NGS, from data to table to figure

From downloading data to final publication ready figures

Background

• I will introduce what I learned from mine supervisors these years. This report also include some knowledge which I learn by myself.

Object

- 1. I hope everyone can follow me, and I ensure you will meet these problems if you do NGS analysis or data analysis.
- 2. If you have any question at any time, please **don't hesitate** to ask a question, actually some of the materials are very difficult to me, and I also learn it while this class is running.
- 3. Although it may take some time to learn, but it will save a lot of time in the future.
- 4. I hope after each class, you can make some practice and read the supplement files.

Object

- 5. All the course martials can be download from https://github.com/regSNPs/from_data_2_table_2_figure
- 6. This is not a advance class, but I confirm you will learn some 'latest knowledge', if you search R question in stackoverflow you will see a lot of question/answers use the knowledge here.
- 7. I think after the report, the students can know how to perform data analysis when supervisor give us a task.

Components of the report

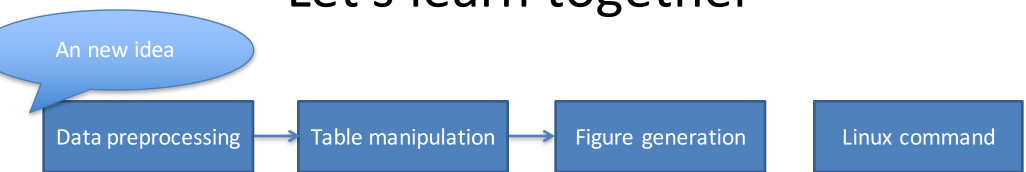
• 1. Data preprocessing (First we need to know where we can got the data and how to preprocess the data)

• 2. Table manipulation (We usually don't use the raw data directly, we need process them into a desired format for further investigation)

• 3. Figure generation (Human beings can't read thousands of rows in table simultaneously, we need figure to visualize them)

• 4. Linux (Besides R, we also need Linux to facilitate some work)

Let's learn together



The required knowledge include:

Biology (understand the biology process of the problem)

Programming skill (understand the code)

Statistic graphic (understand the figure)

The example problem

We will follow a problem during the course. Problem to solve: RNA binding protein binding gene analysis.

We want to study the relationship between multi RBPs and gene expression. i.e the object is to study if the number of RBPs bind the gene can affect the gene expression level in different genes in same cell line.

The example problem

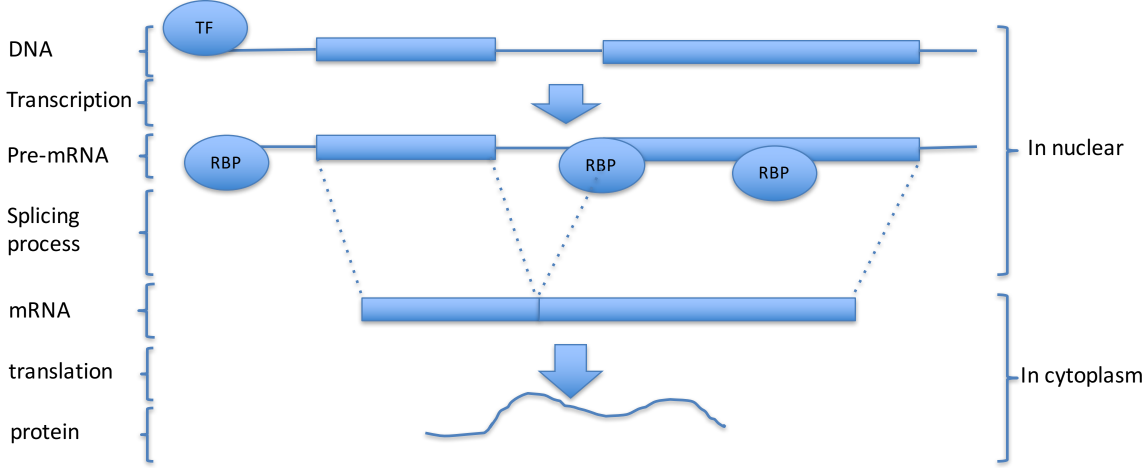
 We can also study the RBP and gene expression analysis in different cell lines, but I didn't have that data currently. It needs NGS data for each cell line.

The biology process of this

Although I'm not very good at biology, but the first thing (most important) is understanding the biology process of the problem. which will make the analysis meaningful.

I will try to explain the biology process of this problem first.

The biology process of gene expression



We all know the TF can affect the gene expression, but will the RBPs in a gene has some correlation with gene expression? I don't know either, let's do it!!

The analysis process

- 1. We need data to know where the RBP binds? This kind of data can be got from **CLIP-seq technique**.
- 2. We need to know how the gene expressed in a cell line. This data can be got from RNA-seq technique.
- 3. We need to ensure that both two kinds of dataset come from the same cell line. After we got the above information, our final analysis is doing correlation between RBP binding count and gene expression.

The CLIP-seq technique

First we need to know where the RBP binds? The below is a demonstration of RBP binding region, in reality, we don't know where the RBP binds.



The technique is called CLIP-seq, which can locate the target of RBP binding sites in the pre-mRNA.

The CLIP-seq experiment

Got from Wikipedia

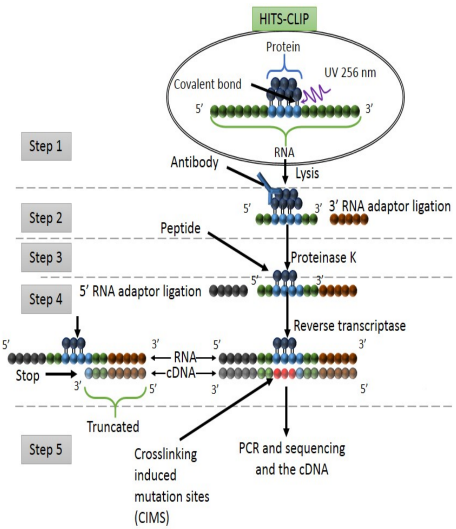


Figure 2: HITS-CLIP

Step 1

HITS-CLIP begins with the in-vivo cross-linking of RNA-protein complexes using ultraviolet light. The cell is lysed and the protein of interest is isolated using immunoprecipitation.

Step 2

Washing is performed to remove free RNA, and RNA adaptors are ligated at the 3' ends.

Step 3

Proteinase K digestion is performed. This leaves a peptide at the cross-link site that modifies the chemical structure of the nucleotide.

Step 4

5` RNA adaptors are ligated and cDNA is synthesized using reverse transcription.

Step 5

PCR and sequencing of the cDNA.

CLIP-seq workflow

It is a technique just like the CHIP-seq, whereas work on premRNA and RBPs.

The goal of the technique is extracting the sequence in the RBP binding sites.

We just need to know that the reads we will see later come from the binding sites.

I will show you the standard protocol later.

Sequence Database and annotation database

 There are two kinds of biology database. We need sequence database to get the sequence and annotation database to get gene annotation.

• 1. The sequence database store the sequence from sequencer, this kind of database is very very large.

• 2. The annotation database stores the genome annotation.

The sequence database

 Once we decide the data (CLIP-seq dataset) we need. Next step is selecting the database that we are going to extract the data.

 There are various kinds of sequence database, eg: Encode, modEncode, GEO, SRA, EBI, GTEX, TCGA, 1000 Genome Project, ESP6500

• I will give a brief introduction for these databases.

The sequence database

- Encode: DNA, RNA and other sequence data for HUMAN.
- modEncode: DNA,RNA and other sequence data for Model organism.
- GEO: gene chip and sequence database.
- SRA: sequence database.
- EBI: European mirror for SRA.
- GTEx: Genotype and Phenotype sequence database.
- TCGA: Tissue specific cancer sequence database.
- 1000 Genome Project: Thousands of people's DNA and RNA sequence.
- ESP6500: 6500 people's Exon sequencing data.

The annotation databases

- There are many annotation database available which include: UCSC, refseq, and Ensembl, Uinprot, PDB et.al.
- I will give a brief introduction to these databases.

- Usually, refseq is more accurate while Ensembl is more comprehensive.
- For refseq hg19 (GRCh37), it contains around 50000 transcripts.
- For ensembl hg19 (GRCh37), it contains around 200000 transcripts.

The annotation databases

- Uniprot: Protein sequence and functional annotation.
- PDB: Protein 3D structure and functional annotation.
- Pfam: Protein Domain information.
- NAR database index: http://www.oxfordjournals.org/our_journals/nar/database/c/
- Bioiformatics webserver directory
- https://bioinformatics.ca/links_directory/
- NAR webserver index:
- http://www.oxfordjournals.org/our_journals/nar/webserver/c

Encode&UCSC

- Encode
- Encode dataset matrix:
- UCSC's encode end with 2012
- <a href="https://www.encodeproject.org/matrix/?type=Experiment&x.limit="https://www.encodep

- UCSC
- Table browser, download various kinds of annotation from different assemble.

