# RNA-seq解析パイプライン: Transcript-based

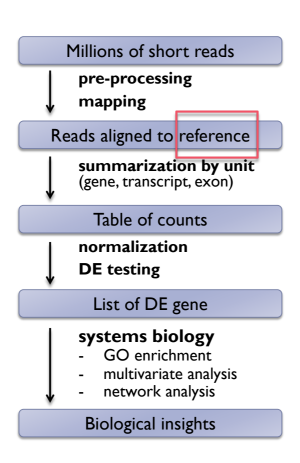
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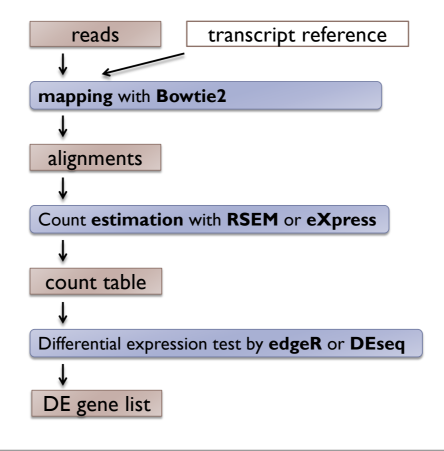


# Two Basic Pipelines

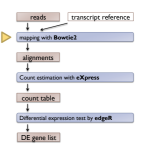
- Choice of reference
  - ▶ **Genome** standard for genome-known species
  - Transcript the only way for genome-unknown species
     -- can be used for genome-known species



### A Pipeline: Transcript-based



# Mapping – alignment software



- For mapping reads onto transcript reference
  - short read mapper (unspliced read aligner) is used
  - Bowtie2

http://bowtie-bio.sourceforge.net/bowtie2/index.shtml

### bowtie2

Bowtie is an ultrafast, memory-efficient short read aligner.

http://bowtie-bio.sourceforge.net/bowtie2/index.shtml

```
(example)
$ bowtie2 -x transcript.fa -U reads.fq -a -S out.sam
```

- Input
  - ▶ Reads (fastq) and reference (bowtie2-db)
- Output
  - ▶ Alignment in SAM format: out.sam

## Let's Try Bowtie2

Align 75-bp Illumina reads with a transcript reference using Bowtie2.

#### Prepare reads and reference genome

Sequences for this exercise are stored in ~/data/SS/.

```
IlluminaReads1.fq - Illumina reads in fastq format
minimouse_mRNA.fa - a set of transcript sequences
```

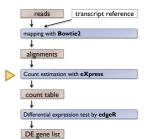
#### **Build index of reference sequence**

```
$bowtie2-build minimouse_mRNA.fa myref
```

#### Align reads with reference

```
$bowtie2 -x myref -U IlluminaReads1.fq -a -S out.sam
```

# Count Reads by Transcript/gene



▶ The simplest way: just count reads by contig.

#### But...

▶ <u>Mapping ambiguity</u> should be taken into consideration.

### Estimate Abundance

- Multimapping issues
  - Isoforms
  - Very similar paralogs
  - Repetitive sequences
  - > => cannot align reads uniquely
- ▶ Mapping ambiguity should be taken into consideration.



- Critical for RNA-seq de novo analysis
- Software: RSEM and eXpress (EM algorithm)

