**High Performance Research Computing** 

A Resource for Research and Discovery



# Introduction to Genome/Transcriptome Assembly, using Sequencing Technologies

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## Outline

- Background
  - Sequencing
- Application of Sequencing in Research
  - Transcriptome assembly
  - Evaluations
  - Hands-on experiments



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## **Primary NGS Applications**

- 1. Alignment
- Today
   2. Assembly (no reference, with a reference)

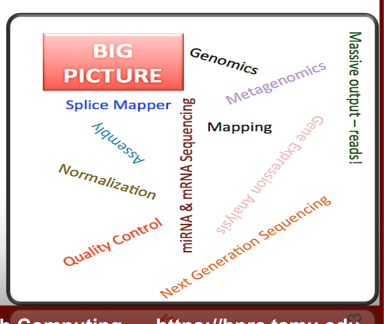
   Genome

   Transcriptome

Last week  $\rightarrow$  3. RNA-Seq

- Next Week  $\rightarrow$  4. Metagenomics
  - 5. ChIP-Seq

Next Week  $\rightarrow 6$ . RADSeq



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# Determining the sequence of nucleotides within a DNA (or RNA) fragment

- Ultimately completing the genome of non-model organisms, e.g. <u>Pacific whiteleg shrimp</u>
- Human genome project, \$3.8 Billion, 13 years to complete (1990-2003), 8-9x, coverage, 27 GBases

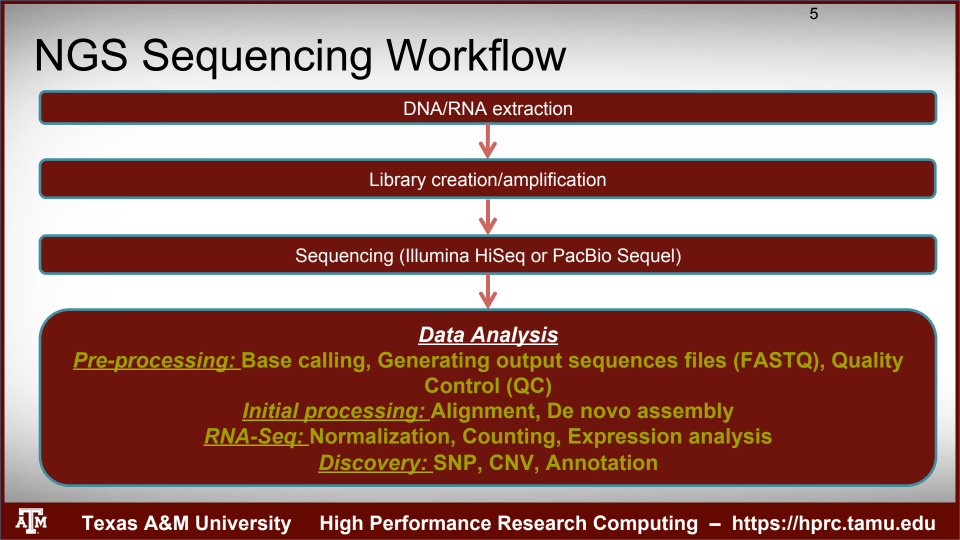
## <u>How?</u>

Using sequencing methods, such as Sanger sequencing, next generation sequencing and single-molecule techniques

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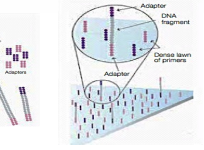
## SHORT READS

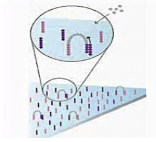
### • Illumina

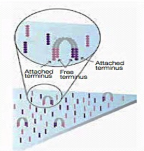
### Illumina next-generation sequencing

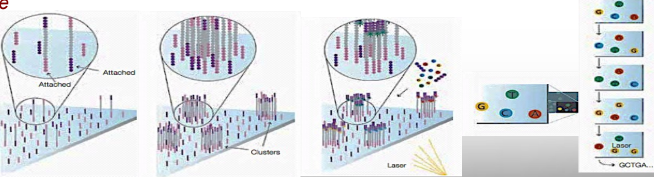
#### Sequencing by Synthesis (SBS) Technology

