Artemis: Raw Reads To Pathway Analsyes In Much Less Time

Timothy J. Triche, Jr, Anthony R. Colombo, Harold Pimentel
15 January, 2016

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1 Introduction

Kallisto is software developed by Nicolas Bray, Harold Pimentel, Pall Melsted, and Lior Pachter (UC Berkeley) that analyzes 30 million unaligned paired-end RNA-Seq reads in less than 5 minutes on a standard laptop computer. Kallisto quantifies transcript abundance from input RNA-Seq reads by using a process, known as pseudoalignment, which identifies the read-transcript compatibility matrix. Artemis is a BioConductor package that extends functions and utilities for RNA-Seq analysis from raw reads to results in minutes.

2 Reads to Quantification to Annotation

Artemis was designed to reduce the programmative steps required to quantify and annotate multitudes of sample directories. Artemis calls Kallisto to perform on- the-fly transcriptome indexing and quantification recursively for numerous sample directories. For RNA- Seq projects with numerous sequenced samples, Artemis encapsulates expensive preparatory routines. Artemis programmatically orders FASTQ files output from DNA sequencers and inputs a list required by Kallisto for processing multitudes of demultiplexed reads. The Artemis function 'runKallisto' recursively indexes transcriptomes and quantifies abundances for any number of samples.

The function 'mergeKallisto' merges quantified output into an object of of subclass a KallistoExperiment-class, SummarizedExperiment-class, SummarizedExperiment-methods are preserved in KallistoExperiment-methods. Gene annotation is performed from user-selected bundled transcriptomes (ERCC, Ensembl, and/or RepBase) simultaneously merging annotated samples into one R object: KallistoExperiment. Artemis annotates genes for Homo-Sapiens GrCh38 and Mouse GrCm38 (NCBI). Routines such as 'annotateBundles' yields annotated genes from transcriptomes such as External RNA Control Consortium (ERCC), Ensembl release 81 of non-coding RNA, coding RNA, and a hg38 repeatome for both species.

2.1 Kallisto Installation

For linux systems, after installing the dependencies, kallisto is installed via:

```
mkdir /KallistoSource
cd /KallistoSource
git clone https://github.com/pachterlab/kallisto.git
cd ./kallisto
mkdir ./build
cd ./build
cmake ..
make
make install
```

3 Gene Wise Analysis

Artemis supports various levels of analysis, namely transcript-level or gene-level analysis which involves the Limma package for differential expression analysis.

Gene Wise Analysis is founded on the idea that groups of transcripts by a fixed Ensembl Gene ID is termed a "gene"; where "gene" counts are defined as the sum of all transcripts identified by the same unique Ensembl Gene Id. Gene Wise analysis generates bundled and aggregated transcripts associated with a specific Ensembl Gene ID. Artemis wraps limma around another method titled "collapseBundles", which collapses transcripts into appropriate groups and sums the quantified transcript counts of the group; these transcript aggregated counts are defined as "gene" counts.

3.1 The Measure Depends On The Level

Not all transcripts have the same function homology. Most folks agree that genes are made up by transcripts defined by the transcripts' coordinate location on the genome. However there are transcipt isoforms in DNMT3A and WT1 that have radically different biological function depending on the transcript isoform that is present. The problem with conducting *only* a gene level analysis is that many genes can have the same total gene level total quantified counts; however the biological mechanisms for the same "gene" can vary greatly by a single transcript isoform.

```
suppressWarnings(suppressPackageStartupMessages(library(artemis)))
suppressPackageStartupMessages(library(artemisData))
jsonFile <- system.file("extdata", "NS.JSON", package="artemis")
appSession <- fetchAppSession(jsonFile) ## a
names(appSession$samples) <- appSession$samples ## so column names get set
appSession$outputPath <- system.file("extdata", "", package="artemisData")
pathBase<-system.file("extdata",package="artemisData")</pre>
```

