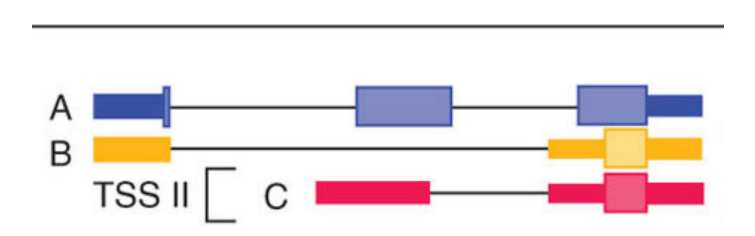
# Tuxedo Pipeline for Novel Transcript Discovery

Dhivya Arasappan

# What is a gene? What is a transcript?

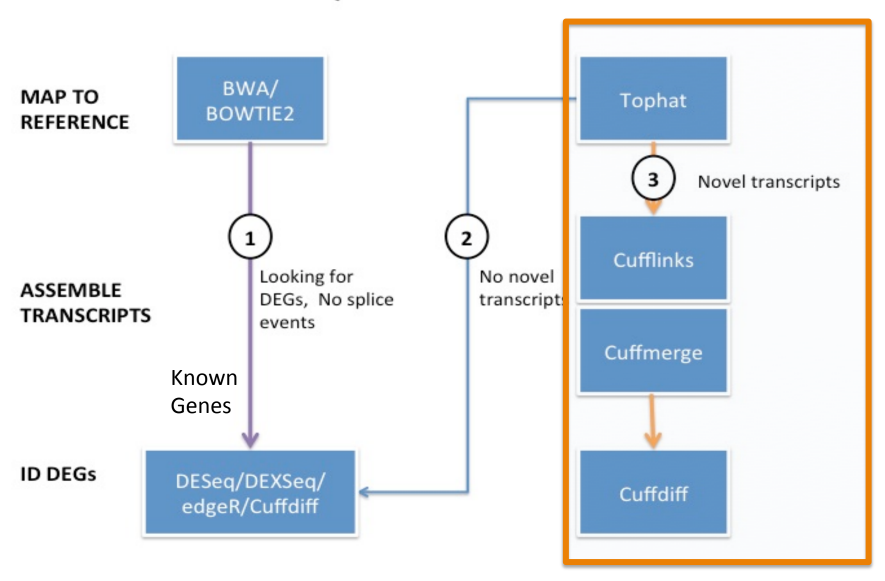
A gene can have multiple transcripts!



 We want to identify all these transcripts, whether annotated or not.

# Back to the big picture...

### **RNA-SEQ ANALYSIS PIPELINES**



# **Tophat** Novel transcripts Cufflinks Cuffmerge Cuffdiff

### **TUXEDO PIPELINE**



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



Cufflinks package

Cufflinks

Assembles transcripts

Cuffcompare

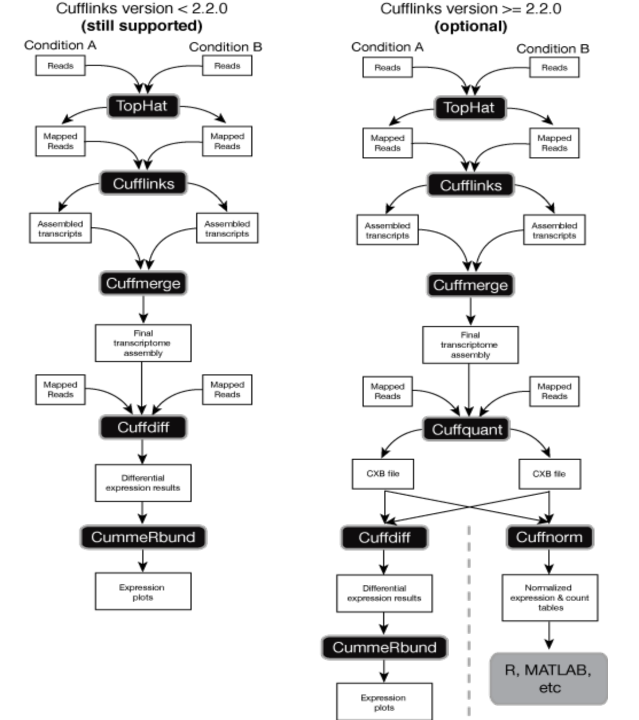
Compares transcript assemblies to annotation

Cuffmerge

Merges two or more transcript assemblies

Cuffdiff

Finds differentially expressed genes and transcripts
Detects differential splicing and promoter use



The pipeline is sequential.

