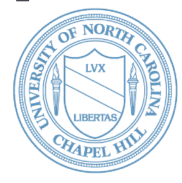
## Statistical methods for RNA-seq data

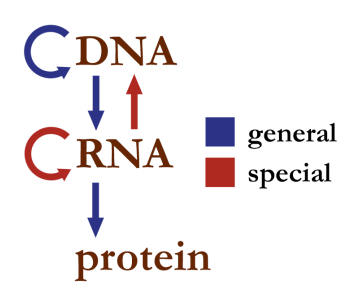
Wei Sun
Department of Biostatistics
Department of Genetics
University of North Carolina, Chapel Hill

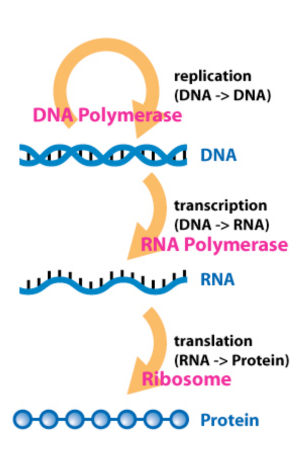


#### Outline

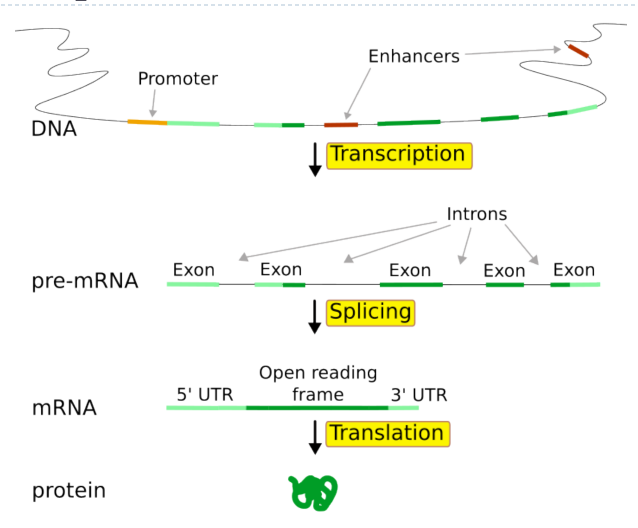
- ▶ I.An brief introduction to RNAseq
- ▶ 2. Allele-specific expression
- ▶ 3. Isoform-specific expression

## Central Dogma

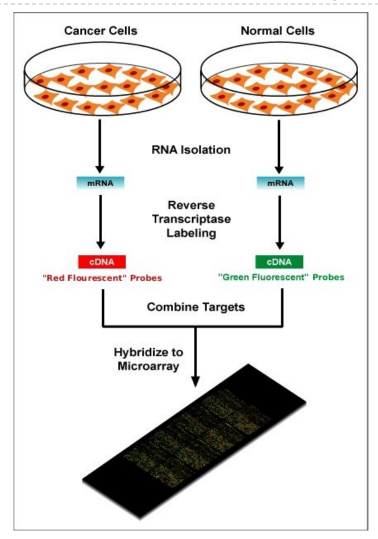




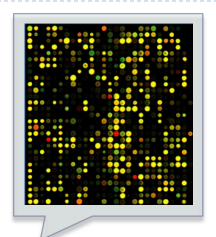
# Transcription and Translation



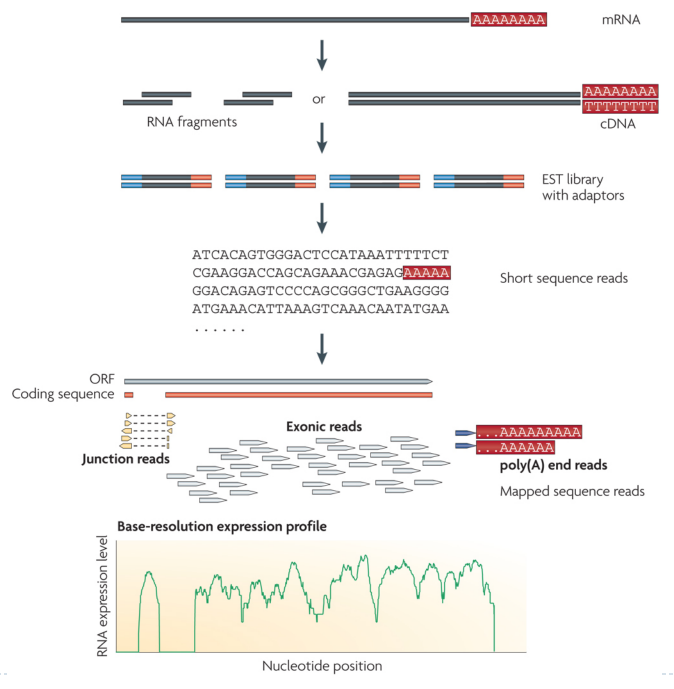
# Gene expression microarray



# Gene expression microarray



	Sample I	Sample 2	•••	Sample n
Gene I	10.45	10.52		10.40
Gene 2	4.63	4.76		4.70
Gene 3	8.80	8.96		8.82
•••				
•••				
Gene p	8.30	8.44		8.27