3.2 Creating The Design Matrix

In order to analyze bundle-aggregated transcripts defined as "genes", we create a design matrix which controls for individual effects and contrasts treatment effects across individual subjects.

```
##
      (Intercept) treatmentTRUE subject2 subject4
## n1
                                0
                 1
## n2
                 1
                                0
                                         1
                                                   0
## n4
                1
                                0
                                         0
                                                   1
                                         0
                                                   0
## s1
                                1
## s2
                                1
                                         1
                                                   0
                 1
## s4
## attr(,"assign")
## [1] 0 1 2 2
## attr(,"contrasts")
## attr(,"contrasts")$treatment
## [1] "contr.treatment"
## attr(,"contrasts")$subject
## [1] "contr.treatment"
```

4 Annotate!

In order to run gene-wise analysis, Artemis requires that the merged KallistoExperiment must be annotated; this is because we must collapse transcripts into groups linked to unique Ensembl Gene Ids.

4.1 Building Annotation libraries

Library Annotations are built using TxDbLite; these annotation databases allow for lite annotations parsing gene names, bio-types and family type from reference fastas from ERCC, Ensembl, or RepBase. Currently exonic, intronic, or other coordinate dependent information is not included in TxDbLite. The supplemental package artemisData stores the ready-to-load annotation libraries under /extdata/Libraries directory. For demonstration, we build the libraries under the artemisData/extdata/fasta/tmp directory.

```
suppressPackageStartupMessages(library(TxDbLite))
suppressWarnings(suppressPackageStartupMessages(library(artemis)))
suppressPackageStartupMessages(library(artemisData))
jsonFile <- system.file("extdata", "NS.JSON", package="artemis")</pre>
appSession <- fetchAppSession(jsonFile)</pre>
names(appSession$samples) <- appSession$samples</pre>
appSession$outputPath <- system.file("extdata", package="artemisData")</pre>
fastaPath<-system.file("extdata", "fasta", package="artemisData")</pre>
appSession$fastaPath<-fastaPath
cd<-appSession$fastaPath
setwd(paste0(appSession$fastaPath,"/","tmp"))
NS <- mergeKallisto(appSession$samples,
                      outputPath=appSession$outputPath)
## Setting transcriptome automatically from Kallisto call string.
fastaTx<-c("ERCC.fa.gz", "Homo_sapiens.GRCh38.81.cdna.all.fa.gz", "Homo_sapiens.RepBase.20_05.merged.fa")
erccDb<-erccDbLiteFromFasta(paste0(appSession$fastaPath,"/tmp/","ERCC.fa.gz"))</pre>