Output of one step becomes input of next step.

Figure from: Trapnell et al, Nature protocols, 2012.

# Of course, Tuxedo Pipeline can be run without looking for novel events

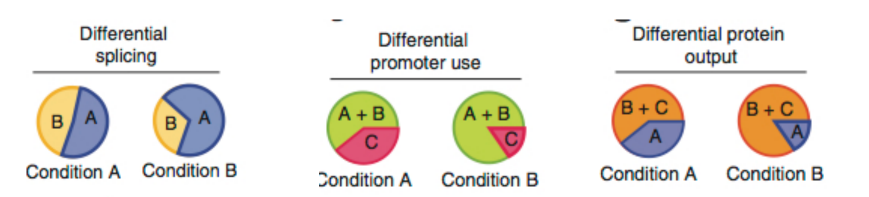
- NO NOVEL JUNCTIONS: Simple differential gene expression analysis against a set of known transcripts.
  - User provides a gff/gtf file containing annotated features. Quantify only the annotated features and id DEGs.
- NOVEL JUNCTIONS ALSO: In addition to known transcripts, novel transcripts should be explored.
  - User provides a gff/gtf file containing annotated features. But you also allow the search for novel variants as well. Both annotated and novel variants are quantified and DEGs are identified.
- ONLY NOVEL/DE NOVO JUNCTIONS: No gff/gtf file is provided.
  Using just the read data and the genome reference, construct
  de novo transcripts, quantify them and id DEGs.

### What do we get at the end of running this pipeline?

A view of how the transcriptome is different between condition C1 and condition C2

- Both in terms of annotated genes and transcripts.
- And novel genes and transcripts

Differential gene expression and so much more...



### **STEP 1: TOPHAT**

What does Tophat do?

Tophat maps your data to your reference in a transcriptome-aware manner, that will also identify junctions. We've already looked at how you can turn on and off its ability to identify novel junctions.

no-novel-juncs	Only look for reads across junctions indicated in the supplied GFF file.
-G <gtf file="" gff3=""></gtf>	Supply TopHat with a set of gene model annotations and/or known transcripts, as a GTF 2.2 or GFF3 formatted file.

# **STEP 2: CUFFLINKS**

What does cufflinks do?

TRANSCRIPT ASSEMBLY (also referred to as transcriptome reconstruction)

Let's see first what that entails.

# Why transcript assembly?

**Transcript assembly** = assembly of mapped reads into transcriptional units.

### Why?

- Define a precise map of all transcripts expressed in a sample.
- How does our transcriptome look in comparison to the known transcriptome?
- Look for novel transcripts between conditions/ samples.
- Look for differences in expression for these novel transcripts between conditions/samples.

# Why is transcript assembly hard?

Difficult to tell which read came from which transcript

- Many Short reads, many transcripts!
- Transcripts are expressed in different amounts.
   So, coverage of reads can be vastly different.
- Reads can come from mature mRNA (exons only) and precursor RNA (containing partial introns).