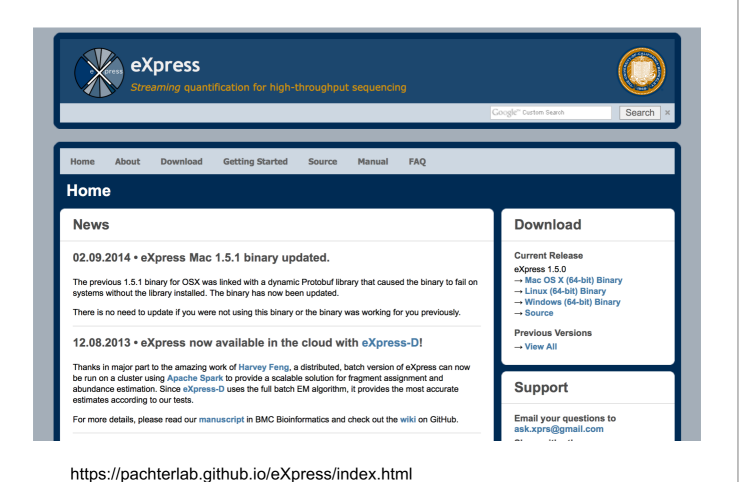
### eXpress

eXpress is a streaming tool for quantifying the abundances of a set of target sequences from sampled subsequences.

http://bio.math.berkeley.edu/eXpress/

```
(example)
$ express transcripts.fasta hits.bam
```

- Input
  - alignment (bam|sam) and reference (fasta)
- Output
  - Count estimation table: results.xprs



# Let's Try eXpress

#### Prepare alignments and reference genome

Sequences for this exercise are stored in ~/data/SS/.

IlluminaReads1.fq - Illumina reads in fastq format
out.sam - this file should be generated in the previous bowtie practice

#### Run eXpress

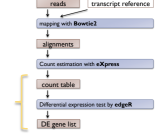
\$ express minimouse mRNA.fa out.sam

Output : results.xprs, params.xprs

# eXpress: output

#### results.xprs

-4	A	В	С	D	E	F	G	Н	I	J	K	L	M	N
1	bundle_id	target_id	length	eff_length	tot_counts	uniq_counts	est_counts	eff_counts	ambig_distr_alpha	ambig_distr_beta	fpkm	fpkm_conf_low	fpkm_conf_high	solvable
2	1	m.245853	621	398.1	807	15	86.2	134.4	9.83E+01		2.34E+01			T
3	1	m.245856	660	442.0	991	199	919.8	1373.4	5.53E+01	5.46E+00	2.25E+02	2.12E+02	2.38E+02	T
4	2	m.42076	1959	1591.7	156	156	156.0	192.0	0.00E+00	0.00E+00	1.06E+01	1.06E+01	1.06E+01	T
5	3	m.60782	291	83.0	12	12	12.0	42.1	0.00E+00	0.00E+00	1.57E+01	1.57E+01	1.57E+01	T
6	4	m.158451	282	64.5	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
7	5	m.337354	219	39.4	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
8	6	m.338934	261	82.3	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
9	7	m.5973	822	719.9	4	4	4.0	4.6	0.00E+00	0.00E+00	6.01E-01	6.01E-01	6.01E-01	T
10	8	m.337793	219	38.7	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
11	9	m.340910	210	40.5	0	0	0.0	0.0	0		0.00E+00	0.00E+00	0.00E+00	T
12	10	m.289784	3177	2521.4	350	350	350.0	441.0	0.00E+00	0.00E+00	1.50E+01	1.50E+01	1.50E+01	T
13	11	m.248666	240	61.8	1	1	1.0	3.9	0.00E+00	0.00E+00	1.75E+00	1.75E+00	1.75E+00	T
14	12	m.90727	240	55.7	13	13	13.0	56.1	0.00E+00	0.00E+00	2.53E+01	2.53E+01	2.53E+01	Т
15	13	m.338727	216	48.1	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
16	14	m.123519	225	43.2	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
17	15	m.328661	251	50.8	1	1	1.0	4.9	0.00E+00	0.00E+00	2.13E+00	2.13E+00	2.13E+00	Т
18	16	m.26062	642	356.1	1	1	1.0	1.8	0.00E+00	0.00E+00	3.04E-01	3.04E-01	3.04E-01	T
19	17	m.1295	240	53.6	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
20	18	m.307626	201	220.2	4	3	3.0	2.7	8.33E+00	4.07E+04	1.47E+00	1.46E+00	1.49E+00	T
21	18	m.307625	204	35.7	301	300	301.0	1718.3	1.02E+01	2.10E-03	9.12E+02	9.05E+02	9.18E+02	T
22	19	m.49789	237	51.9	3	3	3.0	13.7	0.00E+00	0.00E+00	6.26E+00	6.26E+00	6.26E+00	T
23	20	m.33508	162	151.3	1	1	1.0	1.1	0.00E+00	0.00E+00	7.15E-01	7.15E-01	7.15E-01	T
24	21	m.109341	183	286.3	2	2	2.0	1.3	0.00E+00	0.00E+00	7.56E-01	7.56E-01	7.56E-01	T
25	22	m.331919	564	277.3	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
26	23	m.23766	303	98.5	3	3	3.0	9.2	0.00E+00	0.00E+00	3.30E+00	3.30E+00	3.30E+00	T
27	24	m.246777	1149	1152.1	631	29	202.5	202.0	1.58E+02	3.90E+02	1.90E+01	1.65E+01	2.15E+01	T
28	24	m.246852	1323	1315.4	761	156	588.8	592.2	1.22E+02	4.85E+01	4.84E+01	4.50E+01	5.19E+01	T
29	24	m.246633	207	31.8	10	4	5.7	37.1	1.29E+04	3.27E+04	1.94E+01	1.05E+01	2.82E+01	T
30	24	m.246662	192	200.4	6	3	3.0	2.9	1.20E+01	3.22E+03	1.63E+00	1.51E+00	1.74E+00	T
31	25	m.99743	1641	1387.9	470	470	470.0	555.7	0.00E+00		3.66E+01			
32	26	m.335620	234	58.9	0	0	0.0	0.0	0	0	0.00E+00	0.00E+00	0.00E+00	T
33	27	m.16882	528	297.5	14	14	14.0	24.9	0.00E+00	0.00E+00	5.09E+00	5.09E+00	5.09E+00	T
34	28	m.77438	255	81.4	9	9	9.0	28.2	0.00E+00	0.00E+00	1.20E+01	1.20E+01	1.20E+01	Т
35	29	m.131505	450	263.2	18	11	15.8	27.1	8.87E+00	3.95E+00	6.51E+00	4.68E+00	8.35E+00	T
36	29	m.131517	170	195.9	6	0	1.8	1.5	8.17E+00	1.96E+01	9.74E-01	0.00E+00	2.46E+00	T
37	29	m.131504	705	528.2	15	14	14.4	19.2	6.53E+01	1.01E+02	2.95E+00	2.69E+00	3.21E+00	Т