## Extracting spike-in associations...done.
## Creating the database...done.
## Writing the spike-in tables...done.
erccPkg<-makeErccDbLitePkg(erccDb,destDir=paste0(appSession$fastaPath,"/","tmp"))
## Creating package in /home/anthonycolombo/R/x86_64-pc-linux-gnu-library/3.2/artemisData/extdata/fasta
#Create a Ensembl Annotation Db with cdna and ncrna
lapply(fastaTx,function(x) findDupes(x))
## found no duplicated sequence names .... no dupes found
## found no duplicated sequence names .... no dupes found
## Warning in .Call2("fasta_index", filexp_list, nrec, skip, seek.first.rec, :
## reading FASTA file Homo_sapiens.RepBase.20_05.merged.fa: ignored 147
## invalid one-letter sequence codes
## found no duplicated sequence names .... no dupes found
## [[1]]
##
              duplicates
## ERCC.fa.gz
## [[2]]
##
                                          duplicates
## Homo_sapiens.GRCh38.81.cdna.all.fa.gz
## [[3]]
                                         duplicates
## Homo_sapiens.RepBase.20_05.merged.fa
```

```
ensDb<-ensDbLiteFromFasta("Homo_sapiens.GRCh38.81.cdna.all.fa.gz")</pre>
## Loading required package: Biostrings
## Loading required package: XVector
## Loading required package: org.Hs.eg.db
##
## Extracting transcript lengths...done.
## Extracting transcript descriptions...done.
## Extracting genomic coordinates...done.
## Extracting gene and biotype associations...done.
## Tabulating GC content...done.
## Tabulating transcript biotypes...done.
## Tabulating genes.....done.
## Creating the database...done.
## Writing the gene table...done.
## Tabulating gene biotypes...done.
## Writing the gene_biotype table...done.
## Writing the tx table...done.
## Tabulating transcript biotypes...done.
## Writing the tx_biotype table...done.
## Writing the biotype_class table...done.
ensPkg<-makeEnsDbLitePkg(ensDb,destDir=paste0(appSession$fastaPath,"/","tmp"))</pre>
## Creating package in /home/anthonycolombo/R/x86_64-pc-linux-gnu-library/3.2/artemisData/extdata/fasta
repDb<-repDbLiteFromFasta("Homo_sapiens.RepBase.20_05.merged.fa")
## Extracting repeat lengths...done.
## Extracting repeat descriptions...done.
## Creating the database...done.
repPkg<-makeRepDbLitePkg(repDb,destDir=paste0(appSession$fastaPath,"/","tmp"))
## Creating package in /home/anthonycolombo/R/x86_64-pc-linux-gnu-library/3.2/artemisData/extdata/fasta
ErccDbLite(erccDb)
## ErccDbLite :
## |package_name: ErccDbLite.ERCC.97
## |db_type: ErccDbLite
## |type_of_gene_id: N/A
## |created_by: TxDbLite 1.9.100
## |creation_time: Fri Jan 15 11:57:48 2016
## |organism: N/A
## |genome_build: N/A
## |source_file: /home/anthonycolombo/R/x86_64-pc-linux-gnu-library/3.2/artemisData/extdata/fasta/tmp/E
## | 97 spike-in controls from 4 subgroups (no known genes).
```