Encode&UCSC database

- I will show you the two databases.
- One is the encode database, another is the UCSC database.
- For encode, I will show how to get CLIP data for SRSF1 in K562.
 - (SRSF1.web)https://www.encodeproject.org/experiments/ ENCSR432XUP/
- For UCSC, I will show how to get the annotation, which will be used in various kinds of NGS analysis.
 (e.g. How to use the table browser, how to use download)

UCSC

- Besides the web interface, UCSC also provide two kinds of methods to extracting data from their database.
- 1. DAS server_{E.g.http://genome.ucsc.edu/cgi-bin/das/hg19/dna?segment=chr4:35654,35695}
- 2. mysql client:

Users can also use various kinds of mysql client (e.g. mysql workbench, JDBC et.al.)

Mysql -user=genome -host=genom-mysql.cse.ucsc.edu -A

Data formats

The download eCLIP data from Encode can be many different formats.

Including: fastq format (original reads), bam format (alignment file), bed format (peaks region file).

Here we assume the raw format is fastq format, I will show you the above formats. (UCSC format help:

http://genome.ucsc.edu/FAQ/FAQformat.html

Data preprocessing workflow Fastq Bam Bed

From fastq to bam

• The main command here is **read alignment**.

```
e.g bowtie2 [options]* -x <bowtie-index> {-1 read1 -2 read2 | -U
<r>> [-S <sam>]
```

- But there are possibly other steps, eg:
- 1. Demultiplexing (sometimes, multi samples are sequenced once a time, multiplexing is a technique to index them)
- 2. Cutadaptor (adaptor is used to do sequencing, we need to remove the adapters from head or tail of the adaptor)
- 3. FastQC (quality control)
- 4. rmRep (remove repetitive elements)

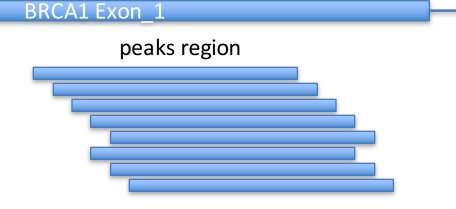
The BAM file we got

A bam file looks like below:

```
chr1 13232 33421 ATAAFCFAD &%dfd2$@$ ... chr2 61323 62121 ATCTTGCGT *&dfd2)a# ...
```

••• •

The bam file can be viewed through IGV viewer. I will show you an IGV image.



Quality control

 Usually, we need to check the quality of the result bam file.

- The possible criterion include:
- Distribution of the quality scores.
- % of reads mapped to the genome.
- % of reads mapped to the exon region.
- % of reads mapped to the positive strand.
- % of reads mapped to the negative strand.

From bam to bed

- Next we need to call peaks in the bam file. It extracted the regions that contain the reads.
- The main command is the Clipper.

- It also include:
- Samtools sort (sort the bam file by position)
- Samtools index (index the bam file to do fast retrieve)
- BigbedTobed (convert Bigbed to bed file)

Alignment and Peak Callers

• For sequence alignment, the candidate softwares include: BWA, Bowtie, Bowtie2, Star et.al. A comprehensive comparison of these tools can be checked at "Evaluation of Alignment Algorithms for Discovery and Identification of Pathogens Using RNA-seq";

 (When I leave Liu lab, our lab just start to use star aligner)

Alignment and Peak Callers

- For peak callers, dozens of algorithms exist, if you are interest in them, you can search it on the web. Like alignment software, these kinds of papers are usually hard to read.
- Both alignment and peak caller need scientist with strong background in **statistic** and other inter - displine skills (Including biology, computer science et.al).

The full pipeline here

• eCLIP_analysisSOP_v1.P.pdf. (the computation pipeline)

eCLIP_SOP_v1.P_110915.pdf. (the experiment pipeline)

• I will show you the standard analysis pipeline in the supplement file.

The data we got (SRSF_rep1.bed)

| chr1 15212 | 15250 | ENSG00000227232.4_0_3 | 15 | - | -1 | - | 0.0292794233248 | 30 |
|-------------|--------|------------------------|-----|---|----|----|-------------------|--------|
| chr1 16239 | 16287 | ENSG00000227232.4_1_4 | 19 | - | -1 | - | 0.0104649442093 | :63 |
| chr1 16441 | 16485 | ENSG00000227232.4_2_5 | 25 | - | -1 | - | 0.00256249972915 | 62 |
| chr1 17451 | 17517 | ENSG00000227232.4_3_10 | 63 | - | -1 | - | 4.50322291742e-07 | 17481 |
| chr1 90235 | 90280 | ENSG00000239945.1_0_4 | 19 | - | -1 | - | 0.0105097488596 | .56 |
| chr1 90240 | 90275 | ENSG00000238009.2_0_4 | 20 | - | -1 | - | 0.00879079737544 | .56 |
| chr1 109158 | 109192 | ENSG00000238009.2_1_5 | 29 | - | -1 | - | 0.00115927397829 | 175 |
| chr1 113824 | 113881 | ENSG00000238009.2_2_8 | 49 | - | -1 | - | 1.21214630854e-05 | 113848 |
| chr1 115708 | 115784 | ENSG00000238009.2_3_22 | 145 | - | -1 | - | 2.6562287759e-15 | 737 |
| chr1 116372 | 116428 | ENSG00000238009.2_4_13 | 74 | - | -1 | - | 3.4175167505e-08 | 397 |
| chr1 135196 | 135226 | ENSG00000237683.5_0_3 | 15 | - | -1 | - | 0.0270886550409 | 213 |
| chr1 135196 | 135226 | ENSG00000268903.1_0_3 | 15 | - | -1 | -: | 0.0274502908066 | 213 |

Column 4 is the Ensembl gene id. The gene where the peaks locate.

Each row is a peak region. **Now we know where the RBP binds!** Column 9 is the **p-values** (The probability of see the reads here when the region is not peak). We can filter some peaks using p-values.

The p-value here

 The p-value is the probability that the observed result has nothing to do with what one is actually testing for. Specifically, the p-value is defined as the probability of obtaining a result equal to or "more extreme" than what was actually observed, assuming that the model (null hypothesis) is true.

Prob(data|HO)

The p-value here

 The p-values are got from statistic hypothesis test and they almost everywhere in today's journals.

The quality scores as we discussed above is also
 10log(p-value). Include the each nuclide acid's quality, and the alignment quality.

Controlling the FDR

 When there are multi test in one experiment, like the example above (The peaks we got above). We will usually need FDR to avoid the multi-test problem.

- For example, if we have 10000 peaks called and select 0.05 as the p-values' cutoff. Although the type I error is 0.05, but the there are still 0.05*10000=500 is wrong.
- One solution is correcting the p-value's cutoff by dividing the samples size.

Basic quality control

 Usually, we need to check the file to ensure the file we got is right, this can be done by basic Linux commands.

- Some possible criterion is:
- The number of peaks in the files.
- How many chromosomes in the file.
- The number of peaks in the chromosome 1.
- The distribution of p-values for the peaks.

Most used Linux command

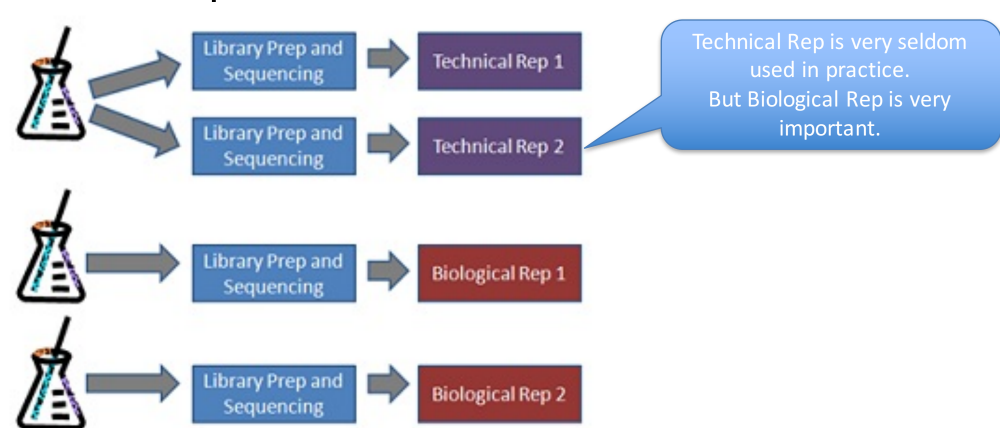
```
#Count how many peaks in the file:
wc -1 SRSF1 rep1.bed; (39116)
#Count how many peaks in chr12:
cut -f 1 SRSF1 rep1.bed | grep chr12 | wc -1 (2133)
#Get the p-value column:
cut -f 9 SRSF1 rep1.bed > SRSF1 rep1 pvalue.txt
#how many chromosome in it
cut -f 1 SRSF1 rep1.bed | sort -u | wc -l
```

