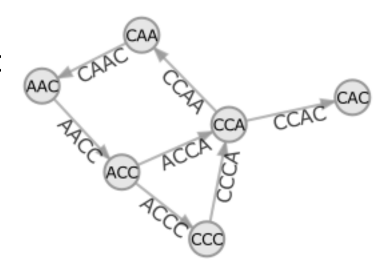
#### De novo transcriptome assembly

How it works, limitations and how to tell if your assembly is any good

#### De novo

- Latin expression meaning "from the beginning," "afresh," "anew," "beginning again."
- De novo, a term for any method that makes predictions about biological features using only a computational model without extrinsic comparison to existing data

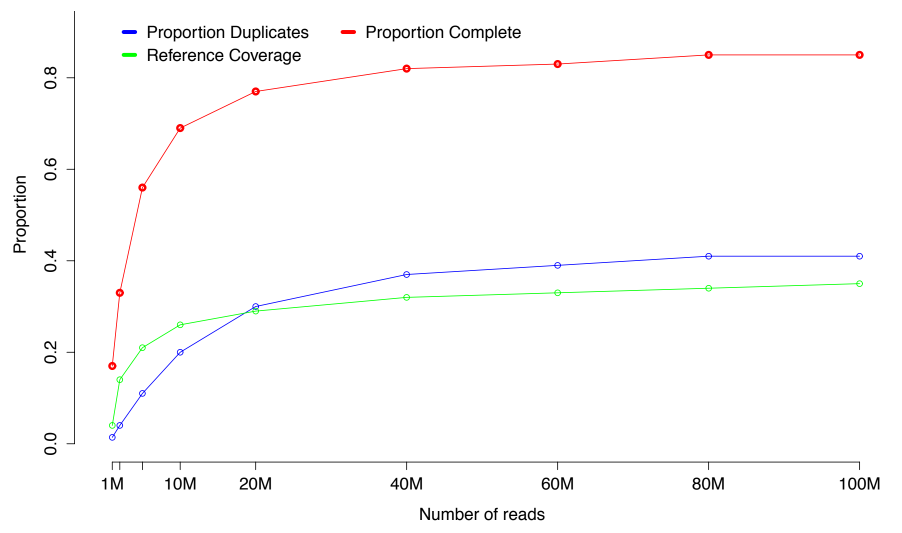


 No reference genome, then you must assemble your reads into genes

### Transcriptome vs Genome Assembly

- Transcriptome is easier
  - Smaller total volume of bases
  - Less low complexity regions
- Transcriptome is harder
  - Alternative splicing
  - Expression variability between tissues/cells
    - Difficult/impossible to fully sample all transcripts
    - Exponentially distributed coverage levels
    - Not the same in every cell
- Because of the very unique properties of transcriptome assemblies, it is important to use an assembler meant for transcriptomes (not genomes)

## How many reads?

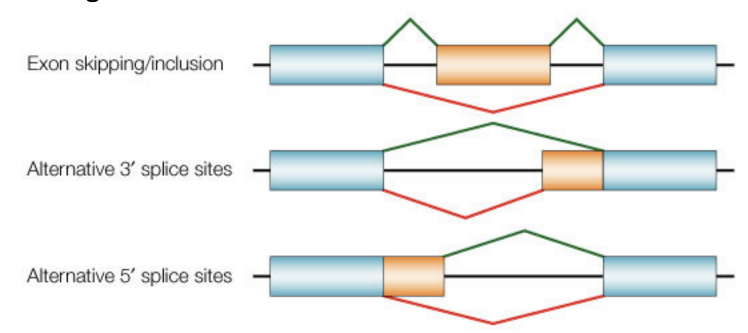


## How many individuals? How many tissues?

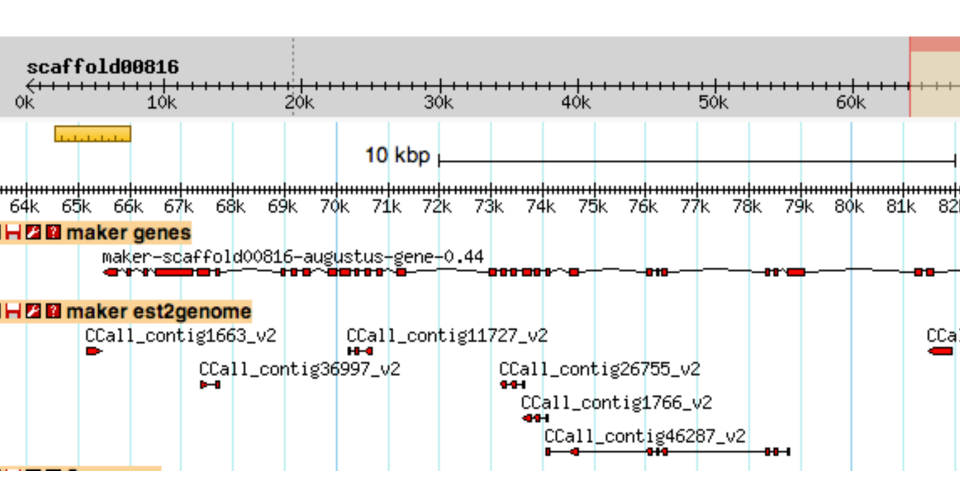
#### Problems with *de novo* assemblies

#### Results

- Highly fragmented assemblies
- Chimeras (can be biological, experimental or computational)
- Paralogs, alleles and alternative splicing variants combined or fragmented



