## *Genome Assembly and Structural Variation Detection from MinION Nanopore Data*



## *Base Calling of Nanopore*



- Hidden Markov model
- Only four options per transition
- Pore type = distinct kmer length



- Form probabilistic path through measured states currents and transitions
	- e.g. Viterbi algorithm

Basecalling currently is performed at Amazon

a CGTAAGAGTACGTCCAGCATCGG-5 ATGGTA AGAGTGATA  $+29 +25$  $n = 0$ рA  $\mathbf b$ Ш 36  $2s$ 32 pA 28 24  $2C$ 30 Time (arbitrary units)<br>TTCTT<br>TTATT TCTTT<br>TATTT TTTTC TTTTT  $CTTTT$ TTTTT **TTTCT** TITTE **ATTTT TTTAT** TTTTA TTTTT



#### **1D and 2D Base Calling**

**The 1D vs 2D barcoding refers to whether the complementary strand is used to improve basecalled data. Basically – it gives two shots when examining the same loci. The advantage being that the complementary strand will have a different kmer profile.**

#### **Read Length Distribution – Ecoli and Yeast**



**Ecoli by UCSC:** [http://www.ebi.ac.uk/ena/data/view/ERS715551-ERS715552/](http://www.ebi.ac.uk/ena/data/view/ERS715551-ERS715552) Yeast by CSH : <http://labshare.cshl.edu/shares/schatzlab/www-data/nanocorr/>

# **Assembly Method**

#### **Sequencing reads:**



*1. Overlap graph*



 $\text{ACCTG}$   $\rightarrow$  CCTGA  $\rightarrow$  CTGAT $\rightarrow$  TGATC

 $\begin{array}{l} \text{AGCGA} \textcolor{red}{\blacktriangleright} \text{GGGAT} \textcolor{red}{\blacktriangleright} \text{CGAT} \end{array}$ 

 $\bigl\text{\rm GATCA}\bigl\rightarrow \text{\rm ATCAA}\bigl\rightarrow \text{\rm TCAAT}\bigl\rightarrow \text{\rm CAATG}\bigl\rightarrow \text{\rm AATGT}\bigl\rightarrow \text{\rm TGTGA}$ 

 $G\Delta$ 

 $C \times 6$ 

*2. de Bruijn graph*

*3. String graph*

#### *The Classic Overlap, Layout and Consensus Method*



2) Layout



3) Consensus

#### CCTATG-TAGTCAGTCG

#### **ATGCTAGTCAG**

#### GCTAGTCGGTCGATCTACC

#### CAGTCGATCTGCCGGT

#### GTCAGTC-ATCTAC-GGTTAGCATTGC

#### CCTATGCTAGTCAGTCGATCTACCGGTTAGCATTGC **Consensus**

## **The Greedy Graph Based Method**

**The greedy algorithms are implicit graph algorithms. They drastically simplify the graph by considering only the high-scoring edges. As an optimization, they may actually instantiate just one overlap for each read end they examine.** 



## **One Contig for The Ecoli Genome**



## *Missing Homoplymers Recovered by Nanopolish from the Event Data*



## *Assembly of Ecoli from Different Methods*



- **(i) Assemblies of 1,2,3 were obtained from ONT data only, while assembly 4 used both ONT and MiSeq reads;**
- **(ii) Assemblies of 1 and 2 were obtained after using nanopolish;**
- **(iii) \* - in Assembly 3, the indel information is the number, rather the bases;**
- **(iv) ^Loman NJ, Quick J, Simpson JT: A complete bacterial genome assembled** *de novo* **using only nanopore sequencing data.** *Nat Methods.* **2015; 12(8): 733–735.**

#### *Single Molecular Integrated Scaffolding (SMIS)*



*SMIS: http://sourceforge.net/projects/phusion2/files/smis/*

#### *Fake Mate Pairs from ONT Reads*



# ONT Assisted Scaffolding

*http://sourceforge.net/projects/phusion2/files/smis/*

Mate pair data is used to scaffold contigs. Contigs, and pairs of contigs connected by pairs, define a bidirectional graph:



Using expected insert size, a estimate of the gap size can be given for each contig.



## Saccharomyces cerevisiae complete genome

*Scaffold N50 858Kb ; Contig N50 330Kb* 



*Yeast W303 Assembly from PacBio Data using PBcB*

## **Data:**

[http://datasets.pacb.com.s3.amazonaws.com/2013/](http://datasets.pacb.com.s3.amazonaws.com/2013/Yeast/) [Yeast/](http://datasets.pacb.com.s3.amazonaws.com/2013/Yeast/)

- **33 contigs and N50 = 777023**
- **12 out of 17 chromosomes are covered with a single contig**
- **99.95 % identity compared with assembly from Miseq**
- **No major homoplymer problems!**

#### Table 3 CSHL W303 Yeast Illumina Reads Used for Assembly<sup>+</sup>



+The dataset was downloaded from <http://labshare.cshl.edu/shares/schatzlab/www-data/nanocorr/>

#### Table 4 W303 Yeast Assembly Stats



*SOAPdenovo\* - reads were processed and base errors corrected using our own tools;*

*SMIS-Merge+ - Scaffolding was performed using SMIS on the merged assembly and contigs were processed using our own tools.*

#### *Methods of Structural Variation Detection*





## *Split Reads – Identifying Breakpoints*



*Parsing the alignment CIGAR strings and looking for common breakpoints with hard or soft clipping "H" or "S"* 

#### *Normalising Insert Variation Factor*

There are N mate pairs of sequences which can be mapped to a reference chromosome. To quantify the likelihood of structural variation for a given pair, we define a normalised insert size variation factor:

$$
p_i = 1 - \left[\frac{c_i - c_{i-1}}{D_i}\right]^{0.3} \qquad 0 \le i < N \text{ and } 0 \le \frac{c_i - c_{i-1}}{D_i} \le 1
$$

where  $C_i$  - Mapping coordinate of the  $i^{th}$  pair on the chromosome;  $D_i$  - Insert size difference between the shredding distance and the value estimated from alignment;

It is seen from the above figures that the noise level of insert size variation was significantly reduced and this makes the detection much easier.

## *CNVs in Yeast Chr8 Comparison – SC288C vs W303*



# *Summary:*

- **Missing homoplymers is the major issue for de novo assembly;**
- **PacBio shows advantages in genome assembly, so far;**
- **Detection of structural variations is still a challenging task, while Oxford MinION data offers exciting chances.**



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