

## Relevant Vocabulary

- <u>Base pairs (bps):</u> nucleotides; ATCG
- <u>Genome</u>: the full collection of genetic material in the nucleus of an organism
- <u>Reads/sequences</u>: string of nucleotides representing a segment of the genome
- <u>DNA sequencing</u>: method to determine the order of nucleotides in a section of DNA
- <u>Genome assembly</u>: the process of reconstructing a genome's sequence using reads
- <u>Kmer</u>: a short sequence of nucleotides of length K (e.g. 5mer, 6mer)
- <u>SCK</u>: Single-Copy Kmer; a sequence that appears once in the genome • <u>Repetitive regions</u>: portions of the genome that appear more than once
- <u>Coverage</u>: the number of times a portion of DNA is sequenced
- <u>Pseudacris feriarum</u>: upland chorus frog, found in the SE United States
- Size of genome: 4.5 billion bases (gigabases, Gb)

## Background

### **Current genome assembly methods**<sup>1</sup>:

- 1. De Bruijn graph (**Fig 1**):
  - Stores all sequences and their connections in one large graph.
- Heuristic methods required to extract genome sequence from graph. 2. Overlap layout consensus method (Fig 2):
  - Use similarities between DNA sequences to create longer consensus sequences.
  - Repetition in large genomes makes it easy for this method to incorrectly overlay two
  - sequences that share a common region but are in different parts of the genome.

### **Problems:**

- Both methods are effective at assembling small genomes, but they do not scale well as genome size and data set size increase.
- Poor scaling occurs because repeats are not initially avoided.
- The *Pseudacris feriarum* genome is large (4.5 Gb) and has numerous repeats, making it difficult to assemble with current methods.

### **Solution:**

- Avoid the issues caused by repeats by
- 1) identifying SCKs (Single-Copy Kmers),
- 2) identifying their relative positions in the genome,







Figure 1 shows an example of a de Bruijn graph. The method breaks sequences into smaller kmers and tracks their connections. The graph contains the entire genome.

Figure 2 represents the Overlap Layout Consensus method. The method aligns reads based on common regions.



Figure 3 represents the pipeline for our proposed method. Our method relies on identifying SCKs, clustering SCKs, tracking their relative positions to one another, and then finally overlaying reads.

# New Genome Assembly Method

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### Methods

#### **1. Identify optimal length for SCKs**

- Two types of sequencing data
- Short, high-quality reads & long, low-quality reads • Issues
- Kmers need to be long enough to be unique in genome
- Kmers need to be short enough to minimize sequencing error effects ( $\sim 13\%$ )
- After intense study, we found 18 base pairs is the optimal kmer length.



Figure 4 shows the probability of observing a kmer 0-10 times by chance. By 18 base pairs long, it is extremely unlikely to see a given kmer by chance.

**Figure 5** shows the probability of finding at least 1 kmer in 25 reads without an error for the given kmer sizes. With an error rate of 0.13 and a kmer size of 18, we expect to see almost every SCK at least once.

#### 2. SCK Distribution

- Identified homozygous and heterozygous peaks using 18mer coverage values.
- Chose homozygous SCKs to avoid phasing issues.
- Selected 18mers within the coverage range (50x-58x), determined using the observed SCK homozygous peak and avoiding overlap with the heterozygous peak (**Fig 6**).



#### **3. Final SCK Selection**

- First filtered SCKs by short, high-quality reads. Selected initial SCKs that appeared within the desired coverage and did not contain homo-polymers likely resulting from sequencing error.
- SCKs from previous step were then filtered using long, low-quality reads. With a 13% sequencing error, each SCK is expected to appear in only 2 reads. To make sure that only SCKs are selected, only kmers that appear 2-6 times in the long reads were selected.

### 4. Chosen SCKs

• After the above steps, we obtained 11,813,926 SCKs to be used for assembly



Figure 6 represents the observed distribution of 18mers over different coverage values. We wrote a method to identify two distinct kmer peaks: heterozygous (left) and homozygous (right). The chosen SCK coverage range (50x-58x) is highlighted in gray.



Figure 7 depicts a simulated representation of long reads from a region of a chromosome. The ticks represent sequencing error in the reads while the dots represent error-free SCKs. To maximize use of long reads, reads will have to be aligned based on shared SCKs.

#### Next steps:

- chromosome.
- grouping of non-repetitive regions.

#### **Computational Costs:**

#### **Future considerations and implications:**

- successfully re-assemble other genomes.

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2. Hotaling, S., Kelley, J. L., & Frandsen, P. B. (2021). Toward a genome sequence for every animal: Where are we now? Proceedings of the National Academy of Sciences, 118(52), e2109019118. https://doi.org/10.1073/pnas.2109019118



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• Use chosen SCK set to estimate relative positions of SCKs along each

• Previous linking using short reads on mouse chromosomes allowed for

• Use linking data and long reads to span large repeat regions.

• Our goal is to reduce the computational cost of genome assembly since current methods require large amounts of disk space and/or RAM. • Currently our kmer selection process can be performed in 7 hours using 300 gigabytes of RAM. Additional steps are expected to be efficient.

• Use previously-assembled genomes to determine if proposed method can

• Only 1% of eukaryotic genomes have currently been assembled<sup>2</sup>. There is need for a new, faster, and computationally-cheaper method.

### References

### Acknowledgements