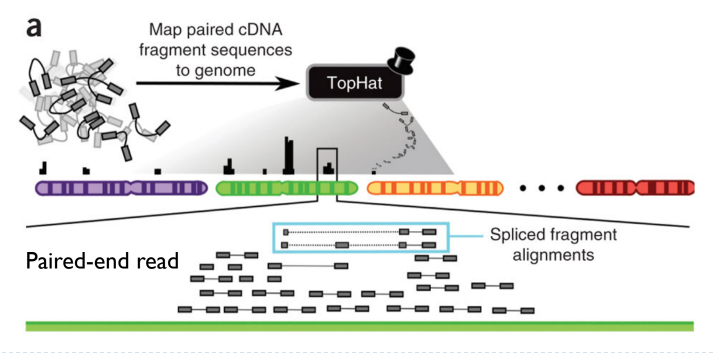


Nat Rev Genet. 2009 Jan; 10(1):57-63

### RNA-seq

RNA are cut into small fragments, select those fragments around certain base pairs (e.g., 400bps).

Sequence one end or both ends of the sequence fragments. Each sequenced part of an RNA-seq fragment is called an RNA-seq read. Map the RNA-seq reads to the genome, then count the # of fragments per gene



#### Bowtie

Extremely fast, general purpose short read aligner

# TopHat Aligns RNA-Seq reads to the genome using Bowtie Discovers splice sites

#### Cufflinks package

Cufflinks

Assembles transcripts

Cuffcompare

Compares transcript assemblies to annotation

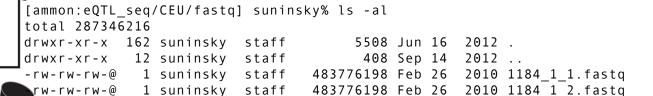
Cuffmerge

Merges two or more transcript assemblies

Cuffdiff

Finds differentially expressed genes and transcripts

Detects differential splicing and promoter use



bsub -M 16 -q week tophat --segment-length 17

- -G ~/research/eQTL\_seq/CEU/anno/Homo\_sapiens.GRCh37.66.updated.gtf
- -o /lustre/scr/w/e/weisun/CEU/tophat/NA12892 -r 8
- ~/bin/bowtie/indexes/hg19 | 184 | 1.fastq | 184 | 2.fastq

cufflinks -p 8 -o Cl RI clout Cl RI thout/accepted hits.bam

Create a file called assemblies.txt that lists the assembly file for each sample. The file should contain the following lines:

 $./CI\_RI\_clout/transcripts.gtf$ 

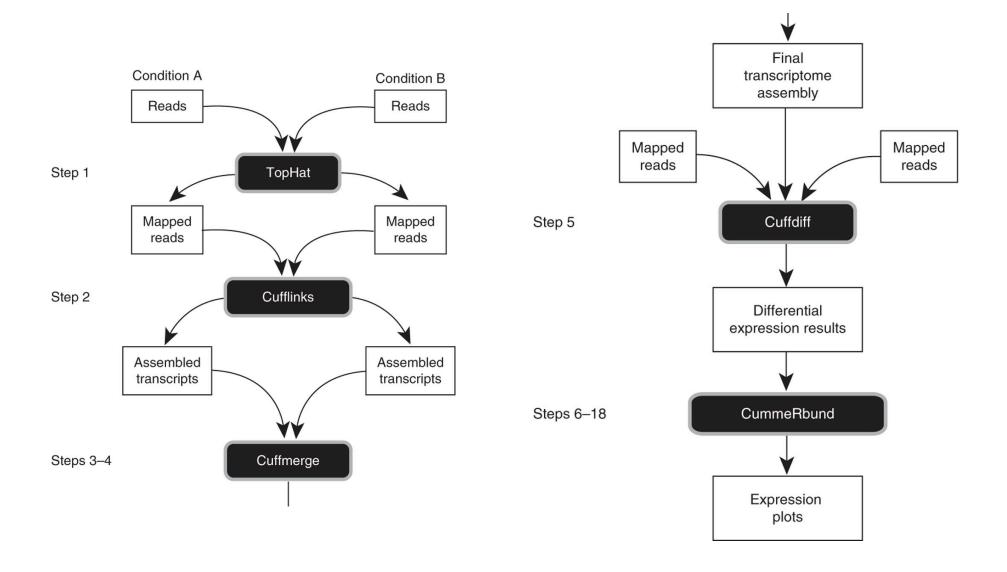
./C2 R2 clout/transcripts.gtf

cuffmerge -g genes.gtf -s genome.fa -p 8 assemblies.txt

cuffdiff -o diff\_out -b genome.fa -p 8 -L C1,C2 -u merged\_asm/merged.gtf

CummeRbund
Plots abundance and differential
expression results from Cuffdiff

Nat Protoc. 2012 Mar1;7(3):562-78.