- Randomly shearing DNA
- Attaching DNA fragments to the flowcell surface
- Cluster generation, "Bridge Amplification"
- Adding four labelled reversible terminators, primers, and D polymerase
- Determining the attached nucleotide, based on the emitted fluorescence









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## Sequence and Quality Scores

**Quality scores** adapter sequence measure the G +В probability that a base = С 0 (a)sequence is called incorrectly. fragment D Α G E G G A flow-cell surface adapter sequence **Quality Score** Read

**Quality Score** 

- Illumina Quality Score
  - Phred-like algorithm: similar to scoring for Sanger sequencing

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- Quality score of a given base, Q, is defined as:
- e: estimated probability of the base call being wrong

$$Q = -10\log_{10}(e)$$

Quality Coore	Probability of Incorrect Base Call	
Quality Score	Call	Inferred Base Call Accuracy
10 (Q10)	1 in 10	90%
20 (Q20)	1 in 100	99%
30 (Q30)	1 in 1000	99.90%

#### **FASTQ** Format

#### 

Read 1

@HWI-EASXXX:96:96:1:1:7939:13150 1:N:0:

TTCTCCCCCCTTCTCCGTTTCATTCCACCCGCCCTATTCCTTCGCCTCCTCTTCCTTG

+

BEHBHGDA(DA>CCAEAHHHHGGHGHADCF@CDCE@EGGGDHH?HG@GGDGFGGGGE=

@HWI-EASXXX:96:96:1:1:14632:1706 1:N:0:

+

HHHHHFGD(GCGECGGHHHBDGEGGGGGGS>HFHDHBG2D8C>C)C-@D?;A>ECECAA0A=;+B0A?+;AD<@DB>5=A@@

Read 2

@HWI-EASXXX:96:96:1:1:7939:13150 2:N:0:

CAAGGAAGAGGAGGCGAAGGAATAGGGCGGGTGGAATGAAACGGAGAAGAGGGGGAGAA

+

4111166664@@@@@@@@@@@@@@@@@@;@44284477778+4666575228884444@

@HWI-EASXXX:96:96:1:1:14632:1706 2:N:0:

ACCTTCTCCTCCATCCTCTCCCCCCCCCTCCCCCTCTCTCTCTGTGACTCCTCCCCATTTCTTCTTCTTCTCGTG

F

-555598888@C@@C@@@@@@@@C444444@@@@@:40::6465689998@:@@@::4447677544:::@@;@@#######

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## **Choosing Illumina Sequencer!**



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http://core-genomics.blogspot.com/2016/01/meet-newest-members-of-family-miniseq.html

# LONG READS

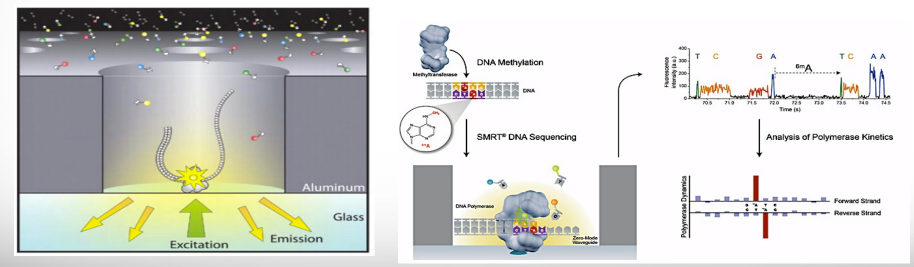
- Pacific Biosciences (PacBio)
- Oxford Nanopore Technologies MinION