EnsDbLite(ensDb)

```
## EnsDbLite :
## |package_name: EnsDbLite.Hsapiens.81
## |db_type: EnsDbLite
## |type_of_gene_id: Ensembl Gene ID
## |created_by: TxDbLite 1.9.100
## |creation_time: Fri Jan 15 12:01:05 2016
## |organism: Homo sapiens
## |genome_build: GRCh38
## |source_file: Homo_sapiens.GRCh38.81.cdna.all.fa.gz
## | 175372 transcripts from 38530 bundles (genes).
```

RepDbLite(repDb)

```
## RepDbLite :
## |package_name: RepDbLite.Hsapiens.2005
## |db_type: RepDbLite
## |type_of_gene_id: RepBase identifiers
## |created_by: TxDbLite 1.9.100
## |creation_time: Fri Jan 15 12:01:05 2016
## |organism: Homo sapiens
## |genome_build: RepBase20_05
## |source_file: Homo_sapiens.RepBase.20_05.merged.fa
## | 1116 repeat exemplars from 68 repeat families (no known genes).
```

transcripts(ErccDbLite(erccDb))

```
## GRanges object with 97 ranges and 9 metadata columns:
                              ranges strand | tx_length gc_content
##
                  segnames
##
                     <Rle> <IRanges> <Rle> | <integer>
                                                         <numeric>
##
     ERCC-00002 ERCC-00002 [1, 1061]
                                                    1061 0.5136664
                                      * |
     ERCC-00003 ERCC-00003 [1, 1023]
                                         * |
##
                                                    1023 0.3264907
##
     ERCC-00004 ERCC-00004 [1, 523]
                                            523 0.3441683
##
     ERCC-00007 ERCC-00007 [1, 1135]
                                        * |
                                                   1135 0.4537445
     ERCC-00009 ERCC-00009 [1, 984]
##
                                        * |
                                                   984 0.4725610
##
            . . .
                       . . .
                                       . . . . . .
                                                    . . .
##
     ERCC-00165 ERCC-00165 [1, 872]
                                                    872 0.5000000
                                         *
                                            ##
     ERCC-00168 ERCC-00168 [1, 1024]
                                            1024 0.3417969
##
     ERCC-00170 ERCC-00170 [1, 1023]
                                            - 1
                                                    1023 0.3372434
     ERCC-00171 ERCC-00171 [1, 505]
                                            - 1
##
                                                    505 0.4772277
##
    ERCC_vector ERCC_vector [1, 2732]
                                         *
                                             2732 0.4989019
##
                             gene_id gene_name entrezid
##
                <character> <integer> <integer> <integer>
##
     ERCC-00002 ERCC-00002
                                <NA>
                                         <NA>
                                                   <NA>
##
     ERCC-00003 ERCC-00003
                                <NA>
                                         <NA>
                                                   <NA>
##
     ERCC-00004 ERCC-00004
                              <NA>
                                        <NA>
                                                   <NA>
                           <NA>
                                        <NA>
##
     ERCC-00007 ERCC-00007
                                                   <NA>
##
     ERCC-00009 ERCC-00009
                                        <NA>
                                                   <NA>
##
                               . . .
                                                   . . .
##
     ERCC-00165 ERCC-00165
                                <NA>
                                        <NA>
                                                   <NA>
     ERCC-00168 ERCC-00168
##
                                <NA>
                                         <NA>
                                                   <NA>
```

```
##
      ERCC-00170 ERCC-00170
                                    <NA>
                                               <NA>
                                                          <NA>
##
      ERCC-00171 ERCC-00171
                                     <NA>
                                               <NA>
                                                          <NA>
##
     ERCC vector ERCC vector
                                    <NA>
                                               <NA>
                                                          <NA>
##
                            tx_biotype gene_biotype biotype_class
##
                           <character>
                                        <character>
                                                        <character>
##
      ERCC-00002
                             SpikeIn D
                                             SpikeIn
                                                            SpikeIn
##
      ERCC-00003
                             SpikeIn D
                                             SpikeIn
                                                            SpikeIn
##
      ERCC-00004
                             SpikeIn_A
                                             SpikeIn
                                                            SpikeIn
##
      ERCC-00007 SpikeIn_unannotated
                                             SpikeIn
                                                            SpikeIn
##
      ERCC-00009
                             SpikeIn_B
                                             SpikeIn
                                                            SpikeIn
##
                                                 . . .
                                                                 . . .
##
      ERCC-00165
                             SpikeIn_D
                                             SpikeIn
                                                            SpikeIn
##
      ERCC-00168
                             SpikeIn_D
                                             SpikeIn
                                                            SpikeIn
                                                            SpikeIn
##
      ERCC-00170
                             SpikeIn_A
                                             SpikeIn
##
      ERCC-00171
                             {\tt SpikeIn\_B}
                                             SpikeIn
                                                            SpikeIn
##
     ERCC_vector SpikeIn_unannotated
                                             SpikeIn
                                                            SpikeIn
##
##
     seqinfo: 97 sequences from N/A genome; no seqlengths
```