Most used Linux command

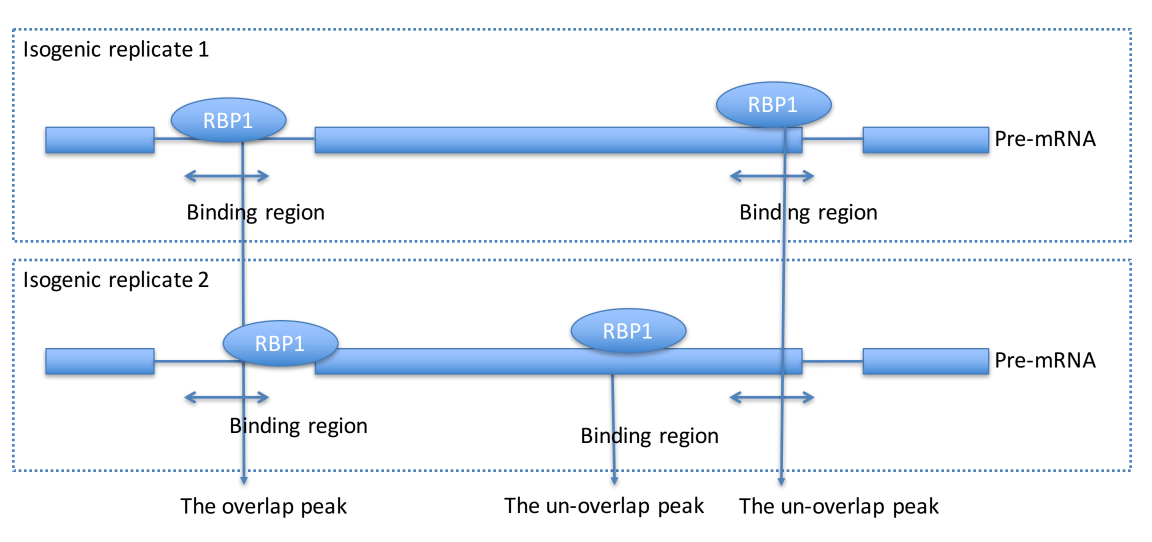
```
#remove the 'chr' in each line
cut -c 4- SRSF1_rep1.bed >SRSF1 rep1 noChr.bed
#Got the top 20 longest peaks in the bed file.
awk '{print $3-$2}' SRSF1 rep1 noChr.bed| sort -unr | head -n 20 (314)
#count the peaks in each of the file and output to a new file
for i in `ls *.bed`
do
   `wc -l $i >> peaks for each file.txt`;
done;
```

Biology replicates

 Usually we use at least one biological replicate to ensure the peaks we found are not due to variance.



Biology replicates



Overlap of two biological replicates

 One possible solution is finding the overlapped regions between the two biology replicates, and treat the overlapped region as peaks.

 There are also advance statistic model to do this, but we won't dive into it.

 But how to find the overlapped regions between two bed files?

The two bed files

Bed_rep_1

| chr1 14924 | 14954 | ENSG00000227232.4_0_3 | 15 - | -1 - | 1 | 0.0286089808565 14938 |
|------------|-------|------------------------|------|------|---|-------------------------|
| chr1 17454 | 17567 | ENSG00000227232.4_1_15 | 95 - | -1 - | 1 | 2.52804288143e-10 17488 |
| chr1 89861 | 89871 | ENSG00000238009.2_0_3 | 16 - | -1 - | 1 | 0.020756022461 89862 |
| chr1 89861 | 89871 | ENSG00000239945.1_0_3 | 16 - | -1 - | 1 | 0.0211020769049 89862 |
| chr1 89871 | 89904 | ENSG00000238009.2_1_3 | 16 - | -1 - | 1 | 0.0246792521386 89885 |
| chr1 89871 | 89904 | ENSG00000239945.1_1_3 | 15 - | -1 - | 1 | 0.0258746724881 89885 |

Bed_rep_2

| chr1 15212 | 15250 | ENSG00000227232.4_0_3 | 15 | - | -1 | -1 | 0.0292794233248 15230 |
|------------|-------|------------------------|----|---|----|----|-------------------------|
| chr1 16239 | 16287 | ENSG00000227232.4_1_4 | 19 | - | -1 | -1 | 0.0104649442093 16263 |
| chr1 16441 | 16485 | ENSG00000227232.4_2_5 | 25 | - | -1 | -1 | 0.00256249972915 16462 |
| chr1 17451 | 17517 | ENSG00000227232.4_3_10 | 63 | - | -1 | -1 | 4.50322291742e-07 17481 |
| chr1 90235 | 90280 | ENSG00000239945.1_0_4 | 19 | - | -1 | -1 | 0.0105097488596 90256 |
| chr1 90240 | 90275 | ENSG00000238009.2_0_4 | 20 | - | -1 | -1 | 0.00879079737544 90256 |

The general problem of overlapping Here we need to find the overlap between the two bed files, actually sometimes we also need to overlap different kinds of files.

The most common file formats we encountered are: BAM, BED, GTF and VCF.

They are all interval range formats, i.e. they all contain a chromosome name and at least a start position. A generic solution to this problem is need.

Find overlap of biology replicates

- R package GenomicRanges:
- Bed file is table range format, also some other file formats like VCF, GFF and GTF. We need efficient methods to overlap between these kinds of files.
- 1. One Linux solution is using Tabix, while a java solution a tribble.
- 2. A more powerful solution is using **R** package **GenomicRanges**. It can efficiently map between ranges format files. GenomicRangesIntroduction.pdf

VCF, BED, GTF file format

• I will show you the above VCF, BED and GTF file formats in UCSC, and show that they are all the general chromosome, coordinate format.

https://genome.ucsc.edu/FAQ/FAQformat.html#format10.1

R package GenomicRanges

To use it, first we need to read the file (either Bed, VCF, GFF or GTF) and then convert them into **GenomicRange** objects. Reference:

```
Codes:
```

```
#First read bed file into table biology_rep_:
SRSF1 rep 1 bed<-read.table("Bed rep 1")</pre>
```

Bed format is 0-based and its left coordinate is inclusive, while the right coordinate is exclusive

```
SRSF1_rep_1_range<-with(SRSF1_rep_1_bed, Granges(seqnames=chr,
+ranges=IRanges(start=start+1,end=end),strand=strand)</pre>
```

```
SRSF1_rep_2_range<-with(SRSF1_rep_2_bed,Granges(seqnames=chr,
+ranges=IRanges(start=start+1,end=end),strand=strand)</pre>
```

R package GenomicRanges

Find the peaks which overlap between the two biology replicates.

```
Codes:
##overlap the two biological replicates.
SRSF1 overlap mch<-findoverlaps(SRSF1 rep 1 range,
SRSF1_rep_2_range);
##get the overlap in replicate 1.
SRSF1 rep 1 overlap<-
SRSF1 rep 1 range[unique(queryHits(SRSF1 overlap mch)),];
##get the overlap in replicate 2.
SRSF1_rep_2_overlap<-
SRSF1 rep 2 range[unique(subjectHits(SRSF1 overlap mch)),];
                                                                   45
```

What the overlap result looks like

What we got currently

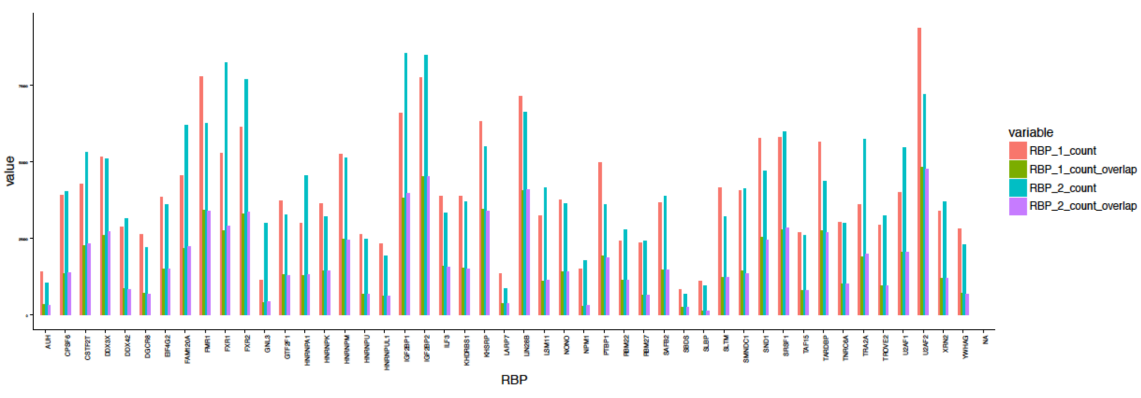
What we got is the overlapped peak regions between the two biological replicates.