\_\_\_\_\_\_

# RNA-seq

@HWI-EAS134:1:1:0:1446#0/1

GGGACTTAATCAACGCAAGCTTATGACCCGCACTT

+

BB<2A81(66B<2>BB0>?B=BA6=80BCBB0?3/

bed-tools

	Sample I	Sample 2	•••	Sample n
Gene I	512	339		286
Gene 2	1043	1212		888
Gene 3	20	12		10
•••				
•••				
Gene p	78	65		42

## Microarray vs. RNA-seq

- ▶ RNA-seq is more accurate
  - Larger dynamic range
- ▶ RNA-seq is more expensive
- ▶ RNA-seq is less high throughput
- What else?
  - Allele-specific expression
  - ▶ RNA isoform specific expression

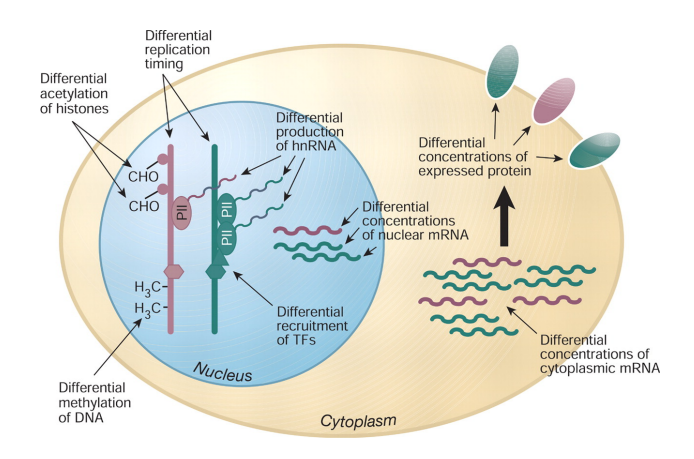
#### Outline

▶ I.An brief introduction to RNAseq

▶ 2.Allele-specific expression

▶ 3. Isoform-specific expression

# Allele-specific expression (ASE)



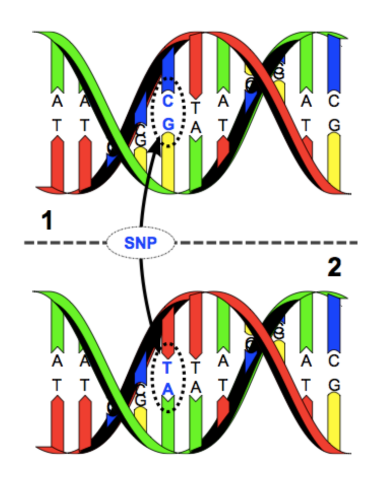
Science 22 October 2004: vol. 306 no. 5696 647-650

#### SNP

A single-nucleotide polymorphism (SNP) is a DNA sequence variation occurring when a single nucleotide — A,T, C or G — in the genome differs between members of a biological species or paired chromosomes in an individual

We have two sets of chromosomes, i.e., two double-helix structure

The genotype of this SNP of this individual is CT (heterozygous)

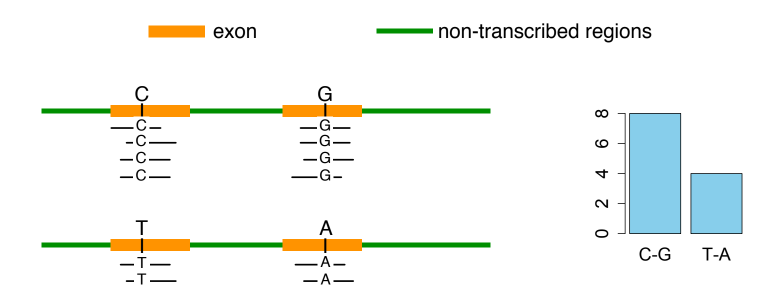


The genotypes at other locations are AA, AA, CC, TT, AA, AA, CC (homozygous)

## Allele-specific expression (ASE)

- ▶ ASE can not be measured by microarray
  - Microarray use the same probe for both alleles of a gene
- ASE can be measured by RNA-seq
  - Only for those RNA-seq reads that overlap with heterozygous SNPs

### Gene-level allele-specific expression (ASE)



- ✓ A pipeline for appropriate sequence alignment.
- ✓ Need haplotype information to combine the ASE at each SNP/indel to obtain gene level ASE
- Sun (2012), A Statistical Framework for eQTL mapping using RNA-seq, Biometrics, 68(1): 1-11

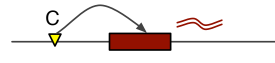
# eQTL (gene expression QTL) Mapping Using ASE

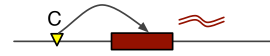
cis-acting eQTL

VS.