transcripts(EnsDbLite(ensDb))

```
##
  GRanges object with 175372 ranges and 9 metadata columns:
##
                                 seqnames
                                                           ranges strand
##
                                    <Rle>
                                                       <!Ranges>
                                                                   <Rle>
     ENST00000000233
##
                                        7 [127588345, 127591705]
##
     ENST00000000412
                                       12 [ 8940365,
                                                        8949955]
                                       11 [ 64305578,
##
     ENST00000000442
                                                       64316738]
##
     ENST0000001008
                                       12 [ 2794953,
                                                         2805423]
##
                                        2 [ 72129238,
     ENST0000001146
                                                       72148038]
##
##
     ENST00000634217
                      CHR_HSCHR11_1_CTG7 [ 2963590,
                                                        2991183]
                      CHR_HSCHR15_4_CTG8 [ 28491559,
##
     ENST00000634219
                                                       28494348]
##
     ENST00000634220 CHR_HSCHR7_1_CTG4_4 [103099776, 103115340]
##
     ENST00000634221
                       CHR HSCHR8 9 CTG1 [ 39107956,
                                                       39151406]
##
     ENST00000634222
                                        6 [ 36754233,
                                                       36757400]
##
                     tx_length gc_content
                                                     tx id
                                                                    gene id
##
                                               <character>
                     <integer>
                                 <numeric>
                                                                <character>
                          1103 0.6092475 ENST00000000233 ENSG00000004059
##
     ENST00000000233
##
                           2756
                                 0.4746009 ENST00000000412 ENSG00000003056
     ENST00000000412
##
     ENST00000000442
                           2215
                                 0.6406321 ENST00000000442 ENSG00000173153
##
     ENST0000001008
                          3732
                                 0.5107181 ENST00000001008 ENSG00000004478
##
     ENST0000001146
                          4732
                                 0.5680473 ENST00000001146 ENSG00000003137
##
                           . . .
##
     ENST00000634217
                                 0.4528620 ENST00000634217 ENSG00000273562
                           594
##
     ENST00000634219
                           508
                                 0.5688976 ENST00000634219 ENSG00000278310
##
                                 0.3500000 ENST00000634220 ENSG00000275723
     ENST00000634220
                           4060
##
     ENST00000634221
                           571
                                 0.4238179 ENST00000634221 ENSG00000275594
                                 0.5077910 ENST00000634222 ENSG00000124772
##
     ENST00000634222
                           2182
##
                       gene_name
                                     entrezid
                                                        tx biotype
##
                                                        <character>
                      <character> <character>
##
     ENST00000000233
                             ARF5
                                          381
                                                    protein_coding
##
     ENST00000000412
                            M6PR
                                         4074
                                                    protein_coding
##
     ENST00000000442
                           ESRRA
                                         2101
                                                    protein coding
##
     ENST0000001008
                           FKBP4
                                         2288
                                                    protein_coding
```

```
##
     ENST0000001146
                         CYP26B1
                                        56603
                                                    protein_coding
##
                 . . .
                             . . .
                                          . . .
                                                    protein_coding
##
     ENST00000634217
                          NAP1L4
                                         4676
##
     ENST00000634219
                            <NA>
                                         <NA> processed_transcript
##
     ENST00000634220
                         NAPEPLD
                                       222236
                                                  retained intron
##
     ENST00000634221
                                       203102
                                                   protein coding
                          ADAM32
##
     ENST00000634222
                          CPNE5
                                        57699
                                                   retained intron
##
                                            gene_biotype biotype_class
##
                                             <character>
                                                            <character>
##
     ENST00000000233
                                          protein_coding protein_coding
##
     ENST00000000412
                                          protein_coding protein_coding
##
     ENST00000000442
                                          protein_coding protein_coding
##
     ENST0000001008
                                          protein_coding protein_coding
##
     ENST0000001146
                                          protein_coding protein_coding
##
##
     ENST00000634217
                                          protein_coding protein_coding
##
     ENST00000634219 transcribed_unprocessed_pseudogene
                                                          pseudogene
##
     ENST00000634220
                                          protein_coding protein_coding
##
     ENST00000634221
                                          protein_coding protein_coding
##
     ENST00000634222
                                          protein coding protein coding
##
##
     seqinfo: 288 sequences from GRCh38 genome; no seqlengths
```

transcripts(RepDbLite(repDb))