How the replicates influence the result

How much the biological replicates influence the peaks region? The details of below program will be elucidate later.

```
##The number of overlapped peaks in each replicate.
overLapNum1<-length(unique(queryHits(oneMtch)));</pre>
overLapNum2<-length(unique(subjectHits(oneMtch)));</pre>
##combine the information
RBPs_replicate_overlap_frame<-c(rbpsnum1,overLapNum1,rbpsnum2,</pre>
overLapNum2)
##melt the data frame
RBPs replicate_overlap_melt<-
melt(RBPs replicate overlap frame,id.vars="RBP");
##plot
p<-ggplot(RBPs_replicate_overlap_melt, aes(x=RBP,</pre>
y=value, fill=variable)) + geom_bar(stat="identity", position="dodge")
```

How many peaks are overlapped between the two biological replicates



Only half of the peaks are overlapped between the two biological replicates,, so you can check that biological replicates is essential to this study, as far as I know, almost all the RNA-seq and CLIP data in ENCODE project have biological replicates. I think Chip-seq will also have biological replicates in it.

R package GenomicRanges

Besides overlap between two genome interval regions. GenomicRanges also include many additional functions. These include:

```
Set operations: ()
  union, setdiff, intersect;
Interval operation: (resize the interval, change the flank region)
  resize, flank, width, shift;
Overlap operations: (overlap with a region)
  countOverlaps, subsetByOverlaps, findOverlaps;
Split operations:split, c(); split a GenomicRanges into multi ones.
Basic operations:tail, head, rev; get subset of the genomicRanges
readGAlignments . Read bam files
                                                                   50
```

Multi RBPs CLIP-seq bed files

Sometimes, we need to analysis two or more RBPs to study their interaction. We need **GRangesList** structure. Here we take HNRNPK as example.

```
##Get the overlap regions of two HNRNPK replicates
HNRNPK_rep_1_overlap<-
HNRNPK_rep_1_range[unique(queryHits(HNRNPK_overlap_mch)),];

##Combine HNRNPK and SRSF1 into a single GRangesList object
RBPs_range_list<-GRangesList(SRSF1_rep_1_overlap,
HNRNPK_rep_1_overlap);</pre>
```

Count RBP binding sites in each gene

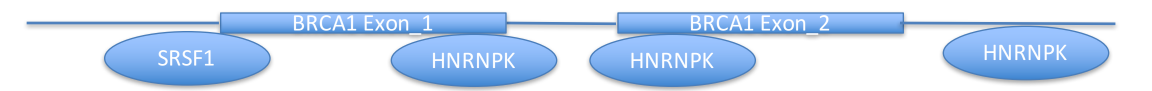
GRangeList object. Next, we want to count RBP binding sites in each transcript and ensure the information we got is enough to do further analysis. We need human genome annotation, here we use hg19_GTF which can be download GTF is 1-based system and both the left and right coordinate are inclusive.

Browser. File: Table Browser

| 1 | pseudogene | gene | 11869 | 14412 | • | + . | gene_id "ENSG00000223972"; |
|---|----------------------|------------|-------|-------|---|-----|-------------------------------|
| 2 | processed_transcript | transcript | 11869 | 14409 | • | + . | gene_id |

....

Count RBP binding sites in each gene



Example: BRCA1 contain 3 HNRNPK binding sites and 1 SRSF1 binding sites.

```
##read the GTF file into R
gtf_frame<-read.table("hg19_ens.gtf",header=F,as.is=T);
##construct a GenomicRange object for the GTF
gtf_range<-with(gtf_frame,Granges(seqnames=chr,</pre>
```

+ranges=IRanges(start=start,end=end),strand=strand)

Codes:

Overlap with RBPs binding sites and genome annotation

Next we overlap the RBPs_range_list with gRBP_range to get number of RBPs in each gene.

```
Code:
GeneName_RBPName<-data.frame(gene_name,RBP_name) #store the result
for(i in 1:2){##for each RBP.
    RBP_overlap_mtch<-
        findOverlaps(RBPs_range_list[[i]], gtf_range);#RBP overlap GTF
    RBP_gtf_overlap<-
        RBPs_range_list[[i]][queryHits(RBP_overlap_mtch),];#overlapped RBP

    Gtf_RBP_overlap<-
        gtf_range[[1]][subjectHits(RBP_overlap_mtch),];#overlapped GTF</pre>
```

Overlap with RBPs binding sites and genome annotation

```
Code:
    gene_names<-seqnames(GRBP_RBP_overlap);#overlapped RBP names</pre>
    RBP names<-names(RBPs range list)[i];#overlapped gene names
    GeneName_RBPName<-</pre>
       cbind(GeneName_RBPName,c(gene_names, RBP_names) );#store the result
>GeneName RBPName
#each line represent a overlap
   Gene names
                     RBP names
   TP53
                     SFRS1
   TP53
                     SFRS1
   TP53
                     HNRNPK
```

• • • •

An brief intro to Bioconductor

- R package GenomicRanges is part of Bioconductor project, which contains many packages related to bioinformatics, including NGS analysis, microarray analysis and so on.
- Besides the GenomicRanges, another general used package I found very useful is the BSGenome, which provide different annotation for many genomes.
- There are also other packages in it and they target on different aspect, like Chip-seq, RNA-seq (EdgeR) and so on.
- I links two papers which describe the Bioconductor packages in the supplement files.

Bioconductor

Many papers are published in Bioconductor by successfully build a package in it for specific use.

I suggest use Rstudio's package build helper to create your own package, very powerful, and a powerful tool is **devtools**.

Build a powerful package need strong IT skill, but biology and statistic knowledge is also required.

R Package reshape2

Next we need to do transformation to count each gene's binding sites.

Here, we first introduce the long table format and short table format.

Long table and short table formats

Short table format

| Gene_name | RBP_name |
|-----------|----------|
| TP53 | HNRNPK |
| TP53 | HNRNPK |
| BRCA1 | SRSF1 |
| TP53 | HNRNPK |
| TP53 | SRSF1 |

As you can see,
The data in the short
table is redundant. I
use more cell to store
the same data.

Long table format

| Gene_name | HNRNPK | SRSF1 |
|-----------|--------|-------|
| TP53 | xxx | xxx |
| BRCA1 | xxx | xxx |

Column name become header

The tidy data

• Usually, we think long table format is more tidy than short table format.

 A tidy table means each row is a observation and each column is a variable.

 Tidy data is very important to make our analysis clean and efficient. As we know, R's data.frame is tidy object.

R package reshape2

R package reshape 2 contains two fundamental function:

melt: Convert long table to short table.

cast: Convert short table to long table.

R package reshape2

```
>GeneName_RBPName
#each line represent a overlap
   Gene names
                 RBP_names
   TP53
                 SFRS1
   TP53
                  SFRS1
   TP53
                  HNRNPK
Codes:
#convert the short table format to long table format
GeneName RBPName cast<-
Cast(GeneName RBPName melt, Gene names~RBP names, length)
#Gene names~RBP names is folumar, where the left side is #row id and
left side is column label
#Where length is the function to count.
```

62

R package reshape2

>GeneName_RBPName_cast

| Gene_name | HNRNPK | SRSF1 | | | | | |
|-------------------------------|--------|-------------|----------|--------|--------|--------|-------|
| TP53 | 3 | 1 | | | | | |
| BRCA1 | 1 | 3 | | | | | |
| #Now we have for each RBP. | | we need, we | got each | gene's | RBP #b | inding | sites |
| # | | | | | | | |
| # | | | | | | | |
| # | | | | | | | |
| # | | | | | | | |

From table to figure

Human beings can't read hundreds of rows simultaneously, we need visualization method to check the dataset.

We now have the table, we want to represent it using figures to do visualization. Maybe it is not very useful for small table, but for large table, it means a lot.

We need R package ggplot2 to convert table to figure.

R package ggplot2

ggplot2 usually receive short table format data and convert it into statistic figures. Its main advantage is productive and high versatile. It make use of graphic grammar.

```
Codes:
```

#ggplot2 receive short table format, we need to melt the #table.

