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Chromium sequencing: the doors open for genomics of obligate plant pathogens

Alistair R McTaggart1,2, Tuan A Duong1,1, Vang Quy Le3, Louise S Shuey1, Werner Smidt1, Sanushka Naidoo1, Michael J Wingfield1 and Brenda D Wingfield*,1
1Department of Biochemistry, Genetics & Microbiology, Tree Protection Co-operative Programme (TPCP), Forestry & Agricultural Biotechnology Institute (FABI), University of Pretoria, South Africa; 2Queensland Alliance for Agriculture & Food Innovation, The University of Queensland, Ecosciences Precinct, Brisbane, Queensland, Australia; 3Section of Molecular Diagnostics, Aalborg University Hospital, Denmark

It is challenging to sequence and assemble genomes of obligate plant pathogens and microorganisms because of limited amounts of DNA, comparatively large genomes and high numbers of repeat regions. We sequenced the 1.2 gigabase genome of an obligate rust fungus, *Austropuccinia psidii*, the cause of rust on Myrtaceae, with a Chromium 10X library. This technology has mostly been applied for single-cell sequencing in immunological studies of mammals. We compared scaffolds of a genome assembled from the Chromium library with one assembled from combined paired-end and mate-pair libraries, sequenced with Illumina HiSeq. Chromium 10X provided a superior assembly, in terms of number of scaffolds, N50 and number of genes recovered. It required less DNA than other methods and was sequenced and assembled at a lower cost. Chromium sequencing could provide a solution to sequence and assemble genomes of obligate plant pathogens where the amount of available DNA is a limiting factor.

Obligate pathogens cannot be cultured. Consequently, there is often insufficient DNA available for whole-genome sequencing. This problem is reflected by the small number of genomes sequenced for obligate fungi in the 1K fungal genomes project [1,2]. Limited DNA quantity is an obstacle to study genomics of fungi and plant pathogens such as those from marine environments, obligate lichen associates, insect associates, downy mildews, powdery mildews, rust fungi and some obligate smut fungi [3,4].

Most genome-sequencing technologies require large amounts of DNA for preparation of a library. For example, 2–7 μg of DNA is recommended for preparation of paired-end and mate-pair libraries for sequencing on an Illumina platform. Long-read platforms, such as PacBio and Nanopore, require 1–15 μg per cell for sequencing; however, many cells are needed for significant sequencing depth of large genomes. Larger amounts of DNA are difficult to obtain for microorganisms that cannot be isolated and cultured. Genomes that have been sequenced for obligate microorganisms are often those of well-studied models, for which material can be cultivated, for example poplar and wheat rust [5].

Rust fungi are one of the most speciose groups of obligate fungal plant pathogens, with genomes that are repeat rich and amongst the largest in the Fungal Kingdom [6]. Despite their diversity and significance as plant pathogens, relatively few genomes of rust fungi have been sequenced. This is partly because of the challenges posed by their obligate nature [3].

There are five near-complete, publicly available genomes of rust fungi from cereal grasses and poplar, assembled from 454 and Illumina sequencing [5,7–11]. However, five other projects have not yet published their assembled genomes of rust fungi because of difficulties encountered from size, repeat content and different haplotypes in the genome [12–16].

Long-read sequencing platforms, such as PacBio and NanoPore, may offer solutions to sequence and assemble repetitive genomes of rust fungi [3]. For example, PacBio sequencing was recently tested to assemble and phase dikaryotic genomes of two rust fungi [17,18]. This research showed that long-read sequencing could resolve challenges for genome assembly of dikaryotic taxa. However, if DNA quantity is a limiting factor for nonmodel plant pathogens, these technologies cannot provide sequencing coverage depth, especially for organisms with large genomes.

Chromium sequencing, developed by 10X Genomics (www.10xgenomics.com), has provided a low-cost approach to assemble human genomes from limited DNA.

**METHOD SUMMARY**

We sequenced and assembled the genome of an obligate fungus to compare Chromium sequencing technology with paired-end and mate-pair libraries. This was the first application of Chromium sequencing for an obligate fungal pathogen, in which DNA quantity is a limiting factor for genomics.
starting material [19]. Small amounts, as low as 1 ng, of high-quality DNA are fragmented into ~100-kb molecules, which are then barcode and amplified within micelles to prepare a Chromium library. Homologous regions of DNA are unlikely to be present in the same micelle, which allows haplotypes to be phased using their micelle-specific barcodes. After amplification, the barcoded library is sequenced on an Illumina platform, and reads are assembled with Supernova Software from 10X Genomics.