### Long reads - PacBio

- Single Molecule Real Time Sequencing (SMRT) Methodology
- Fluorescent dyes

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- Zero Mode Waveguide (ZMW)
- DNA polymerase is immobilized at the bottom of a ZMW



http://www.nature.com/scientificamerican/journal/v294/n1/full/scientificamerican0106-46.html http://science.sciencemag.org/content/323/5910/133.full

### **PacBio Sequel**

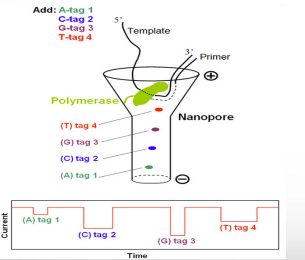


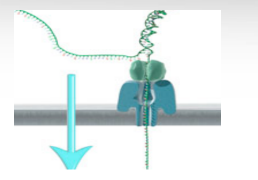
~10 GB per SMRT Cell 1M ZMW/SMRT Cell Up 16 SMRT/week 10 hour run time/SMRT Avg. read 10-15kb

~10x jump over RSII

## Long reads – Oxford NanoPore

- Oxford Nanopore Technologies
- Nanopore: a small hole (nanometer)
  - used to identify DNA sequence, passing through nanopore
- Single DNA molecule is sequenced





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Time in Second

Like electrophoresis, this technique draws DNA toward a positive charge. To get there, the molecule must cross a membrane by going through a pore whose narrowest diameter of 1.5 nanometers will allow only single-stranded DNA to pass [σ]. As the strand transits the pore, nucleotides block the opening momentarily, altering the membrane's electrical conductance, measured in picoamperes (pA). Physical differences between the four base types produce blockades of different degrees and durations (b). A close-up of a blockade event measurement shows a conductance change when a 150-nucleotide strand of a single base type passed through the pore [c].

DNA

**Refining this method** to improve its resolution to single bases could produce a sequence readout such as the hypothetical example at bottom (d) and yield a sequencing Single-stranded technique capable of reading a whole human genome in just 20 hours without expensive DNA copying steps and chemical reactions.

-15 120 pA 500 microsecond -15 pA Hypothetical readout

http://www.nature.com/scientificamerican/journal/v294/n1/full/scientificamerican0106-46.html http://www.kurzweilai.net/single-molecule-electronic-dna-sequencing#!prettyPhoto

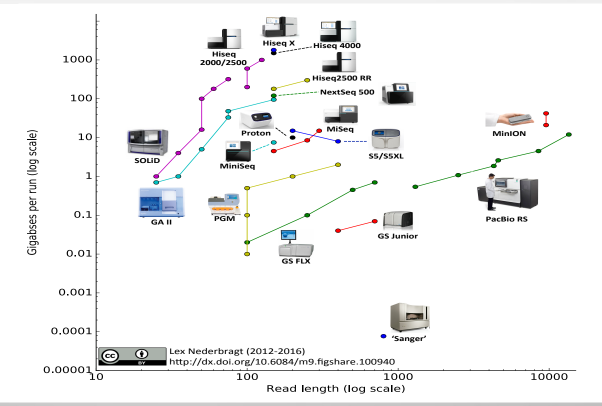




TAMU holds patent Dr. Higgin Bailey TEXAS A&M GRILIFE RESEARCH

## **NGS Read Specifications**

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Lex Nederbragt blog: https://flxlexblog.wordpress.com/2016/07/08/developments-in-high-throughput-sequencing-july-2016-edition/

## **Comparing Sequencing Technologies**

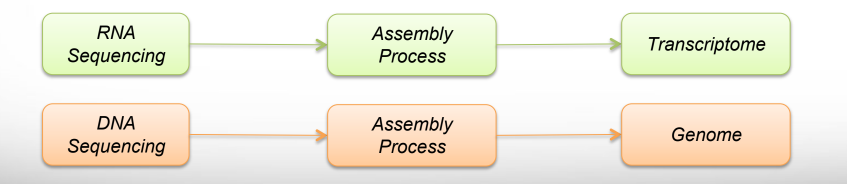
Platform	Read length	Error rates	Technology	Portable?
Illumina	< 400 bp	Low	Sequencing by synthesis	No
PacBio	~ 10-15 Kb	High	SMRT – ZMW	No
Oxford Nanopore Technologies	~ 5-8 Kb	High	Nanopore protein – strand sequencing	Yes