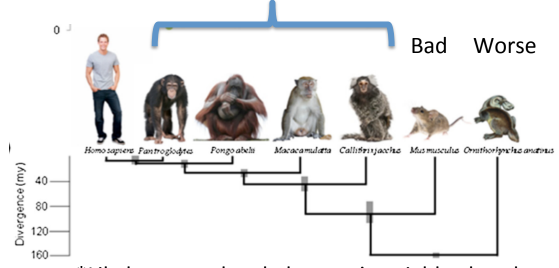
#### Chestnut



## Avoid an entirely *de novo* assembly - Use a distant or close relative

#### Reasonable Distant References

- Is there a close relative with a sequenced genome?
- How close is close enough?
  - Align then assemble
  - Assemble then align

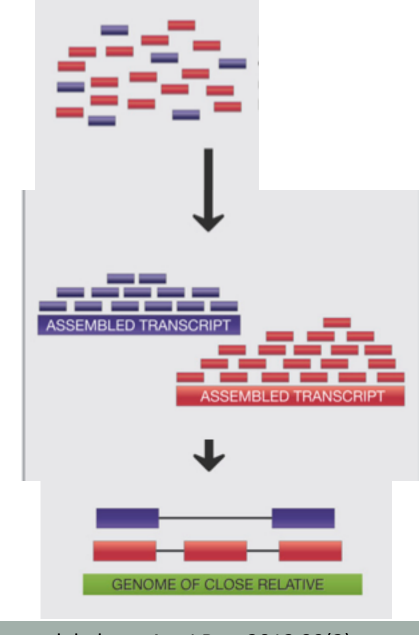


\*Likely to vary by phylogenetic neighborhood

Hornett and Wheat. Quantitative RNA-Seq analysis in non-model species: assessing transcriptome assemblies as a scaffold and the utility of evolutionary divergent genomic reference species. BMC Genomics 2012, 13:361

### Assemble then align

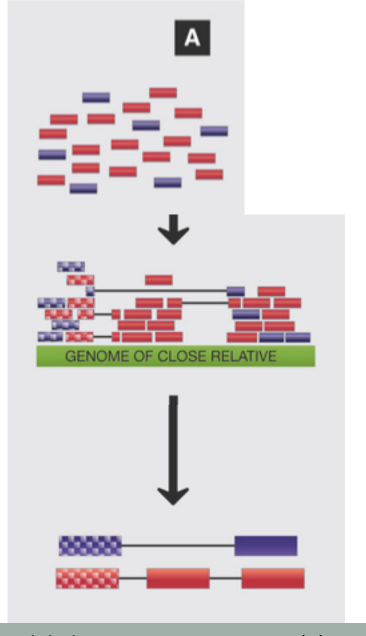
- First, assemble
- Next, align to a close relative
- Main Problems:
  - Fragmented assemblies gene pieces are scattered in a different consensus pieces
  - More difficult to sort out gene family members
- Main Advantages:
  - Alignment to a close relative can identify exon/exon boundaries (sort out alternative splicing)
  - Less bias can discover novel gene sequences



Weber et al. Strategies for transcriptome analysis in nonmodel plants Am J Bot. 2012 99(2): 267-276

### Align then assemble

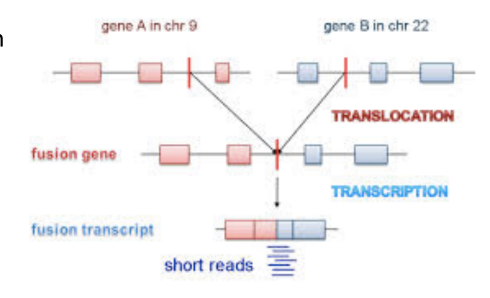
- First, map reads to (distant) reference
- Next, do local assemblies for each gene
- Main Problems
  - Read alignment may be poor due to lack of sequence similarity
  - Gene family expansion/contraction
- Main Advantage
  - Transcript assembly is less likely to be fragmented
  - Even where it is fragmented, you can identify all the fragments that originate from a single locus



