Potential and pitfalls of eukaryotic metagenome skimming:



A test case for lichens

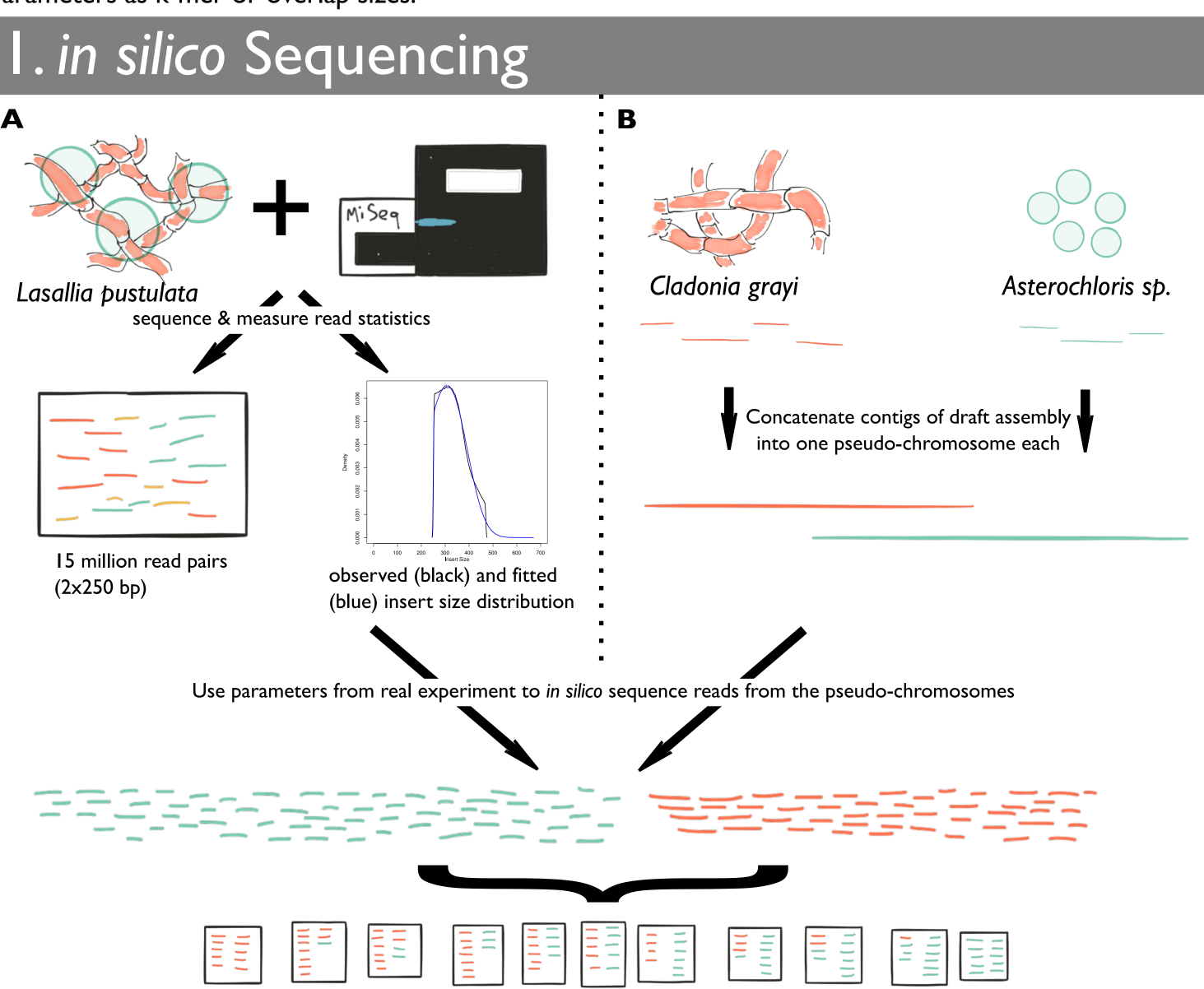
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Summary

Metagenomic sequencing with only a single library layout is used to quickly and cheaply assess the taxonomic and functional complexity of large and diverse microbial communities. We investigate to what extent such metagenome skimming approaches are applicable for the in-depth characterizations of genomes represented in obligate symbiotic communities of eukaryotes, e.g. lichens. It is still unclear how a eukaryotic species mixture, with larger and more repeat-rich genomes, influences different de novo assembly paradigms, such as de Brujin Graph based methods or Overlap Layout based assemblers and how to optimize assembly parameters as k-mer or overlap sizes.