GRanges object with 1116 ranges and 9 metadata columns: ## seqnames ranges strand | tx_length gc_content ## <Rle> <IRanges> <Rle> | <integer> <numeric> ## HERVH HERVH [1, 7713] 7713 0.4602619 X21_LINE [1, 185] ## X21_LINE 185 0.3351351 ## UCON50 UCON50 [1, 133] 133 0.2105263 * -## Charlie22a Charlie22a [1, 491] 491 0.3808554 PrimLTR79 PrimLTR79 [1, 503] ## 503 0.4174950 ## SVA_E [1, 1382] ## SVA_E 1382 0.6099855 * 1 ## SVA F SVA F [1, 1375] * 1375 0.6094545 ## AluYb11 AluYb11 [1, 289] * -289 0.6332180 AluYb10 [1, 288] 288 0.6354167 ## AluYb10 -## AluYb8a1 AluYb8a1 [1, 287] 287 0.6376307 ## tx_id gene_id gene_name entrezid tx_biotype ## <character> <integer> <integer> <integer> <character> ## HERVH HERVH <NA> <NA><NA>ERV1 X21_LINE ## X21_LINE <NA> <NA> <NA> CR1 ## UCON50 UCON50 <NA> <NA> <NA> hAT <NA> <NA> <NA> ## Charlie22a Charlie22a hAT ## PrimLTR79 PrimLTR79 <NA> <NA> <NA> ERV1 ## ## SVA_E SVA_E <NA> SVA < NA ><NA> ## SVA F SVA F <NA> <NA><NA>SVA ## AluYb11 AluYb11 <NA> <NA> <NA> Alu ## AluYb10 AluYb10 <NA> <NA> <NA>Alu ## AluYb8a1 AluYb8a1 <NA> <NA> <NA> Alu ## gene_biotype biotype_class ## <character> <character>

```
##
          HERVH LTR_element
                                      repeat
##
       X21 LINE
                         LINE
                                      repeat
##
         UCON50 DNA_element
                                      repeat
##
     Charlie22a DNA_element
                                      repeat
##
      PrimLTR79 LTR element
                                      repeat
##
          SVA_E other_repeat
##
                                      repeat
##
          SVA_F other_repeat
                                      repeat
##
        AluYb11
                         SINE
                                      repeat
##
        AluYb10
                         SINE
                                      repeat
##
       AluYb8a1
                         SINE
                                      repeat
##
##
     seqinfo: 1116 sequences from RepBase20_05 genome; no seqlengths
files <-dir(paste0(appSession$fastaPath, "/tmp"))[!dir(paste0(appSession$fastaPath, "/tmp")) %in% fastaTx]
lapply(files,function(x) system(paste0("rm -r ",x)))
## [[1]]
## [1] 0
## [[2]]
## [1] 0
##
## [[3]]
## [1] 0
##
## [[4]]
## [1] 0
## [[5]]
## [1] 0
##
## [[6]]
## [1] 0
## [[7]]
  [1] 0
##
##
## [[8]]
```

5 Annotating Merged KallistoExperiment Containers

[1] 0

Artemis has a function "annotateFeatures.R" which annotates ERCC, Ensembl, and RepBase databases for species Homo-Sapiens, Mus-musculus, and Rattus norvegicus. The method "annotateFeatures.R" annotates the merged KallistoExperiment against every TxDbLite library simulatenously. These annotation databases are defined as 'lite' because they do not store exonic or intronic coordinates.

```
suppressPackageStartupMessages(library(artemis))
library(artemisData)
suppressPackageStartupMessages(library(TxDbLite))
```

```
samples<-c("n1","n2","n4","s1","s2","s4")</pre>
pathBase<-system.file("extdata",package="artemisData")</pre>
merged <- mergeKallisto(samples, outputPath=pathBase)</pre>
## Setting transcriptome automatically from Kallisto call string.
libraryPath<-system.file("extdata","Libraries",package="artemisData")</pre>
command<-paste0("sudo R CMD INSTALL ",libraryPath,"/",dir(libraryPath))</pre>
lapply(command,function(x) system(x))
## [[1]]
## [1] 0
## [[2]]
## [1] 0
## [[3]]
## [1] 0
merged <- annotate Features (merged, level="transcript") #annotate features using transcriptomes
## Loading required package: ErccDbLite.ERCC.97
## Loading required package: EnsDbLite.Hsapiens.81
## Warning in .Seqinfo.mergexy(x, y): The 2 combined objects have no sequence levels in common. (Use
     suppressWarnings() to suppress this warning.)
## Loading required package: RepDbLite.Hsapiens.2007
## Warning in .Seqinfo.mergexy(x, y): The 2 combined objects have no sequence levels in common. (Use
     suppressWarnings() to suppress this warning.)
NS<-suppressWarnings(annotateFeatures(NS,level="transcript"))
NS$subject <- factor(substr(colnames(NS), 2, 2))
NS$treatment <- substr(colnames(NS), 1, 1) == "s"
NS$ID <- NULL
design <- with(as(colData(NS), "data.frame"),</pre>
                  model.matrix( ~ treatment + subject ))
rownames(design) <- colnames(NS)</pre>
metadata(NS)$design <- design
#returns a KallistoExperiment at the gene level
GWA<-geneWiseAnalysis(NS,design=design,
                        how="cpm",
                        p.cutoff=0.05,
                        fold.cutoff=1,
                        read.cutoff=1,
                        species="Homo.sapiens")
```