Table 1 | Selected list of RNA-seq analysis programs

Class	Category	Package	Notes	Uses	Input
Read mapping	. 200.000 112.000				
Unspliced aligners <sup>a</sup>	Seed methods	Short-read mapping package (SHRiMP) <sup>41</sup>	Smith-Waterman extension	Aligning reads to a reference transcriptome	Reads and reference transcriptome
		Stampy <sup>39</sup>	Probabilistic model		
	Burrows-Wheeler	Bowtie <sup>43</sup>			
	transform methods	BWA <sup>44</sup>	Incorporates quality scores		
Spliced aligners	Exon-first methods	MapSplice <sup>52</sup>	Works with multiple unspliced	Aligning reads to a reference genome. Allows	Reads and reference genome
		SpliceMap <sup>50</sup>	aligners		
		TopHat <sup>51</sup>	Uses Bowtie alignments	for the identification of	
	Seed-extend methods	GSNAP <sup>53</sup>	Can use SNP databases	novel splice junctions	
		QPALMA <sup>54</sup>	Smith-Waterman for large gaps		
Transcriptome r	econstruction				
Genome-guided Exon identification	Exon identification	G.Mor.Se	Assembles exons	Identifying novel transcripts using a known reference genome	Alignments to reference genome
reconstruction	Genome-guided assembly	Scripture <sup>28</sup>	Reports all isoforms		
ā		Cufflinks <sup>29</sup>	Reports a minimal set of isoforms		
Genome-	Genome-independent	Velvet <sup>61</sup>	Reports all isoforms	Identifying novel genes and transcript isoforms without a known reference genome	Reads
independent reconstruction	assembly	TransABySS <sup>56</sup>			100 mg
Expression quan	tification				
Expression quantification	Gene quantification	Alexa-seq <sup>47</sup>	Quantifies using differentially included exons	Quantifying gene expression	Reads and transcript models
		Enhanced read analysis of gene expression (ERANGE) <sup>20</sup>	Quantifies using union of exons		
		Normalization by expected uniquely mappable area (NEUMA) <sup>82</sup>	Quantifies using unique reads		
	Isoform quantification	Cufflinks <sup>29</sup>	Maximum likelihood estimation of		Read alignments to isoforms
		MISO <sup>33</sup>	relative isoform expression		
		RNA-seq by expectaion maximization (RSEM) <sup>69</sup>			
Differential		Cuffdiff <sup>29</sup>	Uses isoform levels in analysis	Identifying differentially	Read alignments and transcript models
expression		DegSeq <sup>79</sup>	Uses a normal distribution	expressed genes or transcript isoforms	
		EdgeR <sup>77</sup>			
		Differential Expression	Fig	gure :	
		analysis of count data	<u> </u>		
		(DESeq) <sup>78</sup>		rber et al, Nature M	ethods, 2011
		Myrna <sup>75</sup>	Cloud-based permutation method		

#### Most commonly used, if you have a genome.

#### Less resource-intensive

We'll call this graph based approach

Transcriptome reconstruction						
Genome-guided	Exon identification	G.Mor.Se	Assembles exons	Identifying novel transcripts		
reconstruction	Genome-guided	Scripture <sup>28</sup>	Reports all isoforms	using a known reference		
	assembly	Cufflinks <sup>29</sup>	Reports a minimal set of isoforms	genome		
Genome-	Genome-independent	Velvet <sup>61</sup>	Reports all isoforms	Identifying novel genes and		
independent reconstruction	assembly	TransABySS <sup>56</sup>		transcript isoforms without a known reference genome		

If you don't have a genome.