>GeneName_RBPName_cast

| Gene_name | HNRNPK | SRSF1 |
|-----------|--------|-------|
| TP53 | 3 | 1 |
| BRCA1 | 1 | 3 |

melt long table to short table format

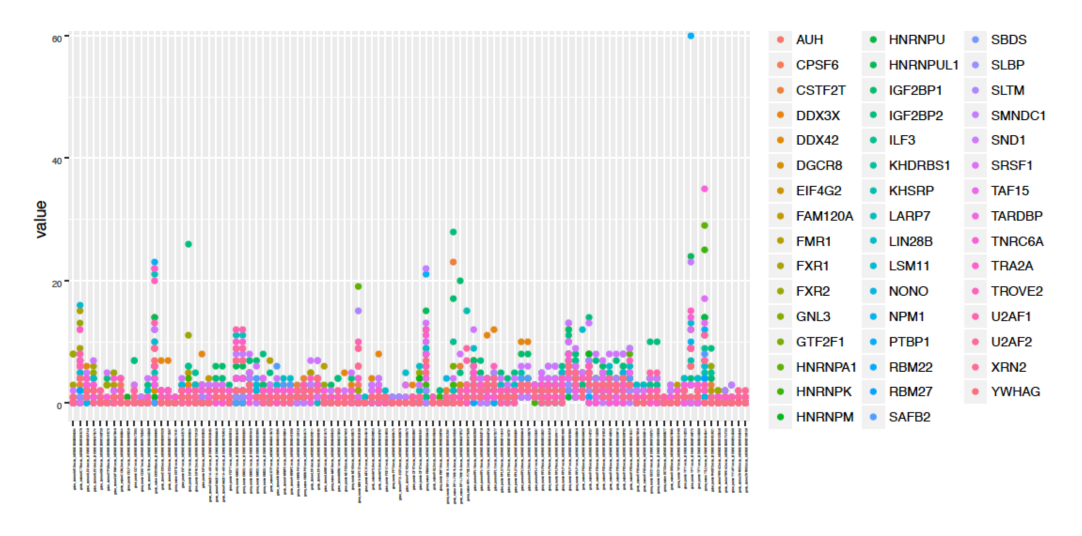
```
Codes:
##convert the long to short fromat
GeneName RBPName melt<-
melt(GeneName_RBPName_cast);
##melt result
>GeneName RBPName melt
Gene name
               RBP_name
                               count
                               3
TP53
               HNRNPK
               SRSF1
TP53
BRCA1
               HNRNPK
BRCA1
               SRSF1
```

Give some ggplot figure example

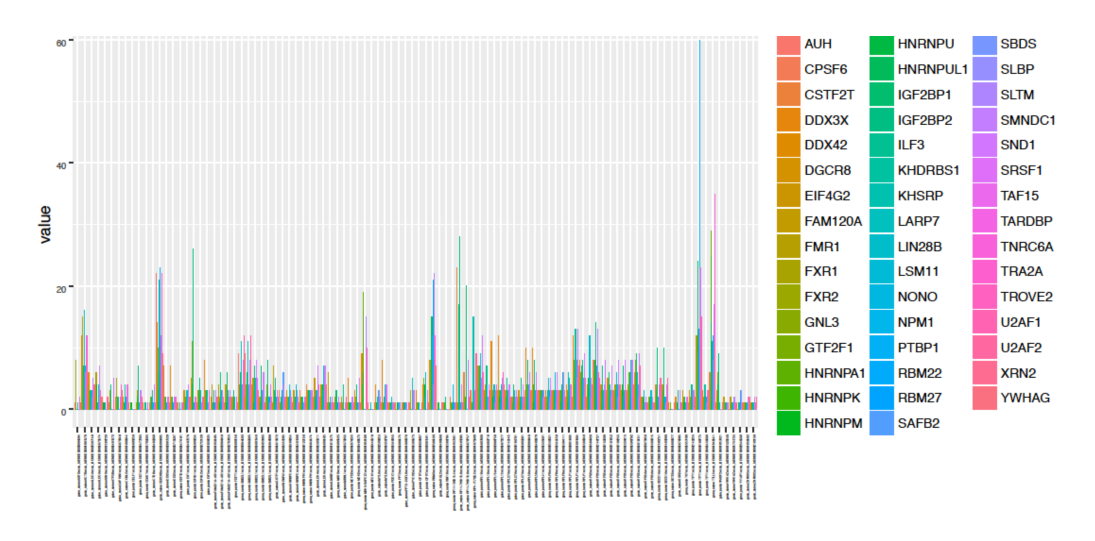
Codes: ##count the number RBP binding in each gene's exon region. GeneName RBPName point2<-ggplot(se target data melt)+ geom point(aes(x=gene name,y=value,color=variable)) ##plot GeneName_RBPName_bar<-ggplot(se_target_data_melt)+</pre> geom bar(aes(x=gene name,y=value,fill=variable),stat="identity",position="dodge" ")+ GeneName_RBPName_tile<-ggplot(se_target_data_melt)+</pre> geom_tile(aes(x=gene_name,y=variable, fill=value))

print(GeneName RBPName p1);

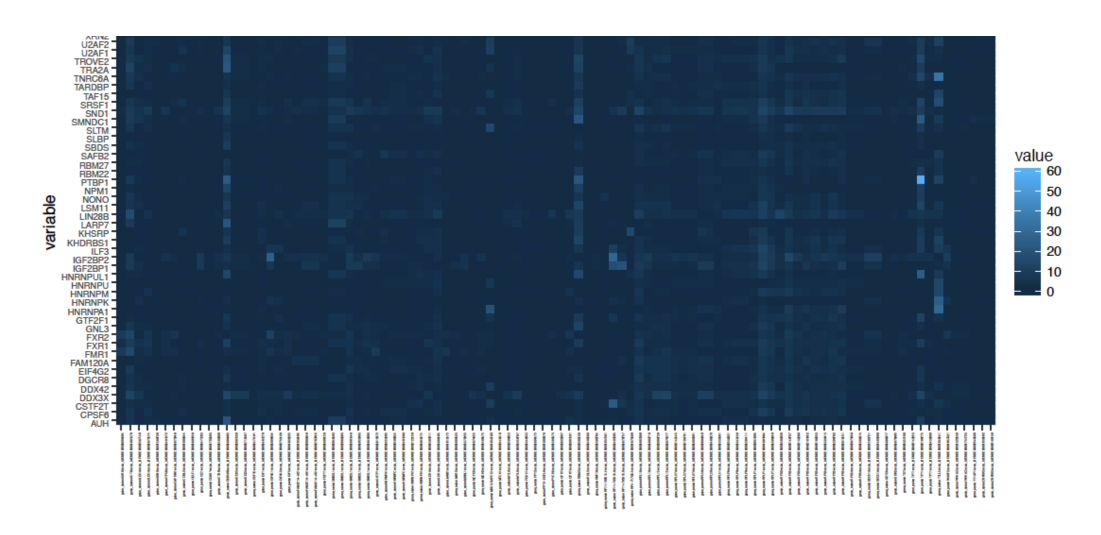
Figure_1



Figure_2



Figure_3



Which one is better?

- I will show you the raw PDF here, which is much clear than this one.
- Scatter plot is very easy to check which gene contain highest count, but if two RBP has same count in the gene, they will overlap. So scatter plot is wrong, and a possible solution is jitter.
- Bar plot avoid the overlap problem by putting each point beside each other, but the 47 colors is very hard to distinguish.
- So I think the best one is the heatmap.

R package ggplot2

You can see the power of ggplot2. only change a few codes can do different kinds of plots, very efficient and productivity.

R package ggplot2

ggplot2 is multi layers figure, as you can see, each '+' add a layer to the final figure.

aes: map variable to figure variable (color, x-axis, y-axis, shape, fill).

geom: The figure type, eg. Bar_plot, histogram, scatter plot.

stat: the statistic transformation, eg. bin, identity.

R package ggplot2

• I list the figure types that support by ggplot2.

Histogram, boxplot, barplot, contour, density, scatter, tileplot, hex plot, violin, pieplot, line and so on.

 If you want to combine two or ggplot2 object in one figure, you may need R function multiplot.r, which can be search and got from web.

R package ggplot2

- · Besides productivity, ggplot is very versatile.
- These include:
- geom_xx (different kinds of figures)
- stat_xx (different kinds of statistics transformation)
- · Scales (change the scales like color, legend, fill et.al.)
- Coordiante (cartesian, polar, flip, trans)
- Faceting (multi plots in one figure)
- Position adjustments (dodge, fill, identity, stack, jitter)
- Annotation (custom, logticks, map, raste)
- Fortify
- Themes (font, size, label orientation et.al.)
- Aesthetics (map from variable to axis)
- Others ()

Graphical Primitives

These geoms plot basic visual objects. You can use them to construct more sophisticated geoms.



geom_blank()

(Useful for expanding limits)



geom_curve()

x, xend, y, yend, alpha, angle, color, curvature, linetype, size



geom_path()

x, y, alpha, color, group, linetype, size



geom_polygon()

x, y, alpha, color, fill, group, linetype, size



geom_rect()

xmax, xmin, ymax, ymin, alpha, color, fill, linetype, size



geom_ribbon()

x, ymax, ymin, alpha, color, fill, group, linetype, size



geom_segment()

x, xend, y, yend, alpha, color, linetype, size



geom_spoke()

x, y, angle, radius, alpha, color, linetype, size



geom_tile()

x, y, alpha, color, fill, linetype, size, width

One Variable

These geoms are useful for visualizing the distribution of values in a single variable.