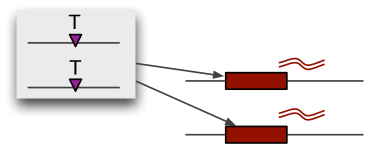
trans-acting eQTL

Sample 1: genotype CC

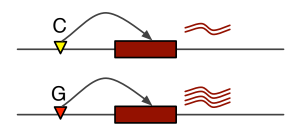




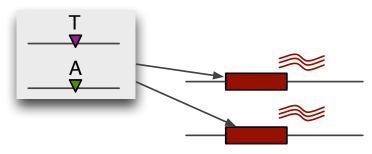
Sample 1: genotype TT

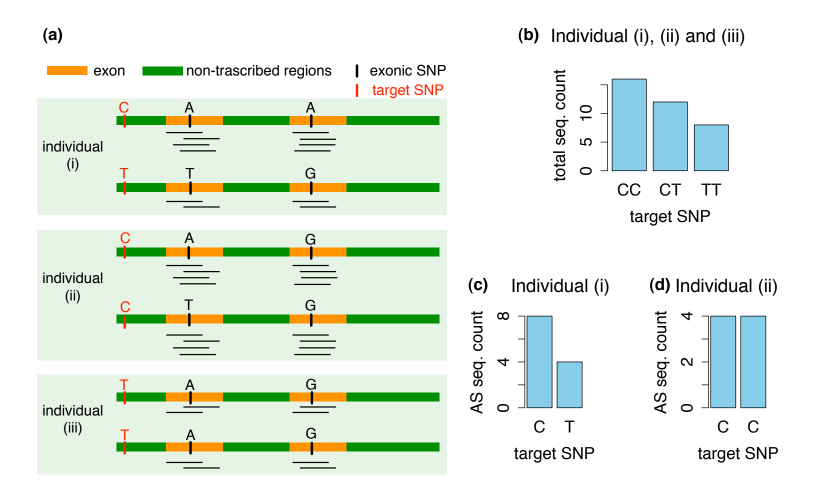


Sample 2: genotype CG



Sample 2: genotype TA

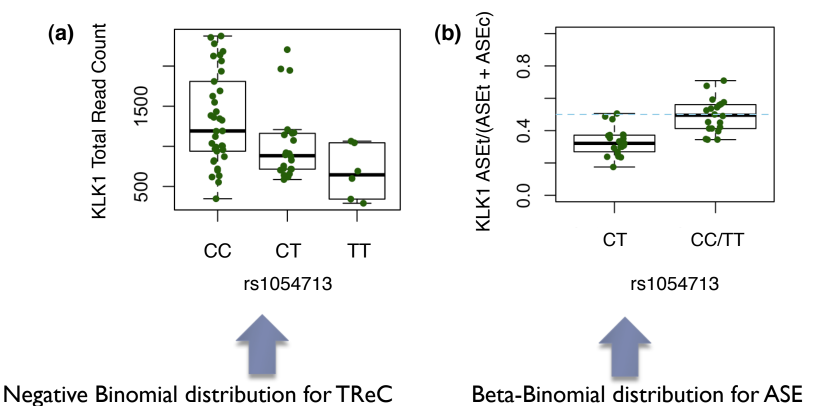




#### Why?

- Combine the information of total expression and ASE to improves the power of eQTL mapping
  - Using ASE, we compare the expression of paternal allele vs. maternal allele within an individual. One allele is an internal control of the other allele, and thus many confounding factors are canceled out.
- Identify cis-acting eQTL that directly influences gene expression in an allele-specific manner

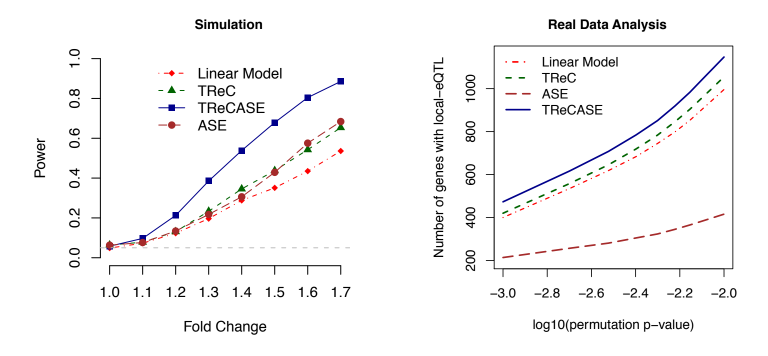
Real data analysis of 65 HapMap YRI samples (Pickrell et al. (2010) Nature 464, 768-772)

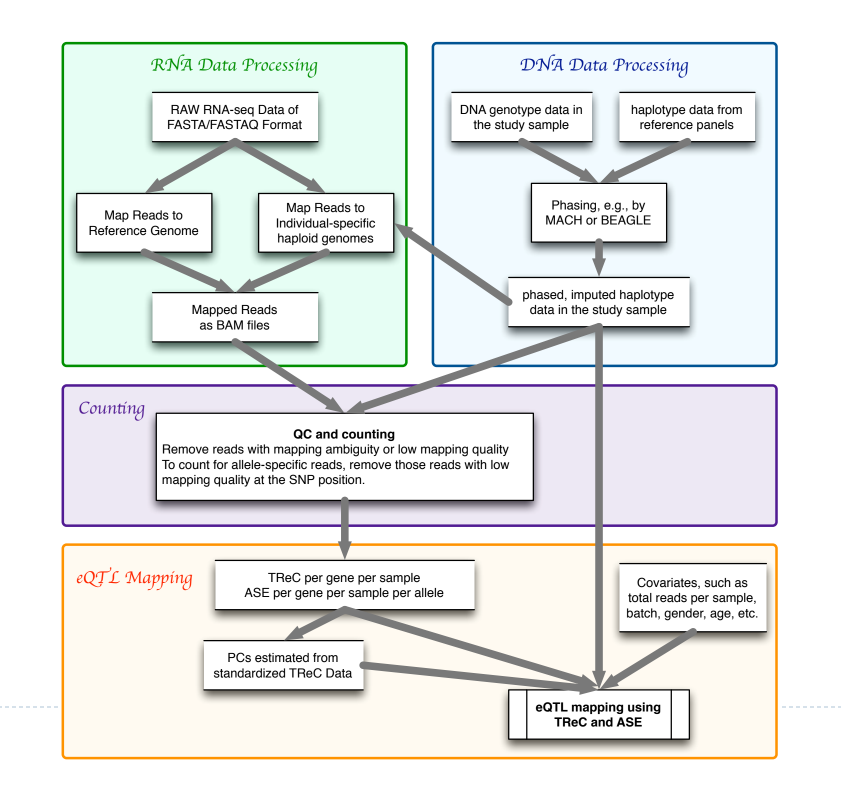


Sun (2012), A Statistical Framework for eQTL mapping using RNA-seq, Biometrics, 68(1):1-11