If you believe your sample has major rearrangements

More CPU and RAM intensive

Figure:
Garber et al, Nature Methods, 2011

# Genome guided transcript assesmbly

### Different assembly methods

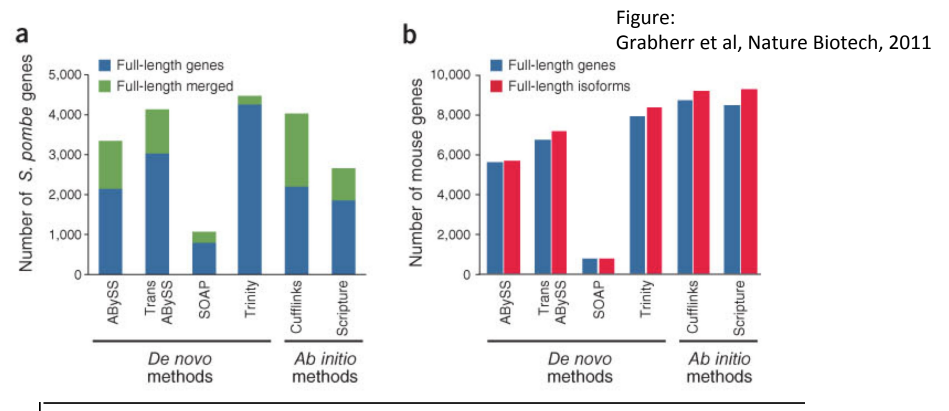
### Exon Identification Method

- First ID putative exons by looking for coverage islands.
- Older method, were meant for shorter read lengths.
- G.MorSe

### Graph Based approach

- Directly uses mappings of spliced reads to reconstruct transcriptome.
- Uses graph topology.
- Cufflinks (part of tuxedo suite), scripture

# How do these tools compare?



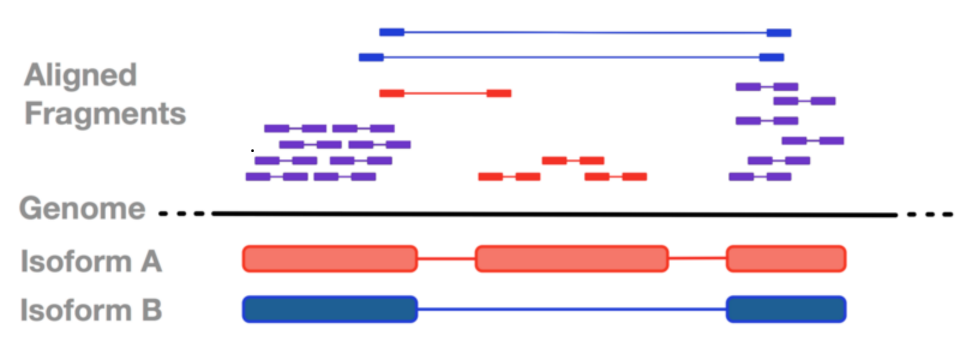
Program combination	vs. orthology annotation		vs. EST annotation	
	Base-level accuracy (%) <sup>1</sup>	Confirmed junctions (%) <sup>1</sup>	Base-level accuracy (%) <sup>1</sup>	Confirmed junctions (%) <sup>1</sup>
TopHat + Cufflinks	83.9	75.8	68.9	63.0
GSNAP + Cufflinks	79.4	71.2	65.7	58.4
GSNAP + Cufflinks (subsample²)	80.3	72.7	60.2	66.3
TopHat + Scripture	70.3	67.9	60.8	62.5

Figure: Palmieri et al, PLOS One, 2012

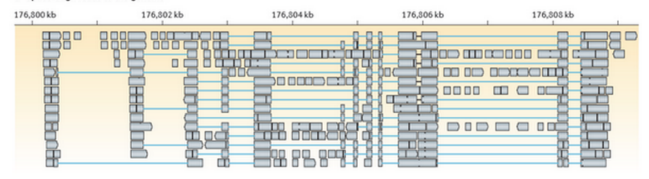
<sup>1</sup>Base level accuracy and percentage of confirmed junctions with different combinations of mapper and assembler on the sample ps94 males compared to the orthology annotation and the EST annotation (<sup>2</sup>based on 48 M reads). doi:10.1371/journal.pone.0046415.t001

### How does Cufflinks do transcript assembly

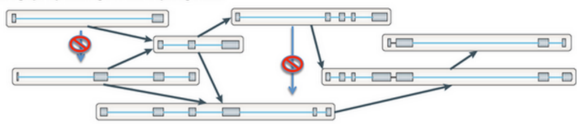
Graph based approach!



