sorted.bam -@ 1
# repeat this for each tophat output

## Step:6. Indexing the sorted bam files.

## Open a linux terminal in the directory containing the tophat output. Run samtools index for indexing the sorted bam files. It will generate a file, accepted\_hits\_sorted.bam.bai. samtools index accepted\_hits\_sorted.bam

# repeat this for each tophat output.

# if parallel is installed in the system, following command can be used instead of running samtools index for each tophat output. j in this command denotes the number of cores to be used by parallel.

# parallel -j 4 'samtools index {}' ::: ./\*/\*sorted.bam

# done

## Step:7. Counting junction reads

## Open an R session in the directory containing fastq files.

# Installing and loading FASE

install.packages('FASE')

library(FASE)

# Declaring folderSRA as current directory and loading the list of sample names (srlist).

folderSRA <- getwd()

load('srlist.Rdata')

# Creating list of junctions.bed files of all samples. This list will be passed to getJunctionCountMatrix to form junction matrix.

jfiles <- unlist(lapply(srlist, function(x) paste(folderSRA, '/' ,x ,'\_tophat\_out/junctions.bed'
,sep = ", collapse = "")))</pre>

# Calling getJunctionCountMatrix. This function requires list of junctions.bed files of all samples and returns a junction matrix (read counts of junctions) using them.

JunctionMatrix <- getJunctionCountMatrix(jfiles)</pre>

# Adding sample names to the junction matrix

colnames(JunctionMatrix) <- c(colnames(JunctionMatrix)[1:5], srlist)
save(JunctionMatrix, file = 'JunctionCounts.Rdata')</pre>

# done

## Step:8. Read count summarization

# Continue in the previous R session (used for making junction matrix)

# Parsing the downloaded GTF file for possible junctions which is required to find intron retention events using intronGTF parser function. An intron parsed GTF file will be generated by default with '\_corrected' appended to the GTF filename, in the same directory as downloaded GTF file. This GTF file should be used in FASE. Note: intronGTF parser needs to be run only once for a GTF file.

gtf <- intronGTFparser('path/Caenorhabditis elegans.WBcel235.99.gtf')

# Creating list of bam files (required for featureCounts)

srlistbam <- paste(folderSRA,'/',srlist,'\_tophat\_out/accepted\_hits\_sorted.bam',sep = ")
# Declaring number of cores to be used for featureCounts.</pre>

p <- 1

# Loading Rsubread package, which contains featureCounts function library(Rsubread)

# Read summarization for exons. Summarization step requires srlistbam, intron parsed gtf file and number of cores. The output of summarization is a matrix of read counts of exons which can be saved as counts exons. Rdata for use in analysis part.

exonCount<- featureCounts(files = srlistbam, isPairedEnd = TRUE, requireBothEndsMapped = TRUE, GTF.featureType = "exon", GTF.attrType = "gene\_id", useMetaFeatures = FALSE, isGTFAnnotationFile = TRUE, annot.ext = gtf, allowMultiOverlap = TRUE, nthreads = p) save(exonCount, file = 'counts exons.Rdata')

# Read summarization for introns. Read count summarization for introns requires srlistbam, intron parsed gtf file and number of cores. It returns a matrix of read counts of introns which can be saved as counts introns. Rdata for use in analysis part.

intronCount<- featureCounts(files = srlistbam, isPairedEnd = TRUE, requireBothEndsMapped = TRUE, GTF.featureType = "intron", GTF.attrType = "gene\_id", useMetaFeatures = FALSE, isGTFAnnotationFile = TRUE, annot.ext = gtf, allowMultiOverlap = TRUE, nthreads = p)

save(intronCount, file = 'counts\_introns.Rdata')

# Read summarization for genes. Summarization step requires srlistbam, intron parsed gtf file and number of cores. It returns a matrix of read counts of genes which can be saved as counts\_genes.Rdata for use in analysis part.

geneCount<- featureCounts(files = srlistbam, isPairedEnd = TRUE, requireBothEndsMapped
= TRUE, GTF.featureType = "gene", GTF.attrType = "gene\_id", useMetaFeatures = FALSE,
isGTFAnnotationFile = TRUE, annot.ext = gtf, allowMultiOverlap = TRUE, nthreads = p)
save(geneCount, file = 'counts\_genes.Rdata')
# done</pre>

## Step:9. Generating readMembershipMatrix (RMM).

# Continue in the previous R session (used for making junction matrix)

# readMembershipMatrix saves RMM.Rdata by default. RMM contains the association of exons with other exons, introns and junctions (skipping and flanking). It is required for finding cassette exon events. It also saves Annotation.Rdata file, which contains annotation for exons and introns. readMembershipMatrix function requires intron parsed GTF file and junction matrix (JunctionCounts.Rdata).