Weber et al. Strategies for transcriptome analysis in nonmodel plants Am J Bot. 2012 99(2): 267-276

### De novo transcriptome assemblies

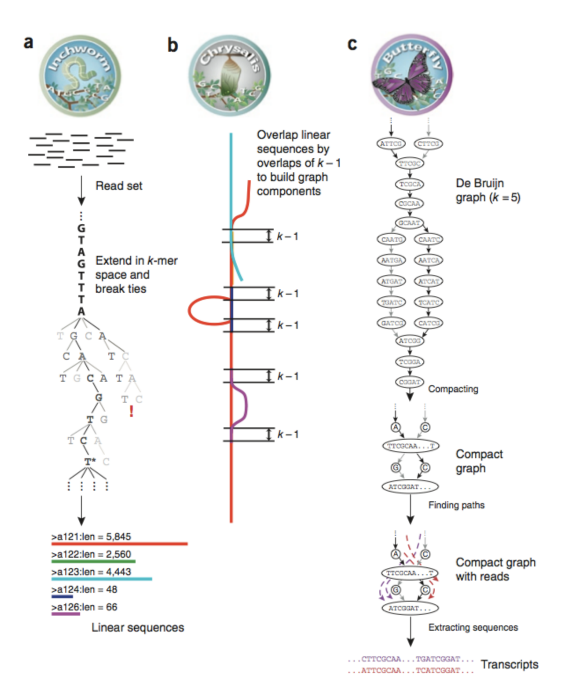
- Completely reference free
- What are they useful for?
  - Transcriptome characterization
  - Enabling proteomics experiments
  - Candidate gene discovery
  - Marker discovery/ development
  - Cancer or other tissues where fusion events are important
  - Metatranscriptomics surveying microbiota