### Table of counts

data import
diagnostics
normalization
DE testing

List of DE gene

evaluation

## edgeR

- A Bioconductor package for differential expression analysis of digital gene expression data
- Model: An over dispersed Poisson model, negative binomial (NB) model, is used
- Normalization: TMM method (trimmed mean of M values) to deal with composition effects
- ▶ **DE test**: exact test and generalized linear models (GLM)

## edgeR (classic)

- input: count data (not RPKM or TPM)
- output: gene table with DE significance statistics (FDR)

```
(example)
$ R
> library(edgeR)
                                            #load edgeR library
> dat <- read.delim("count data.txt", ...) #import count table to R
> group <- c(rep("M", 3), rep("H", 3)) #assign groups</pre>
> D <- DGEList(dat, group=group) #import data to edgeR
> D <- calcNormFactors(D) #normalization (TMM)
> D <- estimateCommonDisp(D) #estimate common dispersion
> D <- estimateTagwiseDisp(D) #estimate tagwise dispersion</pre>
> de <- exactTest(D, pair=c("M", "H")) #DE test</pre>
> topTags(de)
Comparison of groups: H-M
              logConc
                         logFC P.Value
AT5G48430 -15.36821 6.255498 9.919041e-12 2.600872e-07
AT5G31702 -15.88641 5.662522 3.637593e-10 4.083773e-06
AT3G55150 -17.01537 5.870635 4.672331e-10 4.083773e-06
```