Real data analysis of 65 HapMap YRI samples (Pickrell et al. (2010) Nature 464, 768-772)

TReC:Total Read Count, TReCASE=TReC+ASE





#### http://www.bios.unc.edu/~weisun/software/asSeq.pdf

asSeq: A set of tools for the study of allele-specific RNA-seq data

Wei Sun and Vasyl Zhabotynsky

February 22, 2013

#### 1 Overview

#### > library(asSeq)

This vignette describes how to use R/asSeq to perform eQTL mapping using total expression and/or allele-specific expression, including a pipeline for input data preparation.

```
# ------
# 3. getUnique and filtering
# ------
prepareBAM(bamF, sprintf("%s_sorted_by_name", sami), sortIt=FALSE)
```

#### 5.3 Extract the allele-specific reads

R function extractAsReads from our R package asSeq can be used to extract allele-specific sequence reads. For example, the command can be

```
extractAsReads(input, snpList, outTag="myOutput", prop.cut=.5, min.snpQ=10, phred=33)
```

```
chr1 1019668 G A chr1 1020428 T C chr1 1020496 A G chr1 1029889 C T
```

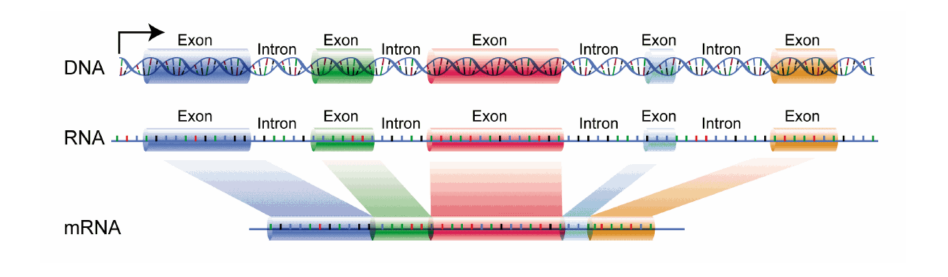
#### Outline

- ▶ I.An brief introduction to RNAseq
- ▶ 2. Allele-specific expression
- ▶ 3. Isoform-specific expression

# Transcription - splicing

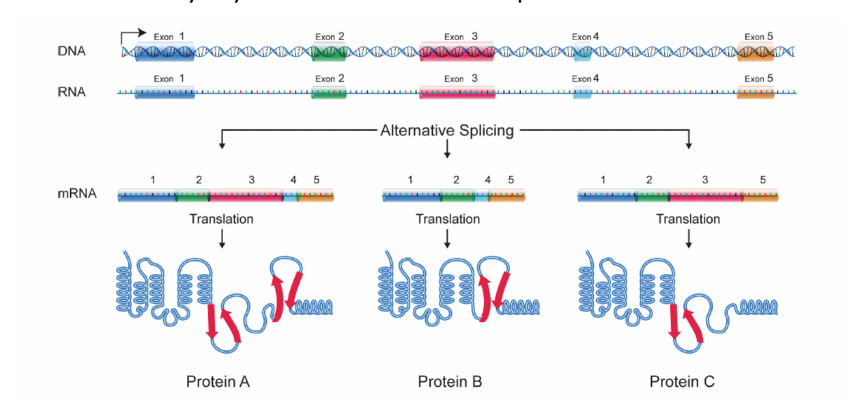
Transcription: DNA → RNA

Splicing: RNA → messenger RNA



## Alternative Splicing

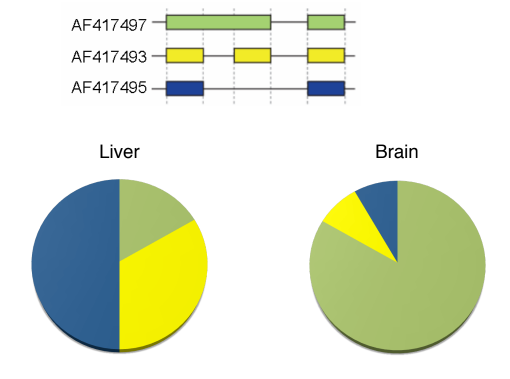
Alternative splicing is a process by which the exons of the RNA are reconnected in multiple ways during RNA splicing. The resulting different mRNAs are **RNA Isoforms** and they may be translated into different protein isoforms.



## Isoform-specific expression

#### Differential Isoform usage

the relative expression of an isoform with respect to the total expression of the corresponding gene



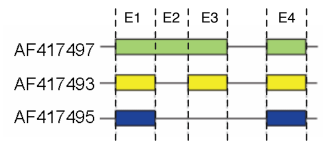
# Differential Isoform Usage

- ► Cufflinks, CuffDiff (Trapnell et al. 2010, NatBiotechnol. 28(5):511-5)
  - Estimate RNA-isoform expression, followed by differential isoform usage testing
- FDM (Singh et al. 2011 bioinformatics vol. 27 2633-40)
  - Testing for differential isoform usage using spliced read alignment, without estimating RNA-isoform expression
- Both CuffDiff and FDM perform two sample test
  - Case vs. Control, treatment vs. placebo
  - Testing for differential isoform usage vs. a continuous variable? e.g., additive genotype, dosage, patient survival time.