Continuous



geom_area(stat = "bin")

x, y, alpha, color, fill, linetype, size



geom_density(kernel = "gaussian")

x, y, alpha, color, fill, group, linetype, size, weight



geom_dotplot()

x, y, alpha, color, fill



geom_freqpoly(binwidth = 5)

x, y, alpha, color, group, linetype, size



geom_histogram(binwidth = 5)

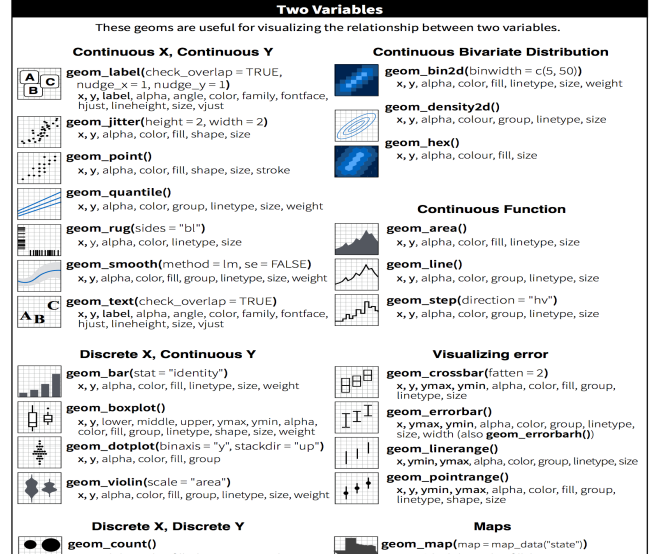
x, y, alpha, color, fill, linetype, size, weight

Discrete



geom_bar()

x, alpha, color, fill, linetype, size, weight





x, y, alpha, color, fill, shape, size, stroke



map_id, alpha, color, fill, linetype, size

Three Variables

These geoms are useful for visualizing the relationship between three variables.



geom_contour()

x, y, z, alpha, colour, group, linetype, size, weight



geom_raster(hjust=0.5, vjust=0.5,
interpolate=FALSE)
x, y, alpha, fill

Check the book: **R for data science** for details.

Calculate the number of genes for each RBP

We want to summarize which RBP binds most genes.

Generalize the problem

The generalize of the problem is **split-aggregate** problem, what we want is spliting the data.frame into multi small ones, and do some operation on each group.

There is a R package called 'plyr' which can be used to do these kinds of job.

The 'for' loop equivalent

R's **for loop** is seldom used in practice. But R provide other function to do iteration:

sapply: apply a function to each element in list.

apply: apply a function to row or column.

lapply: apply a function to each element in list.

Although R provide some functions for doing iteration. But they are not that powerful enough for some problems, the main problem is that the input and output format is ambiguous. While R package plyr solved this problem.

It includes many functions:

Where d means table, I means list, _ means nothing, a means vector, so:

d_ply means input is data.frame, output nothing. ddply means input is data.frame, output data.frame. dlply means input is data.frame, output list. daply means input is data.frame, output vector.

R package plyr is a more powerful solution for this kind of analysis. It group the table and apply a function. (also called split+modify+aggregate)

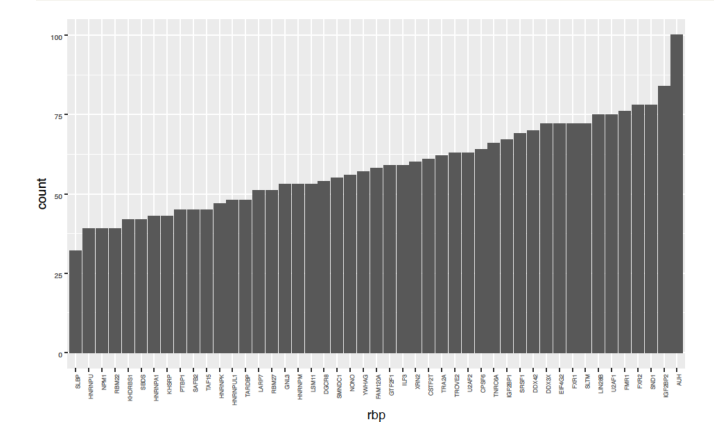
For our problem, we need to treat each RBP a separate group and count the number of genes in each group.

```
RBP_bind_fre<-ddply(GeneName_RBPName, (RBP_names), nrow)</pre>
#each line represent a overlap
  Gene_names RBP_names
  TP53
         SRSF1
  TP53
                 SRSF1
  TP53
                 HNRNPK
>RBP_bind_fre
RBP_names
                 Var1
SRSF1
HNRNPK
```

Calculate number of gene for each RBP

With a little ggplot2, we can do the below figure.

```
rbp_gene_count_plot<-ggplot(rbp_gene_count)+
  geom_bar(aes(x=rbp,y=count),stat="identity")+</pre>
```



· Besides group analysis, R package plyr can also do .

• 1. data.frame operations, like rbind.fill, progress bar,

• 2.generalized for loop, executing a function multitimes and return something (like data.frame).

The gene expression data

 Next we add the gene expression data, which can be got from RNA-seq technique, and do the correlation between gene expression and RBP binding information

 The key things to know is RNA-seq can quantatively determine the expression level of gene.

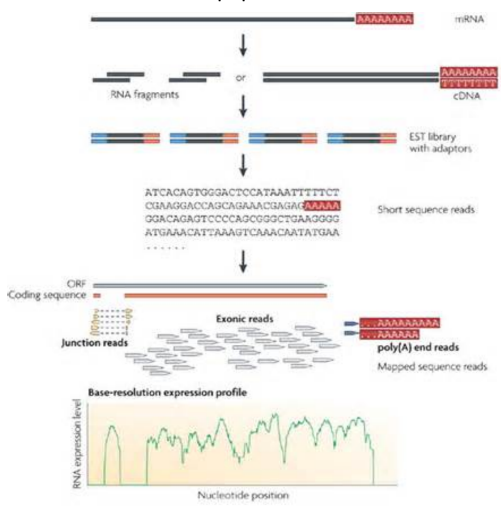
Add gene expression

Here, we use **NGSUtils** to calculate each sample's FPKM, to avoid the bias due to a single sample. We use 115 samples here and take the median FPKM as the gene's expression level.

We have 115 samples and we need to write a small **pipeline** to do it.

RNA-seq workflow

Got from Cufflinks paper



There are many kinds of RNA-seq technique which I also don't know very much.

Another consideration is that different sequencer will also give different result.

91

The RNA-seq data in Encode

The RNA-seq data set also **come from the ENCODE project**, we also assume the download format is fastq format.

Unlike sequence alignment for CLIP-seq, for RNA-seq, we must also consider the junction reads to get the bam file.

One software is the tophat2.

The RNA-seq data set

To ensure the expression data we got doesn't come from big variance, we use 104 samples' RNA-seq dataset from a same cell line (K562).

We use a tool called NGSUtils which can calculate the gene expression data from bam file.

A small Linux pipeline

After we got the bam files, we will try to call the **FPKM** values from the BAM files.

For large samples (like 104 samples here), we usually use pipeline to do batch process.

A lot of language can be used to write pipeline, here, I use bash shell, I think writing pipeline in **bash shell** is a nature way (bash.quickref.pdf).