Fitting bundles...

For the time being, only summing of bundles is supported

finding entrez IDs of top ensembl genes...

head(GWA\$limmaWithMeta,n=20)

```
logFC
##
                                AveExpr
                                                       P. Value
                                                                  adj.P.Val
                                                t
                   2.503936
                                         3.452477 1.909930e-03 3.964324e-02
## ENSG0000000938
                              3.4115048
## ENSG0000000971 -3.222736
                              1.4423090 -3.786282 8.123283e-04 2.105124e-02
## ENSG0000001630 -1.822014
                              6.4316184 -5.308421 1.483109e-05 9.036595e-04
## ENSG0000002822
                   3.750556
                              1.1385317 4.073160 3.850783e-04 1.189385e-02
## ENSG00000003137 -6.183849 -1.8495805 -6.703350 4.074361e-07 5.101421e-05
## ENSG0000003402
                   1.631505
                              8.2001798
                                        7.894025 2.238905e-08 5.078680e-06
  ENSG00000004478 -2.185685
                              3.8978332 -3.916894 5.789056e-04 1.637483e-02
## ENSG0000005187 -1.943089
                              4.9011626 -4.206421 2.715437e-04 9.147383e-03
## ENSG0000005381
                    2.471777
                              6.5678349
                                        7.247126 1.059470e-07 1.745777e-05
## ENSG0000005810 -1.625765
                              7.8181754 -6.920968 2.366541e-07 3.280570e-05
                   2.713230
                                        4.968384 3.634861e-05 1.889533e-03
  ENSG00000005844
                              4.3581900
## ENSG0000006062
                   2.943912
                              4.2249563
                                        5.153493 2.230152e-05 1.254523e-03
                   3.585798 -0.3106464
                                        3.415789 2.095688e-03 4.219900e-02
## ENSG00000006118
## ENSG0000006125 -1.112240
                              6.9416204 -3.925823 5.656129e-04 1.609003e-02
## ENSG0000006831 -2.612343
                              5.8762788 -6.680104 4.319203e-07 5.350451e-05
## ENSG0000007202
                    1.075648
                              6.5488238
                                         3.382974 2.276585e-03 4.495644e-02
## ENSG0000007237
                    3.324875
                              3.7464312
                                        5.593566 7.022544e-06 5.153751e-04
  ENSG00000007384 -4.976736
                             -1.0944191 -4.491820 1.280544e-04 5.177461e-03
                             4.0822873 4.425604 1.525006e-04 5.880027e-03
## ENSG0000008130
                   2.300708
  ENSG00000008283 -2.560269
                              3.9195730 -4.418107 1.555460e-04 5.944953e-03
                            B entrez_id gene_name
                                                       ensembl_id
##
                                   2268
  ENSG00000000938 -1.4095287
                                              FGR ENSG00000000938
  ENSG00000000971 -0.5100927
                                   3075
                                              CFH ENSG00000000971
  ENSG0000001630
                    2.8192731
                                   1595
                                          CYP51A1 ENSG0000001630
## ENSG0000002822
                    0.1513972
                                   8379
                                           MAD1L1 ENSG00000002822
  ENSG00000003137
                    6.1313765
                                  56603
                                          CYP26B1 ENSG00000003137
## ENSG0000003402
                   9.0162886
                                   8837
                                            CFLAR ENSG00000003402
## ENSG00000004478 -0.3685884
                                   2288
                                            FKBP4 ENSG00000004478
## ENSG0000005187
                    0.2188737
                                   6296
                                            ACSM3 ENSG00000005187