### Why assembly?

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#### Generating the consensus of transcriptome or genome of non-model species

Reconstructing the genome and transcriptome of non-model species are essential steps in expanding our understanding of the organism and developing therapeutic approaches to fight disease

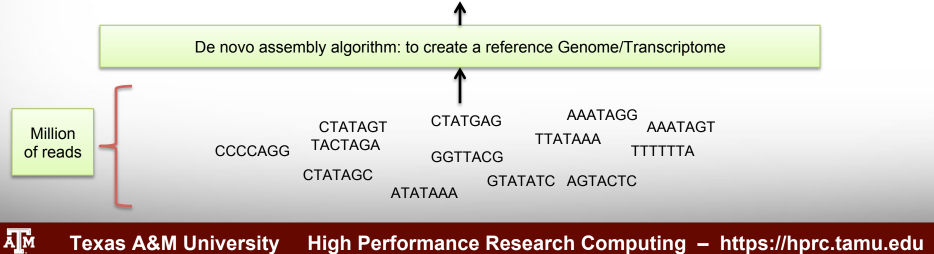


## De novo Assembly

- Pool of reads
- No Reference genome!
- Creating consensus from the reads

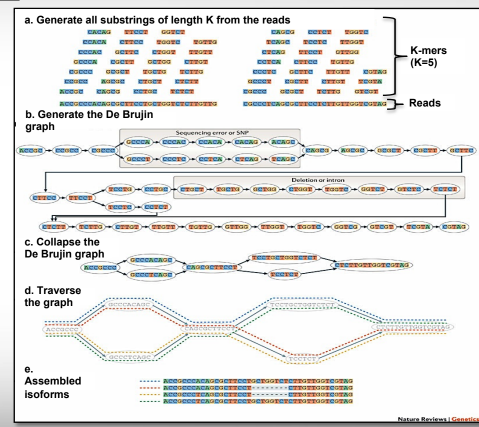
Consensus Genome/Transcriptome

Contig n: ...GATCTACCTATTTTAATCTATCTAGACCCATAAAAAAGTAAAAATTAGTAATTCTTAAGTAATATTAAGTATCGTGG...



## De novo Assembly - 2

- Connection reads by finding common sections of kmers
  - Kmers are made from reads!
- Resolving conflicts
- Complicated process!
- Highly computational resource demanding!



## De novo Assembly - 3

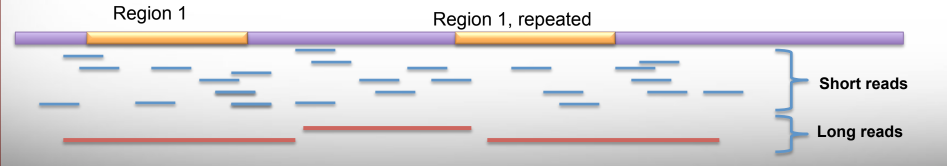
**Reference Genome Generation** 

- Goal: generating the reference genome for a new species, using the genomic DNA data, generated by NGS/Single Molecule platforms
- Main tool: *de novo* assembly algorithms
- Output: annotated reference genome

### De novo Assembly - 4

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Sanger	Next Generation Sequencing	
Low coverage depth	High coverage depth	
High cost for large genomes	Relatively low cost, even for large genomes	
Slow	Fast	
Handles repeats well	Need long reads for repeated regions (e.g. PacBio, Illumina Mate- Pair)	



## Genome Assembly Tools:

- ALLPATHS
- ALLPATHS-LG (Special recipe: fragments + jumping libraries)
- DISCOVAR de novo
- ABySS
- EULER-SR
- SOAPDenovo
- VCAKE
- Velvet