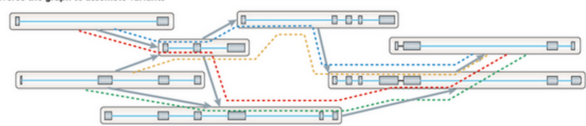
#### a Splice-align reads to the genome



#### b Build a graph representing alternative splicing events



#### C Traverse the graph to assemble variants



#### d Assembled isoforms

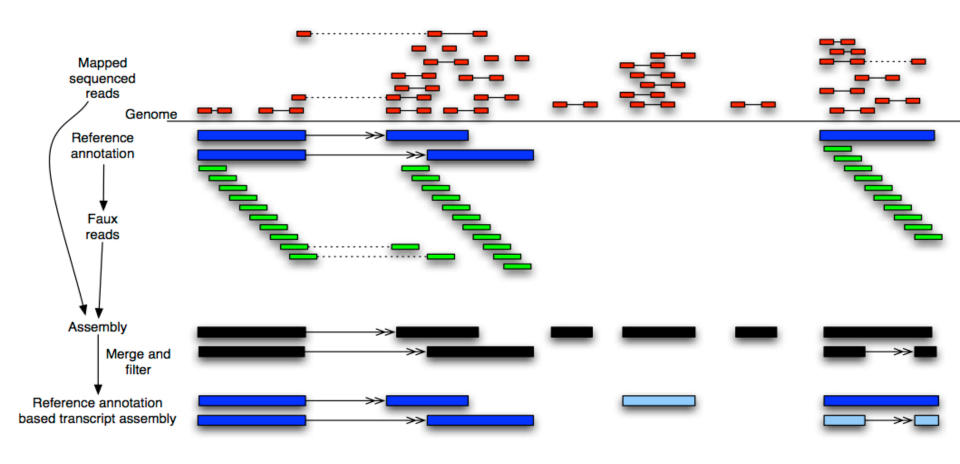


### Figure:

http://sourceforge.net/ projects/trinityrnaseq/files/ misc/RNASEQ\_WORKSHOP/ rnaseq\_workshop\_slides.pdf

## **RABT**

- Reference annotation based transcript assembly (RABT)
  - Uses existing annotation to guide assembly of transcripts.



# After assembly

Calculates abundance for these assembled transcripts.

- Normalized using FPKM (Fragments Per Kilobase of Exon Per Million) (variation of RPKM)
  - RPKM normalizes for transcript length variations and sequencing depth.
  - RPKM= (No.of Mapped reads\*10^9)/ (length of transcript \*total no.of reads)
  - FPKM just exchanges reads with fragments.

# General syntax for cufflinks command

```
cufflinks [options] <accepted_hits.bam>
```

Some of the important options:

```
-p/--num-threads
```

-G/--GTF (quantify only annotated transcripts)

-g/--GTF-guide (both annotated and novel transcripts)

-b/--frag-bias-correct

-u/--multi-read-correct

# General syntax for cufflinks command

-b/--frag-bias-correct

When quantifying abundance, corrects for sequence-specific bias at the ends of reads by 'learning' from the data.

-u/--multi-read-correct

By default, if a read maps to 2 genes it will count as 50% (half a read) towards each gene.

With this flag, it handles this question in a more fine-tuned manner.

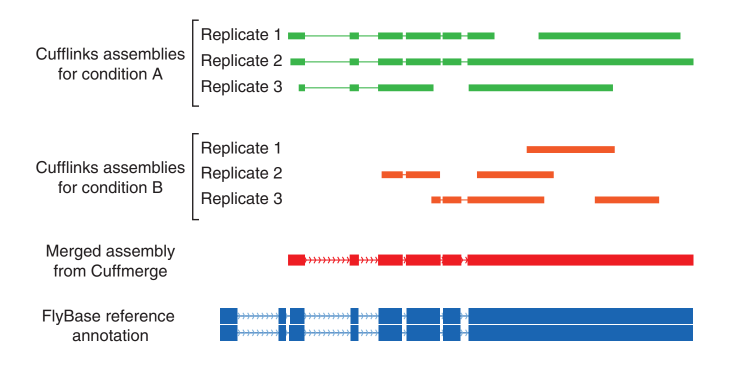
# Let's look at some results from a cufflinks transcript assembly

- Input:
  - Tophat mapped results (bam files)
  - Transcriptome annotation (genes.gtf)

 Let's SWITCH TO THE WIKI for instructions on looking at these results...