## Isoform-specific expression

#### **Construct non-overlapping exons**

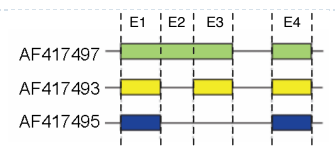
Assume the exon locations and lengths are known, but they may be overlapping with each other. Construct a new set of exons that are non-overlapping



For each gene, count the number of RNA-seq fragments for each exon set

Exon Set	Count
EI	50
E2	20
E3	30
E4	50
E1:E2	10
E2:E3	10
E1:E3	5
E3:E4	18
EI:E4	15

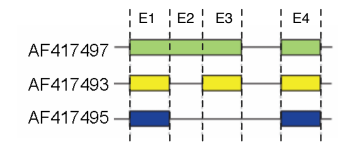
# RNA-isoform selection and abundance estimation



Exon Set	Count
EI	50
E2	20
E3	30
E4	50
E1:E2	10
E2:E3	10
E1:E3	5
E3:E4	18
EI:E4	15

If the expression are all from isoform AF417497, What is the pattern of read count across exon sets?

# RNA-isoform selection and abundance estimation



$$y = b_1 x_1 + b_2 x_2 + b_3 x_3$$

Exon Set	Count
EI	50
E2	20
E3	30
E4	50
E1:E2	10
E2:E3	10
E1:E3	5
E3:E4	18
EI:E4	15

AF417497	AF417493	AF417495
1	1	1
I	0	0
Ī	I	0
I	1	1
I	0	0
I	0	0
0.1	1	0
I	1	0
0.01	0.02	1

#### IsoDOT

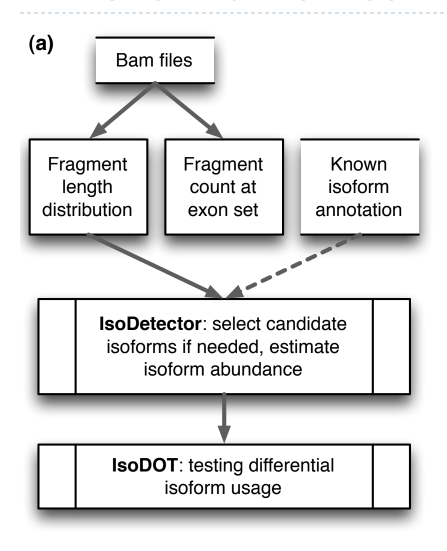
#### Challenges

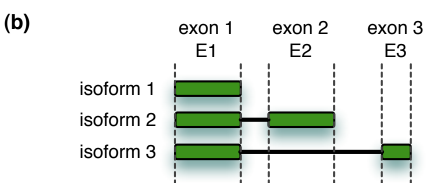
- This is a very challenging (negative binomial) regression problem because
- There is a huge number of possible isoforms, especially when there is no complete annotation on the RNA isoforms of the gene of interest.
- There can be strong correlations of the expression pattern of several isoforms.

#### Solutions

- Select a initial group of candidate isoforms.
- Apply penalized regression.

#### An overview of IsoDOT





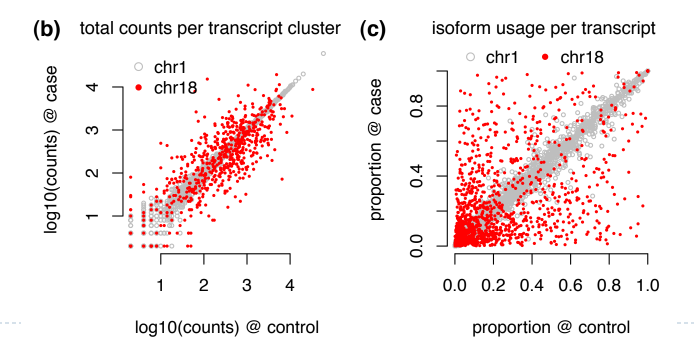
(c)					
(0)	Exon Set	Count	Isoform 1	Isoform 2	Isoform 3
	E1	50	$\eta_{1;1}$	$\eta_{1;2}$	$\eta_{1;3}$
	E2	20	0	$\eta_{2;2}$	0
	E3	10	0	0	$\eta_{3;3}$
	E1:E2	10	0	$\eta_{1,2;2}$	0
	E1:E3	5	0	0	$\eta_{1,3;3}$
	E2:E3	0	0	0	0
	E1:E2:E3	0	0	0	0

$$\mu = x_1b_1 + x_2b_2 + x_3b_3$$
  
y ~ NB( $\mu$ ,  $\phi$ )

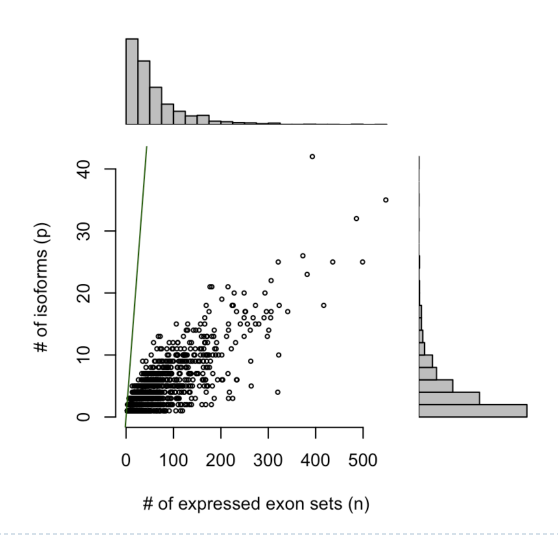
#### Simulation

We simulated two independent sets of ~2 million 76bp paired-end RNA-seq reads by Flux Simulator, using the transcriptome annotation of chromosome I and I8 of mouse genome.