## Let's try edgeR

- edgeR classic
  - ex5: Differential expression analysis with edgeR (pairwise)

Advanced

# edgeR (GLM)

- input: count data (not RPKM or TPM)
- output: gene table with DE significance statistics (FDR)

```
(example)
$ R
> library(edgeR)
                                      #load edgeR library
> dat <- read.delim("count data.txt", ...) #import count table to R
> treat <- factor(c("M", "M", "M", "H", "H", "H")) ")</pre>
> treat <- relevel(treat, ref="M)</pre>
> design <- model.matrix(~treat)</pre>
> rownames(design) <- colnames(y)</pre>
> D <- DGEList(dat, group=treat) #import data to edgeR
> D <- calcNormFactors(D, method="TMM") #normalization (TMM)
> D <- estimateDisp(D, design) #estimate dispersion</pre>
> fit <- glmFit(D, design)</pre>
                                      #fitting to model
> lrt <- glmLRTt(D, coef=2))
                                     #DE test
> topTags(lrt)
```

# Let's try edgeR (GLM)

- edgeR linear model [advanced]
  - ▶ ex12-1: Differential expression analysis with edgeR (GLM)
  - ▶ ex12-2: Differential expression analysis with edgeR (GLM; considering batch effect)

A transcript-based pipeline (alignment-free method)

new

transcript reference reads mapping alignments **Ultrafast!** Count estimation with kallisto or salmon count table Differential expression test by edgeR or DEseq DE gene list

new

# Alignment-free RNAseq quantification

#### Software

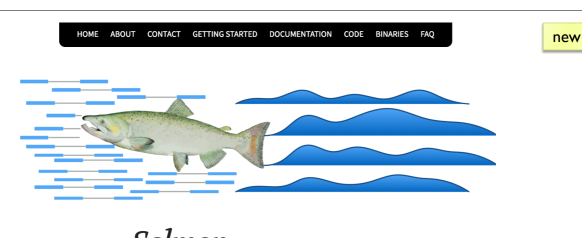
- Salmon
- Kallisto
- Sailfish

### Concept

- Precise alignments are not required to assign reads to their origins.
- > => "psudo-alignment" using a de bruijn graph information (kallisto), a k-mer approach (Sailfish old ver.) or a "quasi-mapping" (Salmon)

#### Benefit

- Ultrafast
- Computationally cheap
- Accuracy: similar or better than mapping-based methods



# Salmon —Don't count . . . quantify!

#### **Overview**

Salmon is a tool for quantifying the expression of transcripts using RNA-seq data. Salmon uses new algorithms (specifically, coupling the concept of quasi-mapping with a two-phase inference procedure) to provide accurate expression estimates very quickly (i.e. wicked-fast) and while using little memory. Salmon performs its inference using an expressive and realistic model of RNA-seq data that takes into account experimental attributes and biases commonly observed in real RNA-seq data.

### **Citing Salmon**

new

### Salmon

Salmon is a tool for quantifying the expression of transcripts using RNA-seq data. Salmon uses new algorithms to provide accurate expression estimates very quickly.

```
(example)
$salmon index ... # step 1. build index
$salmon quant ... # step 2. quantification
```

- Input
  - reference (fasta) and reads (fastq)
- Output
  - Count estimation table: quant.sf

ex14

new

# Let's Try Salmon

Align 75-bp Illumina reads with a transcript reference using Bowtie2.

### Prepare reads and reference genome

Sequences for this exercise are stored in ~/data/SS/.

```
IlluminaReads1.fq - Illumina reads in fastq format
minimouse_mRNA.fa - a set of transcript sequences
```

### **Build index of reference sequence**

```
$salmon index -t minimouse_mRNA.fa \
  -i minimouse_mRNA.fa.salmon_quasi.idx --type quasi -k 31
```

#### Quantification

```
$salmon quant -i minimouse_mRNA.fa.salmon_quasi.idx \
-l U -o salmon_out -r IlluminaReads1.fq
```