# Loading junction matrix

load('JunctionCounts.Rdata')

RMM <- readMembershipMatrix (gtf, JunctionMatrix)

# done

## Step:10. Generating intronMembershipMatrix (iMM).

# intronMembershipMatrix saves iMM.Rdata by default. iMM contains the association of introns with exons, introns and junctions (skipping and flanking). It is required for finding intron retention events. intronMembershipMatrix function requires annotation file generated and saved by readMembershipMatrix as Annotation.Rdata.
# Loading Annotation.Rdata

load('Annotation.Rdata')

iMM <- intronMembershipMatrix(annotation = annotation)</pre>

# done

## Step:10. Generating Gcount matrix. Gcount matrix is a list of gene-wise read count summarization of meta-features times samples. It is required for finding both cassette exon and intron retention events. countMatrixGenes requires the junction matrix (saved as JunctionCounts.Rdata), intron read counts (saved as counts\_introns.Rdata), exon read counts (saved as counts\_exons.Rdata) and annotation (saved as Annotation.Rdata by readMembershipMatrix). It saves Gcount by default as Gcount.Rdata.

# Loadind exon counts, inron counts, annotation and junction matrix. load('counts\_exons.Rdata') load('counts\_introns.Rdata') load("Annotation.Rdata") load("JunctionMatrix.Rdata")

Gcount <- countMatrixGenes(JunctionMatrix, annotation = annotation, intronList = intronCount, exonList = exonCount)

# pre-processing done

# Now downstream analysis can be performed on this dataset.

## Step:1. Preparing design matrix, contrast matrix and groups for limma. # Design matrix: design matrix describes the samples in the experiment. In this tutorial, WT-R1 and WT-R2 belong to control group and eat-R1, and eat2-R2 belong to eat2 group. designM <- matrix(c(rep(1,2), rep(0,4), rep(1,2)), byrow = F, ncol = 2, nrow = 4) colnames(designM) <- c('control', 'eat2') rownames(designM) <- c('WT\_R1', 'WT\_R2', 'eat2\_R1', 'eat2\_R2')

# Contrast matrix describes which sample groups should be compared. In this tutorial, eat2 is being comapred with control samples.

contrastM <- matrix(c(-1, 1), ncol = 1)
rownames(contrastM) <- colnames(designM)
colnames(contrastM) <- 'control\_vs\_eat2'</pre>

# Groups vector shows the groups to which the samples belong to (in sequence).

Groups <- c(1,1,2,2) save(designM, contrastM, Groups, file = 'DCmatrix.Rdata') # done

## Step:2. Alternative splicing: cassette exon events. (EP)
# EPrnaseq requires Gcount matrix and RMM, generated by
ppAuto/ppFASE/readMembershipMatrix. It also requires a design matrix, contrast matrix
and Groups. These matrices can be prepared as mentioned in Part 2 step 1. Threshold
parameter in EPrnaseq is the minimum number of reads that should map to a meta-feature
(intron/exon/junction).

# loading Gcount and RMM

load('Gcount.Rdata')

#### load('RMM.Rdata')

# fitting the data to EPrnaseq function, which returns a ranked list of cassette exon events for all contrasts.

fit<- EPrnaseq(Gcount = Gcount, RMM = RMM, designM = designM, contrastM = contrastM, Groups = Groups, p = p, threshold = 3)

# getPvaluesByContrast returns EPrnaseq ranking for a given contrast. Any contrast from contrast matrix can be passed as shown below. In this case, there is only one contrast 'control vs eat2'.

control vs eat2 <- getPvaluesByContrast(fit, 'control vs eat2')</pre>

# saving the ranking for given contrast in csv format.

write.csv(control\_vs\_eat2, file = 'EP\_control\_vs\_eat2.csv')

# OPTIONAL step.

# cpmCountsEP saves CPM counts and log2CPM expression values for ranked cassette exon events of a particular contrast (in csv format). cpmCountsEP function should be passed with the file containing ranking for a contrast, design matrix and Groups.

cpmCountsEP('EP\_control\_vs\_eat2.csv', designM = designM, Groups = Groups)
# (EP done)

## Step:3. Alternative splicing. intron retention events. (IP)

# iPrnaseq requires Gcount matrix and iMM, generated by

ppAuto/ppFASE/intronMembershipMatrix. It also requires a design matrix, contrast matrix and Groups. These matrices can be prepared as mentioned in Part 2 step 1. Threshold parameter in iPrnaseq is the minimum number of reads that should map to a meta-feature (intron/exon/junction).