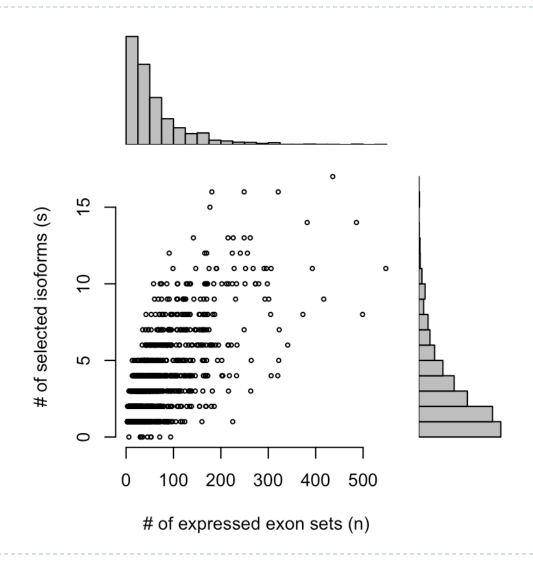
A case and a control sample were generates such as all the genes of chrl are equivalently expressed between the case and the control and all the genes of chrl8 are differentially expressed between the case and the control.



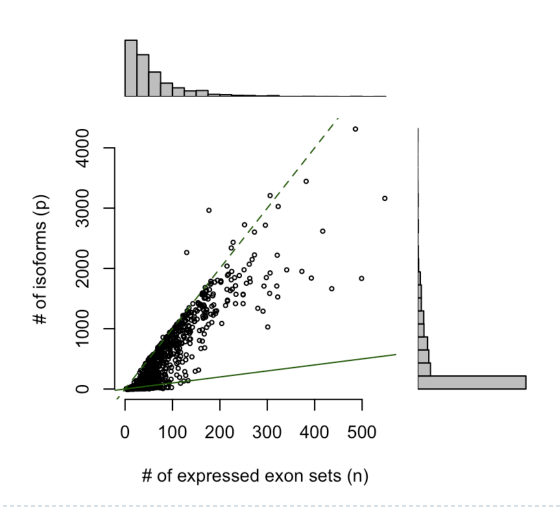
# The dimension of this problem: known isoforms



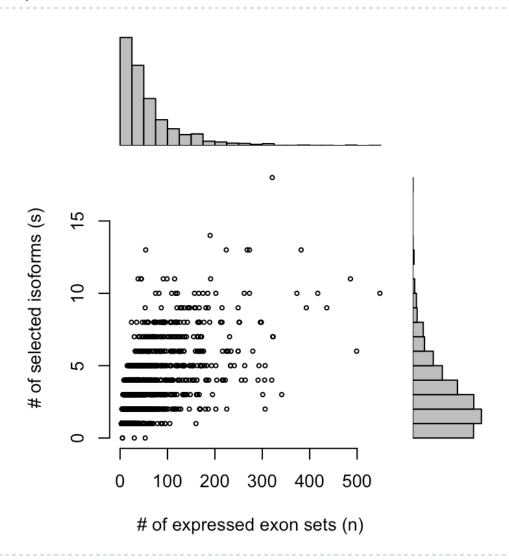
# The dimension of this problem: known isoforms, after isoform selection



# The dimension of this problem: unknown isoforms



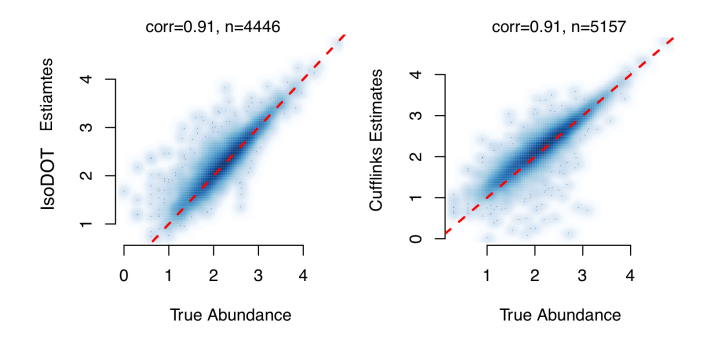
# The dimension of this problem: unknown isoforms, after isoform selection



#### IsoDOT

Compare IsoDOT with Cufflinks in terms of RNA isoform abundance estimation, using simulated data from flux-simulator, with isoform annotation