# **STEP 3: CUFFMERGE**

 Cuffmerge is used to merge all the transcripts that cufflinks assembled into one file.



# **STEP 3: CUFFMERGE**

- Input: All cufflinks assembly files (in gtf format)
- Input: Optionally: Annotated genes (in gff/gtf format)
- Compares your assembled transcripts to a reference annotation.
- Output: merged.gtf
  - Your very own gtf file, containing all the transcripts found in your samples (both novel and otherwise).
- SWITCH TO THE WIKI for instructions on viewing these results

# **STEP 4: CUFFQUANT**

- Optional but recommended step
- Computes gene and isoform expression quantification values and stores them in a structure that can be used by cuffdiff or cuffnorm.
- Input: result from cuffmerge
- Output: abundances files
- SWITCH TO THE WIKI for instructions on viewing these commands and results

# **STEP 5: CUFFDIFF**

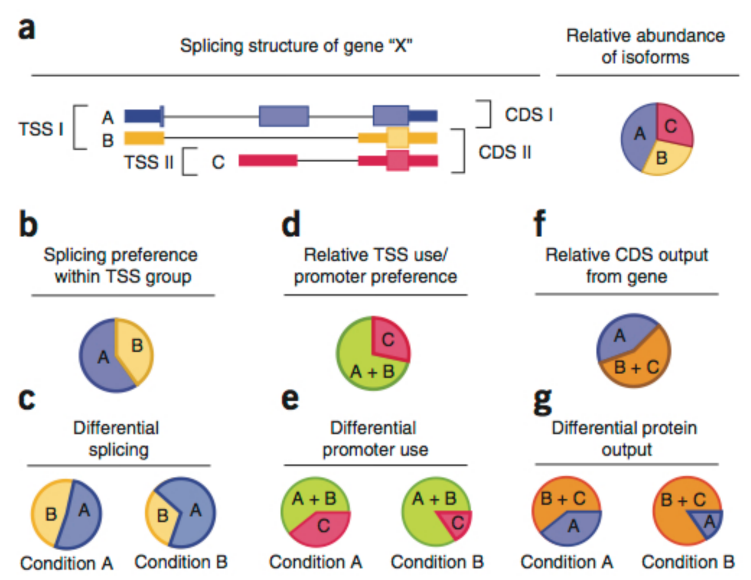
- Calculates differential expression!
- Input:
  - Our newly created merged.gtf file or a gtf file we downloaded (genes.gtf)
  - Our newly created cuffquant abundances file
  - Our newly created merged.gtf file or a gtf file we downloaded (genes.gtf)
  - Mapped bam files
- Counts the number of fragments(reads) generated by each isoform to obtain isoform-level expression.
- Calculates difference in isoform-level expression among conditions.
- If the chance of seeing this difference is small enough under the chosen statistical model, it is deemed signficantly differentially expressed.

# Other differential expression tools vs cuffdiff

Others	Cuffdiff
Raw count method for assigning counts to genes	Isoform deconvolution method for assigning counts to genes
Count the reads mapping to <b>exons</b> of each gene/normalization factor = expression for gene	Count the reads that map to each isoform of the gene/normalization factor = expression for gene
If all isoforms of the gene are up/down, works fine	If all isoforms of the gene are up/down, works fine
If some isoforms of the gene are up and some are down, inaccurate results	If some isoforms of the gene are up and some are down, works fine

### **STEP 5: CUFFDIFF**

Figure from: Differential analysis of gene regulation at transcript resolution with rNA-seq, Trapnell et al, Nature Biotechnology, 2013



# **STEP 5: CUFFDIFF**

 SWITCH TO THE WIKI for instructions on viewing these results

# Limitations of the Tuxedo Pipeline

A Reference is needed.

- Not quick.
  - For a human dataset with say, 60 million reads,
     each step can take 12-24 hours on lonestar and on stampede, probably 6-12 hours.
  - Some steps (cufflinks, cuffdiff) can run out of memory on large jobs.