# loading Gcount and iMM

load('Gcount.Rdata')

load('iMM.Rdata')

# fitting the data to iPrnaseq function, which returns a ranked list of intron retention events for all contrasts.

fit<- iPrnaseq(Gcount = Gcount, iMM = iMM, designM = designM, contrastM = contrastM, Groups = Groups, p = p, threshold = 3)

# getPvaluesByContrast returns iPrnaseq ranking for a given contrast. Any contrast from contrast matrix can be passed as shown below. In this case, there is only one contrast 'control vs eat2'.

control vs eat2 <- getPvaluesByContrast(fit, 'control vs eat2')</pre>

# saving the ranking for given contrast in csv format.

write.csv(control\_vs\_eat2, file = 'IP\_control\_vs\_eat2.csv')

# OPTIONAL step.

# cpmCountsEP saves CPM counts and log2CPM expression values for ranked intron retention events of a particular contrast (in csv format). The function requires the file containing ranking for a contrast, design matrix and Groups.

cpmCountsEP('IP\_control\_vs\_eat2.csv', designM = designM, Groups = Groups)
# (IP done)

##Step:4. Transcript Structure

#transtruct reauires Gocunt, RMM, iMM, annotation, geneID, EP event ID, IP event ID, design matrix and Groups. These can be generated during pre-processing using ppFASE or ppAuto.

#loading Gcount, iMM, RMM and annotation.

load('Gcount.Rdata') load('RMM.Rdata') load('iMM.Rdata') load('Annotation.Rdata')

#providing geneID and ep event ID

geneID <- 'WBGene00011848 ep.event <- 'EX104010'

#extracting matrices corresponding to geneID

index\_gene <- match(geneID, names(RMM))
RMM <- RMM[[index gene]]</pre>

index\_gene <- match(geneID, names(iMM))
iMM <- iMM[[index gene]]</pre>

index\_gene <- match(geneID, names(Gcount))
Gcount <- Gcount[[index\_gene]]</pre>

index\_ann <- match(annotation\$genes, geneID)
annotation <- annotation[!is.na(index\_ann),, drop = FALSE]</pre>

#defining whether or not we want to use flanking intron(s) for finding seed exon(s) keep.intron <- FALSE

#main function

ts <- transtruct(ep.event = ep.event, RMM = RMM, iMM = iMM, Gcount = Gcount, designM = designM, Groups = Groups, annotation = annotation, keep.intron = keep.intron)

##transcript structure done

##Step:5. Transcript Concentration

#transconc finds concentration of each transcript structure generated by transtruct, both sample-wise and condition-wise.

transtruct <- ts

tc <- transconc(transtruct = transtruct, designM = designM)

##transcript concentration done

##Step:6. Survival analysis

# For survival analysis, we are using expression data from Pani et. al. (2021) and corresponding clinical data from TCGA. Normally, clinical data needs to be filtered and matched according to the expression data. However, for this tutorial, the required files can be downloaded from https://github.com/harshsharma-cb/FASE/survivaldata.Rdata.

survivaldata.Rdata contains Gcount matrix, clinical data, RMM, and exon ID for a cassette exon event in Exon8 of CerS2 gene in the LuminalB sub-type. #running survival analysis.

load('survivaldata.Rdata')

survival.result <- survFASE(Gcount = Gcount, clinical.data = clinical.data, rmm = rmm, imm=NULL, eventID = exonID, threshold = threshold, design = designM, Groups = Groups) ##survival analysis done

##Step:7. Network analysis

#FASE uses alternative splicing results to generate splice index which can either directly be used for network analysis (e.g. as heat score in HotNet2) or can be modified into a similarity score to be used as edge weight for an unweighted network (e.g. ClusterONE).