IsoDOT and cufflinks have comparable performance

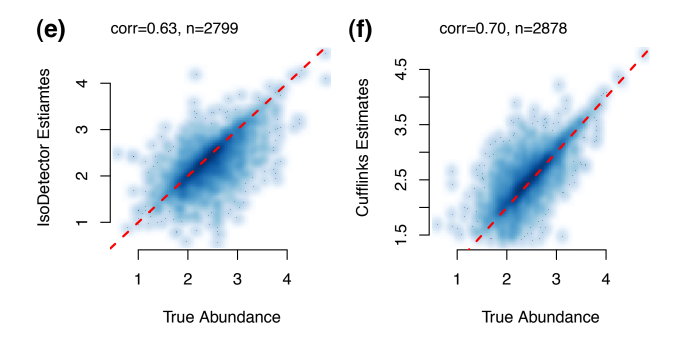


Sun et al. (2012), manuscript

#### IsoDOT

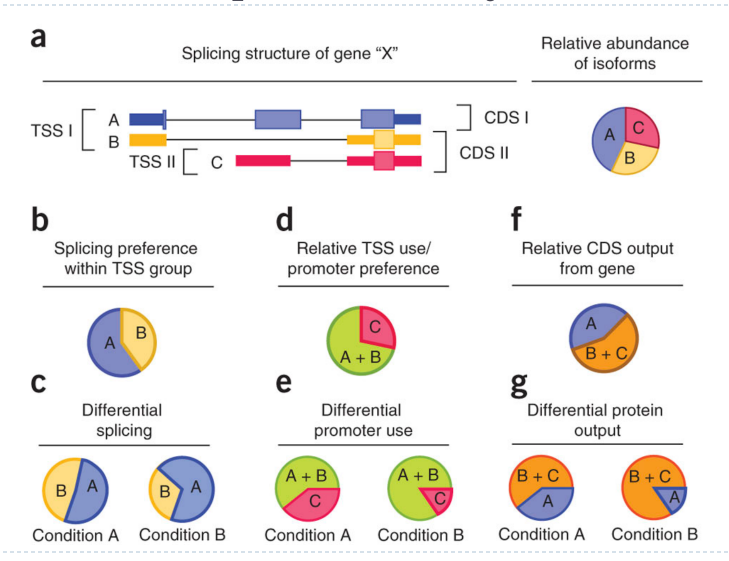
Compare IsoDOT with Cufflinks in terms of RNA isoform abundance estimation, using simulated data from flux-simulator, without isoform annotation

IsoDOT and cufflinks have comparable performance

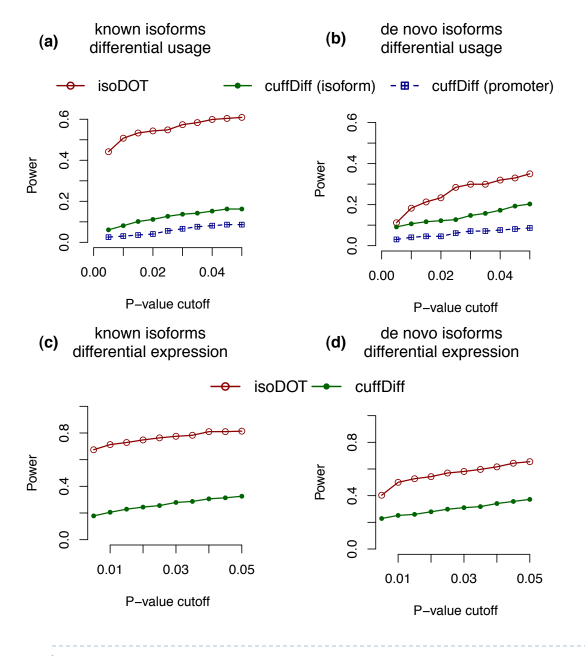


Sun et al. (2012), manuscript

### Different tests performed by Cufflinks



Nat Protoc. 2012 Mar1;7(3):562-78.

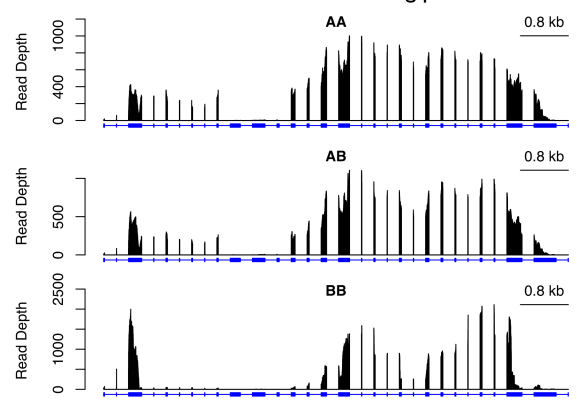


Test for differential isoform usage using simulated RNA-seq data from one case vs. one control

# Isoform-specific eQTL

Real data analysis on 60 CEU HapMap samples

ENSG0000065978, YBX I, Y box binding protein I



#### http://www.bios.unc.edu/~weisun/software/isoform\_files/isoform.pdf

isoform: A set of tools for RNA isoform study using RNA-seq data.

Wei Sun

February 22, 2013

#### 1 Overview

#### > library(isoform)

This vignette describes how to use R/isoform to estimate isoform abundance and assess differential isoform usage between cases and controls using an example. We are working on applying this method to assess differential isoform usage with respect to continuous covariate, e.g., isoform-specific eQTL mapping. Such examples will be added soon.

sorting and some basic QC using function prepareBAM from R package asSeq.

```
cmd2 = sprintf("samtools sort -n %s %s_sorted_by_name", bami, sami)
system(cmd2)
bamF = sprintf("%s_sorted_by_name.bam", sami)
prepareBAM(bamF, sprintf("%s_sorted_by_name", sami), sortIt=FALSE)
```

The following set of codes can be used to count the number of reads per exon set.

```
library(isoform)
```

```
bedFile = "Mus_musculus.NCBIM37.67.nonoverlap.exon.bed"
bamFile = "mm9_simu_set1_paired_reads_sorted_by_name.bam"
ctFile = "mm9_simu_set1_counts_paired_reads.txt"
```

countReads(bamFile, bedFile, ctFile)