The pipeline for NGSUtils here

```
for i in `ls -d dirContainBAMFiles`
do
  echo $i;
  sample=`basename $i`;
  my i dir=${i/.bam};
  mkdir $i dir;
  samtools sort $i $i.sort;
  samtools index $i.sort;
  script="bamutils count -gRBP Homo_sapiens.GRCh37.75_4.gRBP -rpkm -norm all $i.sort >
$sample.count";
  echo $script > $sample.count.job;
  `$script`
  #qsub -l nodes=1:ppn=1,walltime=4:00:00,mem=4gb -M li487@iupui.edu -d
/N/dc2/scratch/liulab/limeng/RBP network/ -m ae -N $sample.count $sample.count.job
```

done;

Process Gene expression

For each sample, now we got a gene expression table.

```
#gene geneid isoid chrom strand txstart txend length count count (CPM)
WASH7P ENSG00000227232 ENSG00000227232 chr1 - 14362 29806 2073
                                                                 309
                                                                       1.76326905472  0.850588063058
DDX11L1 FNSG00000223972 FNSG00000223972 chr1 + 11868 14412 1756 41
                                                                       0.23396126616  0.133235345194
MIR1302-10
            ENSG00000243485 ENSG00000243485 chr1 +
                                                    29553 31109 1021 65
                                                                         0.370914202449
0.363285212977
FAM138A ENSG00000237613 ENSG00000237613 chr1 - 34553 36081 1219 224 1,27822740536 1,04858687889
OR4G4P FNSG00000268020 FNSG00000268020 chr1 + 52472 54936 947
                                                                 10 0.0570637234537 0.0602573637315
OR4G11P ENSG00000240361 ENSG00000240361 chr1 + 62947 63887 940
                                                                      0.0 0.0
                                                                0
OR4F5 FNSG00000186092 FNSG00000186092 chr1 + 69090 70008 918
                                                               3
                                                                    0.0171191170361 0.0186482756385
RP11-34P13.7 ENSG00000238009 ENSG00000238009 chr1 - 89294 133566 3569 2050 11.698063308 3.2776865531
RP11-34P13.8 ENSG00000239945 ENSG00000239945 chr1 - 89550 91105 1319 561 3.20127488575 2.42704691869
```

Process gene expression

Next, we need to combine multi sample's gene expression into one table. This can be done in multi ways since each file is already ordered by the gene name, including using R read each file. Use linux command cut and paste.

| X.gene | ENCFF000DV | ENCFF000DV | ENCFF000DV | ENCFF000DV | ENCFF000DY | ENCFF000DY |
|-------------|------------|------------|------------|------------|-------------------|------------|
| WASH7P | 148.049887 | 167.275891 | 401.776058 | 428.998374 | 19.3728753 | 68.0487607 |
| DDX11L1 | 2.33384883 | 2.42390186 | 0 | 0.61342398 | 1.11666424 | 0.25012929 |
| MIR1302-10 | 0.90201027 | 1.44107576 | 0.36055293 | 2.11003431 | 3.88415134 | 2.11511547 |
| FAM138A | 0.83104818 | 1.16389709 | 0 | 1.59057468 | 0.2938757 | 0 |
| OR4G4P | 0.09724947 | 0 | 0 | 0 | 0.11945795 | 0.23190445 |
| OR4G11P | 0 | 0 | 0 | 0 | 0.00668597 | 0 |
| OR4F5 | 0 | 0 | 0 | 0 | 0.02738482 | 0 |
| RP11-34P13. | 25.8042164 | 24.1905387 | 83.6506025 | 74.306493 | 2.14483193 | 4.22530905 |
| RP11-34P13. | 6.07451601 | 8.4458933 | 0 | 0 | 2.5301266 | 0.38850003 |
| RP11-34P13. | 59.1633562 | 73.6970671 | 155.569658 | 144.734635 | 11.2270541 | 13.258401 |
| RP11-34P13. | 30.8065091 | 33.6749976 | 81.0133226 | 75.8572195 | 15.2915755 | 4.76860331 |
| RP11-34P13. | 32.7517539 | 46.144623 | 30.7176767 | 29.3904024 | 14.7089653 | 1.35743453 |
| CICP27 | 0.06039825 | 0.01378484 | 0.19313984 | 0 | 1.07329875 | 1.67072191 |
| AL627309.1 | 16.6470554 | 22.1368215 | 17.3617548 | 16.4348757 | 8.526187 | 1.07289579 |

Gene expression quality control

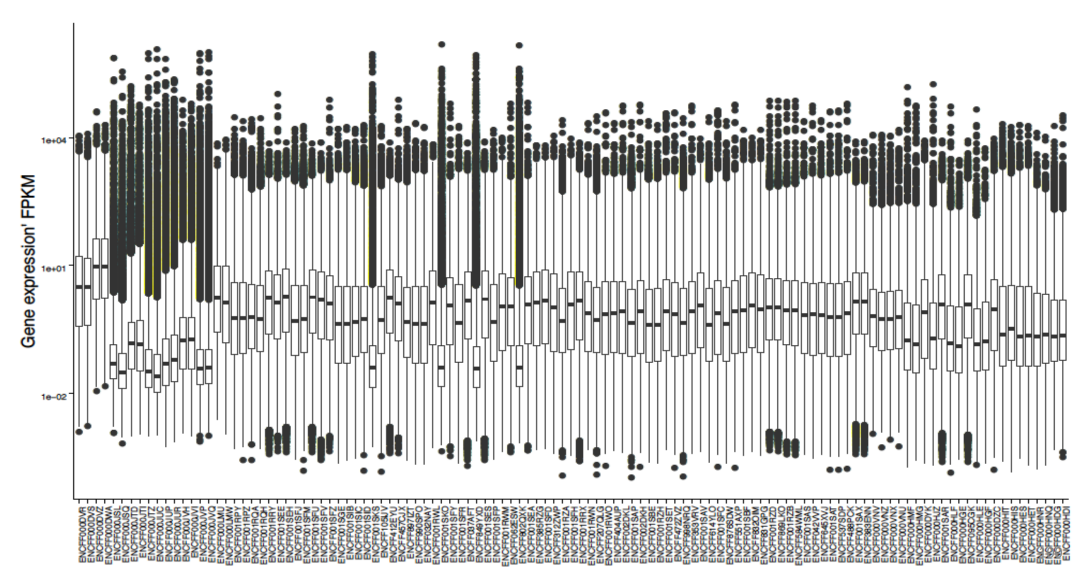
Usually, before we do a experiment, we need to validate the result we got and the feasibility of next step. (quality control)

Here we use heatmap and boxplot to do quality control.

Boxplot code

```
gene expression tbl<-
read.table("gene.expression.tsv", sep="\t", as.is=TRUE);
#ggplot receive short format table
gene_expression_tbl_melt<-melt(gene expression tbl);</pre>
gene_exp_box_plot<-ggplot(gene_expression_tbl_melt)+</pre>
 #geom boxplot are used to plot box plot.
  geom_boxplot(aes(x=variable, y=value ) ) +
  theme classic()+scale_y_log10();
```

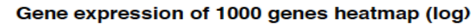
Figure

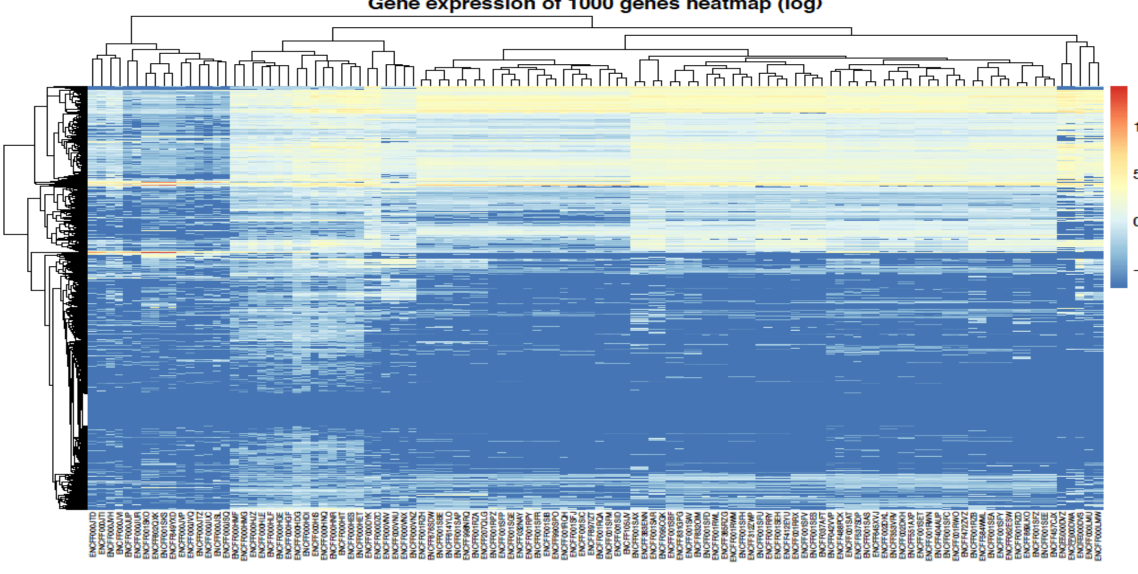


Heatmap

A more generalize way to check gene expression is using heatmap, however, ggplot2 doesn't provide this, here I use R package 'pheatmap'.

Figure





What we can see from the figures?