#heatscore function calculates splice index using expression data of EP and IP events.

heatscore <- heatscore(ep = "EP\_control\_vs\_eat2.csv", ep.exp =
"EP\_control\_vs\_eat2\_log2cpm.csv", ip = "IP\_control\_vs\_eat2.csv", ip.exp =
"IP\_control\_vs\_eat2\_log2cpm.csv")</pre>

#simscore function calculates similarity score for gene pairs.

simscore <- simscore(ep = "EP\_control\_vs\_eat2.csv", ep.exp =
"EP\_control\_vs\_eat2\_log2cpm.csv", ip = "IP\_control\_vs\_eat2.csv", ip.exp =
"IP\_control\_vs\_eat2\_log2cpm.csv")</pre>

##network analysis done

##Step:8. Differential gene expression (DEG)

# DEG is a wrapper function for limma, which requires summarized read counts for genes (saved as counts\_genes.Rdata by ppAuto/ppSumEIG), design matrix, contrast matrix and Groups. The output of DEG is an object of MArrayLM, which stores the result of fitting genewise linear models to the normalized intensities or log-ratios. The output is futher processed to obtain ranking of differentially expressed genes, their CPM read counts and log2CPM expression values.

# loading geneCount

#### load('counts genes.Rdata')

# fitting the data using DEG. It returns an object of MArrayLM with several statistics related to differential expression like p-value, adjusted p-value, t-statistic, etc.

fit <- DEG(geneCount = geneCount, designM = designM, contrastM = contrastM, Groups =
Groups)</pre>

# addAnnotationDEG adds annotation to the DEG fit object, according to given contrast. The function requires geneCount (generated by ppAuto/ppSumEIG in counts\_genes.Rdata), output of DEG and a contrast from contrast matrix. It returns ranking of differentially expressed genes for the given contrast, which can be saved in csv format.

control\_vs\_eat2 <- addAnnotationDEG(geneCount, fit, 'control\_vs\_eat2')
# saving the ranking for given contrast in csv format.</pre>

write.csv(control\_vs\_eat2, file = 'DEG\_control\_vs\_eat2.csv')
# OPTIONAL step.

# cpmCountsDEG saves CPM counts and log2CPM expression values for ranked differentially expressed genes of a particular contrast (in csv format). cpmCountsDEG function should be passed the file containing ranking for a contrast and geneCount.

cpmCountsDEG(geneCount, 'DEG control vs eat2.csv')

## Step:9. Differential junction expression

# DEJ is a wrapper function for limma to find differentially expressed junctions. Before adding annotation to ranked differentially expressed junction using addAnnotationDEJ, the junction matrix should be annotated using JunctionMatrixAnnotation (output is saved by default as JunctionMatrixAnnotation.Rdata). DEJ requires junction matrix (generated by ppAuto/ppRawData and saved as JunctionCounts.Rdata), design matrix, contrast matrix and Groups.

# loading junction matrix

# load('JunctionCounts.Rdata')

# annotating junction matrix using GTF file and junction matrix. The output will be saved as JunctionMatrixAnnotation.Rdata.

# JunctionMatrixAnnotation(gtf = gtf, JunctionMatrix)

# fitting the data using DEG. It returns an object of MArrayLM with several statistics related to differential expression like p-value, adjusted p-value, t-statistic, etc.

# fit <- DEJ(JunctionMatrix = JunctionMatrix, designM = designM, contrastM = contrastM, Groups = Groups)</pre>

# addAnnotationDEJ adds annotation to the DEJ fit object, according to given contrast. addAnnotationDEJ function requires annotated junction matrix JunctionMatrixA (generated by JunctionMatrixAnnotation as JunctionMatrixAnnotation.Rdata), output of DEJ and a contrast from contrast matrix. It returns ranking of differentially expressed junctions for the given contrast, which can be saved in csv format.

control\_vs\_eat2 <- addAnnotationDEJ(JunctionMatrixA, fit, 'control\_vs\_eat2')
# saving the ranking for given contrast in csv format.</pre>

write.csv(control\_vs\_eat2, file = 'DEJ\_control\_vs\_eat2.csv')
# OPTIONAL step.

# cpmCountsDEJ saves CPM counts and log2CPM expression values for ranked differentially expressed junctions of a particular contrast (in csv format). cpmCountsDEJ function requires the file containing ranking for a contrast and junction matrix.

cpmCountsDEJ(JunctionMatrix, 'DEJ\_control\_vs\_eat2.csv')
# DEJ done

## FASE done