## ENSG0000005381
                    7.7419880
                                   4353
                                              MPO ENSG00000005381
## ENSG0000005810
                    6.6960183
                                  23077
                                           MYCBP2 ENSG00000005810
                                            ITGAL ENSG00000005844
## ENSG0000005844
                    2.2455964
                                   3683
## ENSG0000006062
                    2.7250540
                                   9020
                                          MAP3K14 ENSG0000006062
## ENSG00000006118 -1.3272816
                                  54972
                                         TMEM132A ENSG00000006118
## ENSG00000006125 -0.8679923
                                    163
                                            AP2B1 ENSG00000006125
## ENSG0000006831
                   6.4144336
                                  79602
                                          ADIPOR2 ENSG00000006831
## ENSG0000007202 -2.1223845
                                   9703
                                         KIAA0100 ENSG00000007202
                    3.8426315
## ENSG0000007237
                                   8522
                                             GAS7 ENSG00000007237
  ENSG00000007384
                    1.0616307
                                  64285
                                           RHBDF1 ENSG00000007384
                                             NADK ENSG00000008130
  ENSG00000008130
                                  65220
                    0.8776605
##
   ENSG00000008283
                    0.8797450
                                   1534
                                           CYB561
                                                  ENSG00000008283
##
                     gene_biotype
                                  biotype_class
## ENSG0000000938 protein_coding protein_coding
  ENSG00000000971 protein_coding protein_coding
  ENSG0000001630 protein_coding protein_coding
## ENSG00000002822 protein_coding protein_coding
```

```
## ENSG00000003137 protein_coding protein_coding
## ENSG0000003402 protein_coding protein_coding
## ENSG0000004478 protein coding protein coding
## ENSG0000005187 protein_coding protein_coding
## ENSG00000005381 protein coding protein coding
## ENSG00000005810 protein coding protein coding
## ENSG00000005844 protein coding protein coding
## ENSG0000006062 protein coding protein coding
## ENSG0000006118 protein coding protein coding
## ENSG0000006125 protein_coding protein_coding
## ENSG0000006831 protein_coding protein_coding
## ENSG0000007202 protein_coding protein_coding
## ENSG0000007237 protein_coding protein_coding
## ENSG0000007384 protein_coding protein_coding
## ENSG00000008130 protein_coding protein_coding
## ENSG00000008283 protein_coding protein_coding
```

6 Gene Wise Analysis

Gene wise analysis collapses transcripts into groups related to specific ensembl "gene" Ids. The package TxDbLite parses the Ensembl, or RepBase transcript fasta files and stores the respective gene id's associated with the given transcript documented in the transcript fasta header. Artemis' method for gene wise analysis calls "collapseBundles.R" which then calculates the aggregated total counts of transcripts for each unique gene id association. Thus the "gene" count is defined as the sum of all quantified transcripts associated with a specific gene identifier.

6.1 Understanding Gene Wise Analysis Output

The output contains a list of limma derived expression values, and enrichment data derived by biomaRt. ##Expression Results

The expression results were generated by limma/voom and have the meta biotype, gene name, etc information included in the gene wise analysis results.

6.2 Understanding Gene Wise Analysis Output

The output contains a list of limma derived expression values, and entrezID, gene name, and gene biotypes derived by biomaRt and TxDbLite respectively. The expression results were generated by limma/voom and have the meta biotype, gene name, etc information included in the gene wise analysis results.