The medical community has largely adopted Chromium sequencing for genomics and transcriptomics; however, at least three studies have used it to assemble, scaffold and phase genomes of plants [20–22]. Chromium sequencing has not yet been tested for assembly of fungal genomes; there would be few advantages to sequence haploid and culturable fungi with this approach because there is no need to phase haplotypes and quantity of DNA is not limiting. However, obligate dikaryotic plant pathogens with large genome sizes are perfect candidates to sequence, assemble and phase genomes with Chromium sequencing.

We tested whether Chromium 10X provided a competitive platform for sequencing and de novo genome assembly of a rust pathogen, *Austropuccinia psidii*, here used as a model for other microorganisms. The genome assembly from a Chromium library was compared with a combined assembly from one paired-end and one mate-pair library sequenced on Illumina.

**Materials & methods**

**Selection of isolate & DNA extraction**

To obtain enough material from a single genotype of *A. psidii*, a single-pustule isolate was grown on *Syzygium jambos* to extract DNA from approximately 500 mg of urediniospores. This is a technically challenging and laborious process that is not always possible with other species of rust fungi. The urediniospores were ground in liquid N₂ and DNA was extracted following the protocol of a Machery Nagel NucleoSpin® Plant II DNA extraction kit. The taxon selected, *A. psidii*, has an unreleased genome sequenced with paired-end reads on Illumina MiSeq [16], and a publicly available genome, approximately half-complete, sequenced with paired-end reads on Illumina HiSeq [23].

**Genome sequencing**

Genomic DNA was sent to Macrogen Korea for library preparation and sequencing. Three libraries were prepared, a Chromium 10X library, a mate-pair library with a 5-kb insert, and a paired-end library with a 350-bp insert. Macrogen prepared the paired-end and mate-pair libraries from 2 and 7 µg of DNA, respectively (~100 and 350 mg of spores), and the Chromium 10X library from 1 µg of DNA (~50 mg of spores) with fragments above 40 kb size-selecting using BluePippin (Sage Science Inc.). The Chromium 10X and paired-end libraries were run on Illumina HiSeqX, and the mate-pair library was run on Illumina HiSeq4000.

**Genome assembly**

**Assembly of Chromium 10X sequencing data**

A single run of a Chromium library on a HiSeq X Ten yielded 812.77 million 150-bp reads (~120 gigabytes of raw data). The Chromium data were processed (quality filtering and trimming) and assembled using Supernova v 1.2.2 from 10X Genomics [19]. The Supernova run on barcoded micelles resulted in a weighted mean molecule size of 24.47 kb. The assembly yielded 1443 long scaffolds (scaffolds ≥10 kb), with a total scaffold size of 74,872 Mb (only scaffolds ≥10 kb) and a N50 value of 81.25 kb. The final assembly was exported using the Supernova mkoutput function, with minimum size of scaffolds set to 500 bp, and this assembly has been deposited in GenBank (accession: QOUH00000000).

**Assembly of paired-end & mate-pair sequencing data**

Runs on Illumina HiSeq X Ten and HiSeq4000 yielded 43 gigabytes of paired-end and 29 gigabytes of mate-pair data, both of which were 150-bp reads. Trimomatic v 0.36 [24] and NxTrim v 0.4.2 [25] were used with the default options to trim paired-end and mate-pair reads. We used ABySS 2.0 [26] to assemble (k-mer = 99) paired-end reads, and to scaffold with mate-pair reads. Only scaffolds ≥500 bp were retained. The statistics obtained from the assembly with ABySS were reported in Table 1.

**Comparison of Supernova & ABySS assemblies**

We used the NUCmer function in MUMmer [27] to compare the 18 largest scaffolds (>5 Mb) assembled by Supernova (from Chromium reads) to their homologous scaffolds (>5 Mb) assembled by ABySS (from combined paired-end and mate-pair reads). We used Circos [28] to visualize the homology between 18 scaffolds from Supernova and 220 scaffolds from ABySS (Figure 1).