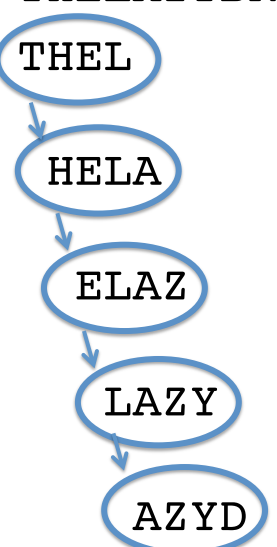
### Trinity strategy

#### Three stages

- 1. Inchworm
- 2. Crysalis
- 3. Butterfly

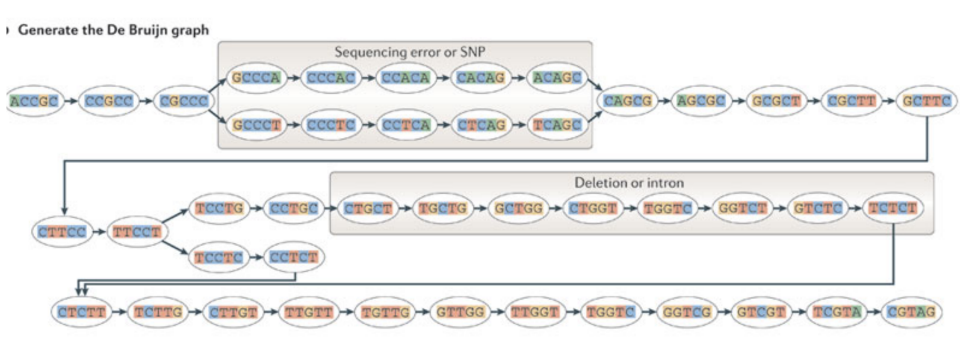


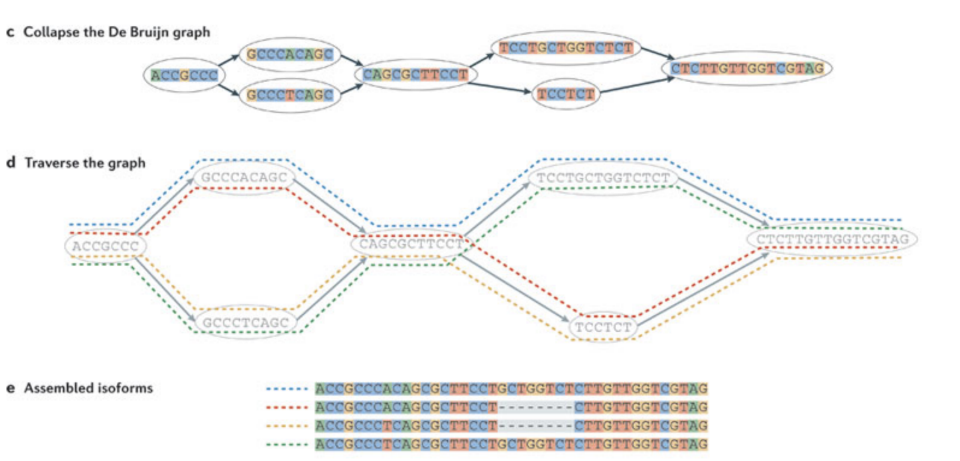
#### THELAZYBROWNDOG



## De Bruijn Graph

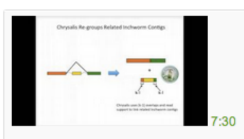
Directed acyclic graph of kmers

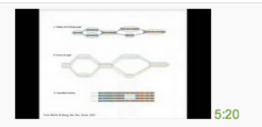




## A Collection of new RNA-Seq Videos from The Broad Institute

Posted by: RNA-Seq Blog Administrator In Presentations ( ) October 10, 2013 ( ) 1,134 Views





### BroadE: Trinity – How it works

BroadE: The General Approach to De novo RNA-Seq Assembly Using De Bruijn Graphs





BroadE: Introduction to De Novo RNA-Seq Assembly using Trinity

BroadE: Strand-specific RNA-Seq is Preferred

#### Videos!

http://www.rnaseqblog.com/acollection-of-new-rnaseq-videos-from-thebroad-institute/

# Trinity output – deciphering the naming

 An example Fasta entry for one of the transcripts is formatted like so:

>c115\_g5\_i1 len=247 path=[31015:0-148 23018:149-246]

Component –
a collection of
contigs that are
likely to be
derived from
alternative splice
forms or closely
related paralogs

Gene – best guess at an individual locus Isoform – alternative splicing events and alleles

These divisions are guesses only!

II. Improving the assembly and checking quality

# How to figure out if your assembly is good

#### BUSCO

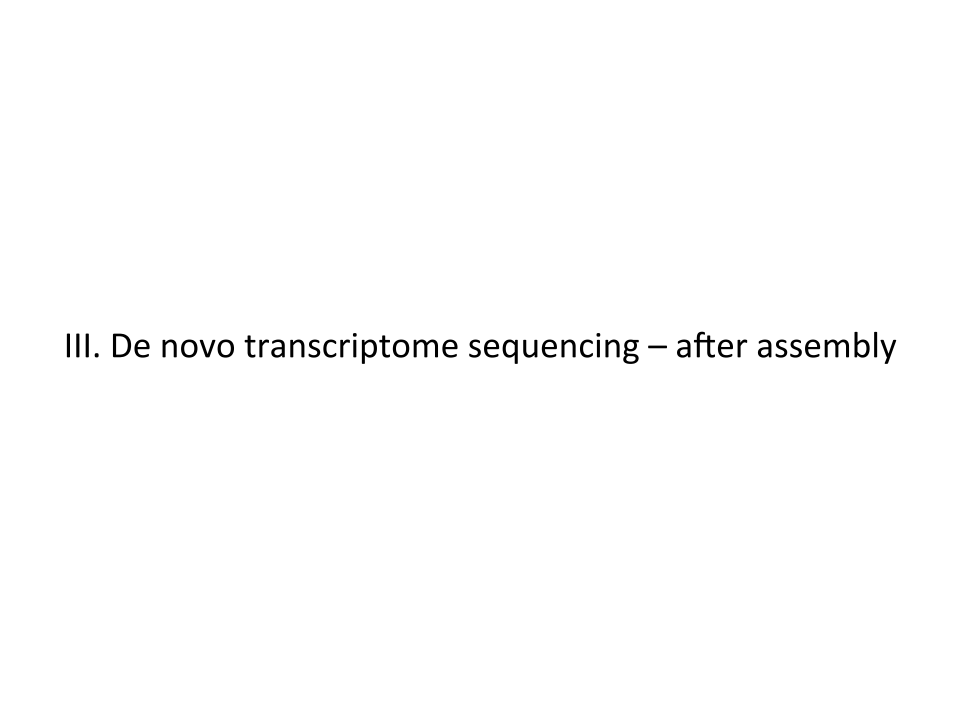
- Benchmarking Universal Single-Copy Orthologs
- based on evolutionarily informed expectations of gene content from near-universal single-copy orthologs selected from OrthoDB.
- Use to assess completeness of transcriptome
- <a href="http://busco.ezlab.org/">http://busco.ezlab.org/</a>



# How to figure out if your assembly is good

- Map reads back and see what % are captured in the assembly
- Transrate
  - analyses a transcriptome assembly in three key ways:
    - by inspecting the contig sequences
    - by mapping reads to the contigs and inspecting the alignments
    - by aligning the contigs against proteins or transcripts from a related species and inspecting the alignments





## ORF Finding - TransDecoder

- Searches all frames for ORFs, start codons and stop codons
- Maximizes length and log-likelihood score of ORF
- a single transcript can report multiple ORFs (allowing for operons, chimeras, etc).