## DESeq/edgeR output vs Tuxedo pipeline output

- Yesterday we generated differential expressed genes too. So, why the big fuss?
  - They were all from annotated genes. So, they all has flybase ids.
  - Now our output has genes with ids 'CUFF...' they are novel.
  - In addition to differential gene expression, we also have results for differential regulation.
  - We also have results telling us where our novel transcripts are with respect to the annotated ones.

### If You Don't Have A Genome

#### Transcriptome reconstruction

The second secon	A SO TO THE RESIDENCE OF THE PROPERTY OF THE P			
Genome-guided reconstruction	Exon identification	G.Mor.Se	Assembles exons	Identifying novel transcripts
	Genome-guided assembly	Scripture <sup>28</sup>	Scripture <sup>28</sup> Reports all isoforms	
		Cufflinks <sup>29</sup> Reports a minimal set of isoforms		genome
Genome-	Genome-independent	Velvet <sup>61</sup>	Reports all isoforms	Identifying novel genes and
independent reconstruction	assembly	TransABySS <sup>56</sup>		transcript isoforms without a known reference genome

- Trinity for assembly
  - Wrapper script to parallelize some parts of trinity:
     <a href="https://wikis.utexas.edu/display/bioiteam/assemble\_trinity">https://wikis.utexas.edu/display/bioiteam/assemble\_trinity</a>
  - assemble\_trinity -a <your\_allocation> -l <R1\_reads.fq> -r <R2\_reads.fq> -o <output\_directory>
- Annotate using trinotate or Blast2GO
- Map reads to the assembled transcriptome and simply quantify transcripts by parsing the SAM file:
  - cut –f 3 samfile|sort|uniq -c

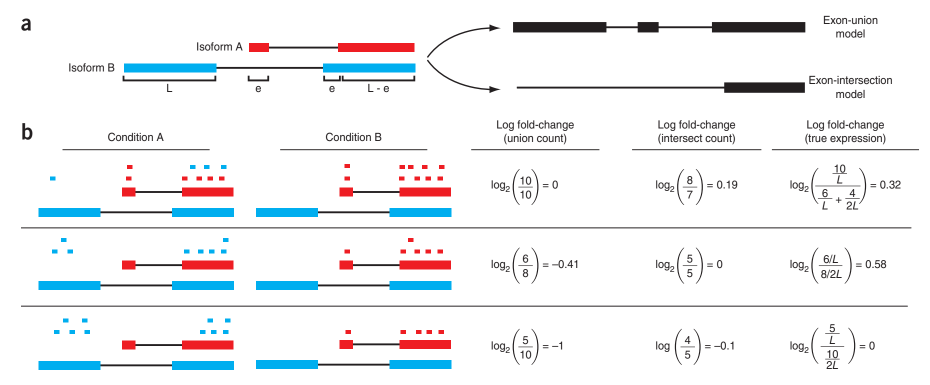


Figure 1 Changes in fragment count for a gene does not necessarily equal a change in expression. (a) Simple read-counting schemes sum the fragments incident on a gene's exons. The exon-union model counts reads falling on any of a gene's exons, whereas the exon-intersection model counts only reads on constitutive exons. (b) Both of the exon-union and exon-intersection counting schemes may incorrectly estimate a change in expression in genes with multiple isoforms. The true expression is estimated by the sum of the length-normalized isoform read counts. The discrepancy between a change in the union or intersection count and a change in gene expression is driven by a change in the abundance of the isoforms with respect to one another. In the top row, the gene generates the same number of reads in conditions A and B, but in condition B, all of the reads come from the shorter of the two isoforms, and thus the true expression for the gene is higher in condition B. The intersection count scheme underestimates the true change in gene expression, and the union scheme fails to detect the change entirely. In the middle row, the intersection count fails to detect a change driven by a shift in the dominant isoform for the gene. The union scheme detects a shift in the wrong direction. In the bottom row, the gene's expression is constant, but the isoforms undergo a complete

Figure from: Differential analysis of gene regulation at transcript resolution with rNA-seq, Trapnell et al, Nature Biotechnology, 2013