**Results & discussion**

The first challenge to sequence genomes of obligate pathogens is the availability of DNA for preparation of a library. Chromium sequencing overcame this hurdle in the present study where a library was prepared from 1 µg of DNA. This amount of DNA was obtained from approximately 50 mg of spores. Long-read sequencing technologies and paired-end and mate-pair libraries are prepared from larger amounts of DNA; they are not practical options to obtain sequencing coverage depth when DNA quantity is a limiting factor.

Repeat-rich regions present another challenge to sequence and assemble genomes of obligate plant pathogens. Chromium 10X libraries outperformed combined paired-end and mate-pair libraries.

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**Table 1. Statistics of the genomes assembled from different sequencing libraries.**

<table>
<thead>
<tr>
<th>Assembler</th>
<th>Data used</th>
<th>Amount of starting material (µg)</th>
<th>Cost of data generation (€)</th>
<th>No. of scaffolds (&gt;500 bp)</th>
<th>N50 (kb)</th>
<th>L50</th>
<th>Estimated genome size (bp)</th>
<th>Completeness of genome (BUSCO fungi; %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supernova</td>
<td>Chromium</td>
<td>1</td>
<td>3000</td>
<td>147,937</td>
<td>60.61</td>
<td>4702</td>
<td>1.20 × 10⁶</td>
<td>80.30</td>
</tr>
<tr>
<td>ABySS</td>
<td>Mate-pair and paired-end</td>
<td>7</td>
<td>6430</td>
<td>390,143</td>
<td>12.58</td>
<td>18,904</td>
<td>998.4 × 10⁶</td>
<td>75.20</td>
</tr>
</tbody>
</table>

Comparison of genomes sequenced and assembled from Chromium with combined mate-pair and paired-end sequence data for *Austropuccinia psidii*. The genome of *A. psidii* assembled from the Chromium library in this study has been deposited at DDBJ/EMBL/GenBank under the accession QOUH00000000.
combined paired-end and mate-pair sequencing. The connected ribbons indicate the homologous relationship between scaffolds as determined using the NUCmer tool in MUMmer, which only included alignments larger than 500 kb.

Figure 1. Circos plot showing 18 largest scaffolds (>0.5 Mb, indicated as S1 to S18) from the Supernova assembly (Chromium sequencing) and their 220 homologous scaffolds from the ABySS assembly (combined paired-end and mate-pair sequencing). The connected ribbons indicate the homologous relationship between scaffolds as determined using the NUCmer tool in MUMmer, which only included alignments larger than 500 kb.

The genome size of A. psidii was first reported as 103–145 megabases [16], whereas it was estimated as >1.2 Gb in the present study. The difference between the two sequenced genomes may be that they are different strains, the pandemic strain sequenced by Tan et al. [16] and the South African strain [30] used here. However, intra-specific genomic diversity in P. striiformis f. sp. tritici and P. triticina was attributed to differences in sequencing platforms used, rather than differences between strains of rust [7–10,31,32]. We hypothesize the size difference of ~1 Gb between the strains is a reflection of the sequencing technology used, a Chromium 10X library sequenced on HiSeqX10 in the present study compared with MiSeq in Tan et al. [16].

Chromium 10X sequencing provided an effective method to sequence and assemble the genome of an obligate plant pathogen from a small quantity of starting material. The Chromium library preparation of A. psidii used less DNA at a cheaper cost to obtain a better assembly, in terms of number of scaffolds (Figure 1) and BUSCO genes recovered [33], than paired-end and mate-pair libraries (Table 1). Chromium sequencing has opened the door for de novo genome assembly, comparative genomics and phylogenomics of obligate plant pathogens and microorganisms, especially when DNA is a limiting factor.

**Author contributions**

AM, TD, VQ and BW designed the study. TD, VQ, WS and LS performed the experiments and analyzed the data. AM and TD wrote the manuscript. SN, MW and BW supervised the study and corrected the manuscript.

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**Competing & financial interests disclosure**

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Address correspondence to: Brenda D Wingfield, Department of Biochemistry, Genetics & Microbiology, Tree Protection Co-operative Programme (TPCP), Forestry & Agricultural Biotechnology Institute (FABI), University of Pretoria, South Africa; Brenda. Wingfield@fabi.up.ac.za

1Authors contributed equally

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