Seems the there are **two different groups of samples**, although they come from only one cell line 'K562'. It turns out the some of the samples come from a independent culture 'K562'.

Gene Expression

Now we take the median of gene expression among samples.

```
And the gene expression.

Gene_name Sample1 Sample2 Sample3 ...

TP53 113.0 313 112

BRCA1 313 113.1 132
.....
```



And the gene expression.

Gene_name Median(FPKM) ..

TP53 213.4

BRCA1 313.5
.....

Combine the RBP and gene expression table

```
Now we have a table contain RBP binding information.

Gene_name SRSF1 BX1

TP53 1 3

BRCA1 3 1

.....
```

```
And the gene expression.

Gene_name Median(FPKM) .

TP53 213.4

BRCA1 313.5
.....
```

Generalize of the problem

- Combine two tables is a daily task for SQL, and it include many kinds of operations, like left_join, inner_join, right_join and full_join.
- Left_join: keep all the data in left table in the result table.
- Right_join: keep all the data in the right table in the result table.
- Inner_join: keep the data in both the tables.
- Full_join:keep the data in either the table.

Here we use R package dplyr. Table manupilation language. A very powerful tool to process tables in R. It implements the above join operations in R, and make multi tables analysis more easily.

Its function include:

filter - select a subset of the rows of a data frame.

arrange – works similarly to filter, except that instead of filtering or selecting rows, it reorders them.

select - select columns of a data frame.

mutate - add new columns to a data frame that are functions of existing columns.

distinct – select distinct rows from a data frame.

summarize - summarize values.

group_by - describe how to break a data frame into groups of rows.

SQL JOIN - sql style joining multi tables together.

Join two tables

```
>geneName_RBPName_cast
Gene_name SRSF1 BX1
TP53 1
BRCA1
>gene expression tbl melt
Gene_name Median(FPKM)...
TP53 213.4
BRCA1 313.5
Gene_RBP_exp_join<-</pre>
inner_join(geneName_RBPName_cast,
gene_expression_tbl_melt,by=c("Gene_name"="Gene_name"));
```

Join two tables

```
Gene_name Median(FPKM) SRSF1 BX1

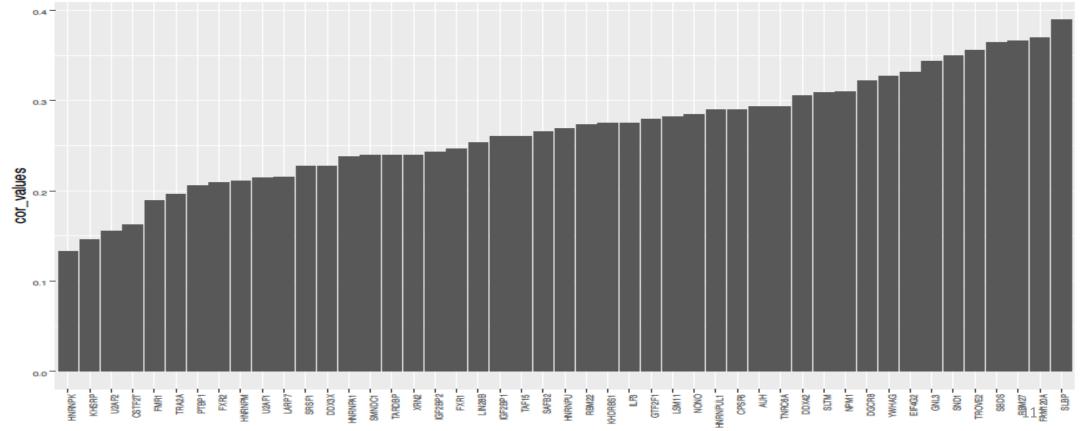
TP53 213.4 1 3

BRCA1 313.5 3 1
.....
```

Now we have all the data in one table.

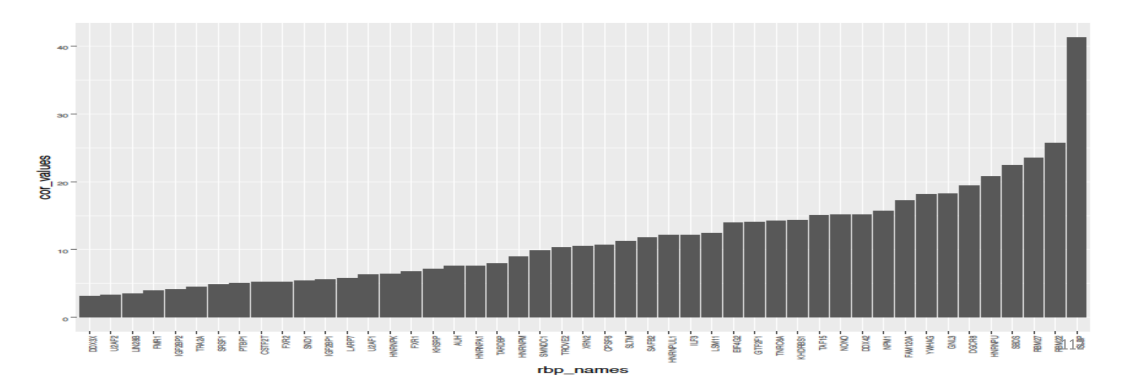
Correlation of RBP and Gene expression

```
With a little ggplot:
rbp_cor_values_data_plot<-
ggplot(rbp_exp_data)+geom_bar(aes(x=rbp_names,y=cor_values),stat="identity")</pre>
```



Linear regression of RBP and Gene expression

Another consideration is longer gene region has higher possibility to contain more RBPs, so we need to consider the gene's length.



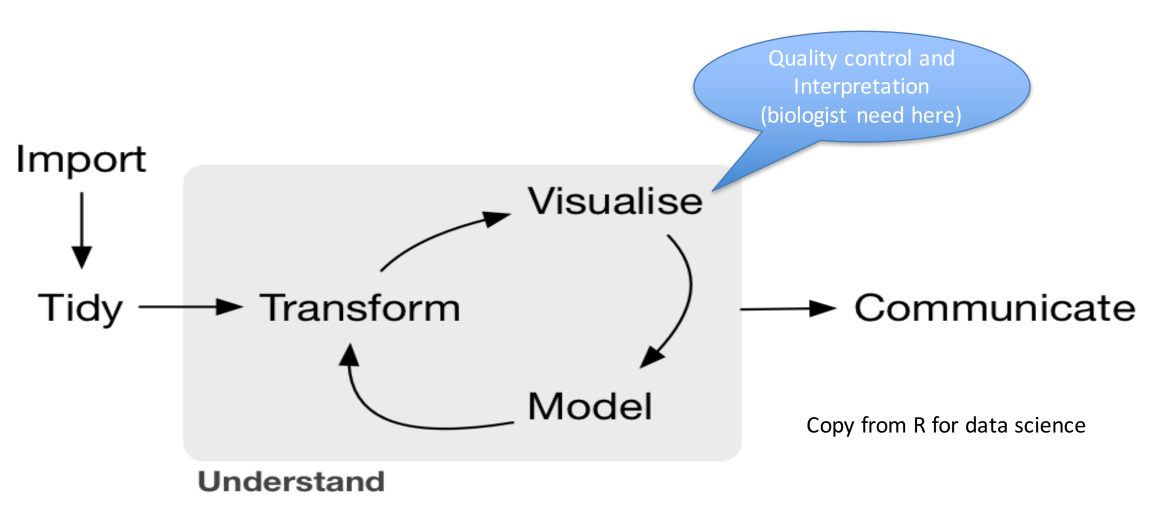
The pipe

 Usually, we will do many modification on the raw data to get the final data we need, which will make the code very complicate.

 Linux has a pipe which facilitate complicate operation. R also has a pipe command, and here it is: %>%

Check the magnittr package for more details.

Summary: a daily life for data scientist



A few advice to newbie when do science

Actually I also am a newbie, but I also some advice to the more newer ones.

- 1. Tidy up the workspace of your project. (like make some folders: result, code, figures, reference, document, discussion, tables, annotation, data, et.al.)
- 2. Record every command ever used. (So if result go wrong, we know where we do mistake, thus can avoid make same mistake twice)
- 3. Double check every step!!

Further reading

http://hadley.nz

Advance R. http://adv-r.had.co.nz/

R for data science. http://r4ds.had.co.nz/pipes.html

ggplot2: elegant graphics for data analysis. (This one is not free)

ggplot2: http://docs.ggplot2.org/0.9.3.1/geom_bar.html

Bimedical data science: http://genomicsclass.github.io/book/

Bioconductor tutorial: https://www.bioconductor.org/help/course-materials/

I recommend read the "R for data science" first.

Acknowledgement

冯伟兴教授和其它实验室的老师和同学们。