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- MaSuRCA
- Canu (will cover in practical portion today)
- CLC Bio Genomics Workbench

## Transcriptome Assembly Tools:

- SOAPdenovo-Trans
- Trans-ABySS
- Velvet + Oases
- Trinity (will cover in practical portion today)
- Rnnotator
- CLC Bio Genomics Workbench

### High Quality Assembly

- Hybrid Approach
- High Coverage
- Merging
  - Metassembler



## **Practical Portion**

# Logging in to the system

## • SSH (secure shell)

 The only program allowed for remote access; encrypted communication; freely available for Linux/Unix and Mac OS X hosts;

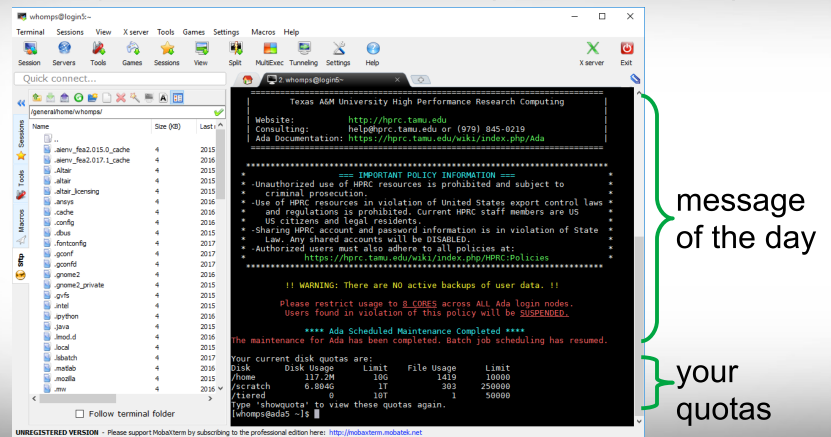
## • For Microsoft Windows PCs, use MobaXterm

- https://hprc.tamu.edu/wiki/HPRC:MobaXterm
  - You are able to view images and use GUI applications with MobaXterm
- or Putty
  - https://hprc.tamu.edu/wiki/HPRC:Access#Using\_PuTTY
    - You can not view images or use GUI applications with PuTTY

# Your Login Password

- Both state of Texas law and TAMU regulations prohibit the sharing and/or illegal use of computer passwords and accounts
- Don't write down passwords
- Don't choose easy to guess/crack passwords
- Change passwords frequently

## Using SSH - MobaXterm (on Windows)



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## Using SSH to Access Ada

#### ssh user\_NetID@ada.tamu.edu

#### https://hprc.tamu.edu/wiki/Ada:Access

You may see something like the following the first time you connect to the remote machine from your local machine: Host key not found from the list of known hosts. Are you sure you want to continue connecting (yes/no)?

Type yes, hit enter and you will then see the following: Host 'ada.tamu.edu' added to the list of known hosts. user\_NetID@ada.tamu.edu's password:

# Contact the HPRC Helpdesk

Website:

hprc.tamu.edu

Email:

help@hprc.tamu.edu

Telephone:

(979) 845-0219

### Help us, help you -- we need more info

- Which Cluster
- UserID/NetID
- •Job id(s) if any
- Location of your jobfile, input/output files
- •Application used if any
- Module(s) loaded if any
- •Error messages
- •Steps you have taken, so we can reproduce the problem

### Second self-organized official Software Carpentry Workshop at TAMU!

- Topics covered: Unix shell, Version Control with Git, R Programming
- Date: February 7-8, 2019, Time: 9:00 AM 5:00 PM
- Instructors: Noushin Ghaffari, Ramalingam Saravanan, David Bapst, and Shichen Wang
- Fee: \$35
- Lunch: provided (the fee includes the lunch, coffee and refreshments for both days)
- Application link: <u>goo.gl/forms/qQSwtO95EVeY007o2</u>
- Workshop webpage: <u>swang8.github.io/2019-01-24-tamu/</u>

# Any question? nghaffari@tamu.edu

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