Merge reads simulated from either reference genomes into L. pustulata twin data sets with varying coverage ratios for the two genomes.

Figure 1: Workflow for generating twin data sets, resembling a real sequencing data set with respect to insert size distribution, read number and read length.

DNA from a thallus of Lasallia pustulata was sequenced using Illumina MiSeq technology, yielding 15 million read pairs with a read length of 250 bp. To estimate the insert size distribution, we joined overlapping read pairs using FLASH [1] and fitted a censored Weibull distribution to the observed insert size distribution (Figure 1,A).

The scaffolds of the genomes of Cladonia grayi [2] and Asterochloris sp. chromosome, respectively (Figure I, B). Both were checked for repeat content & self-similarity using Repeatmasker [4] (Box I) and Gepard [5] (Fig 2.)

were each concatenated to create a contiguous pseudo-

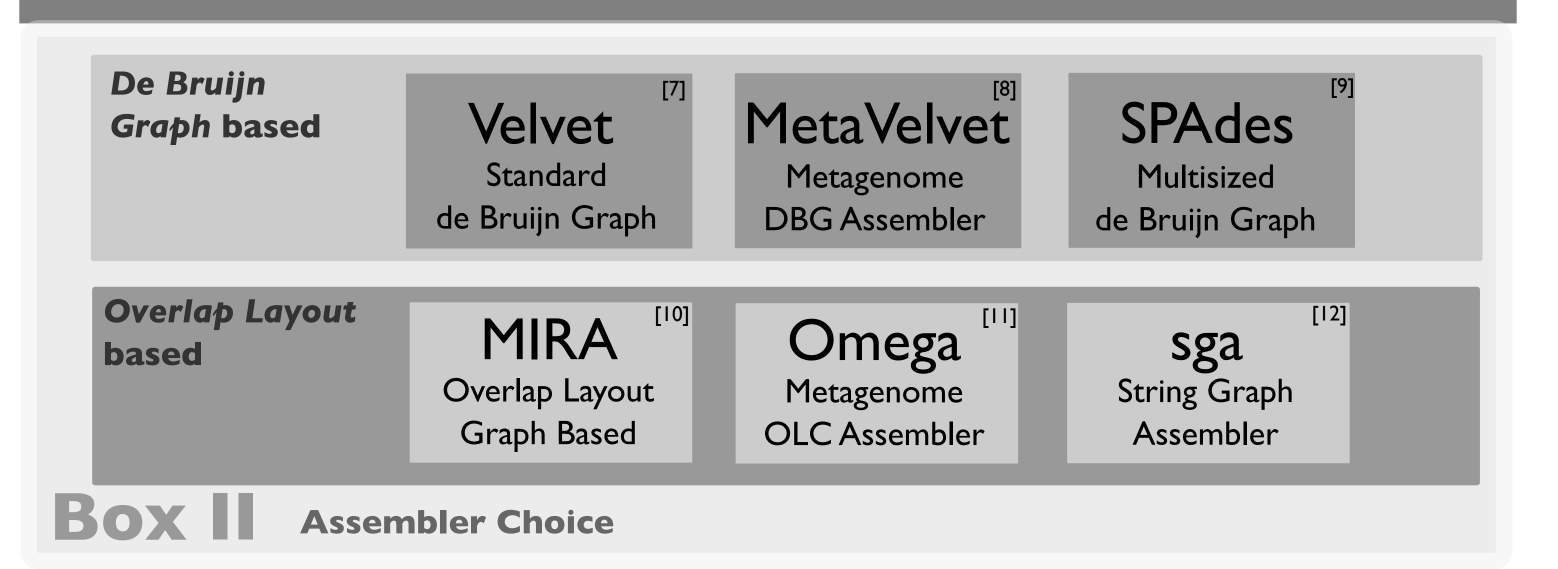
Asterochloris sp. Cladonia grayi **Number of Scaffolds** 1506 153 **Total Length** 38 Mbp 55 Mbp **GC** content 44 % 58 % % Repetitive 5 % 2.8 % Reference Genomes

Using the pseudo-chromosomes as templates, we simulated reads using ART [6], parameterized with the values estimated from the L. pustulata data. The reads were used to compile II twin data sets by mixing fungal and algal reads in varying ratios (Table 1).

Cladonia grayi	Asterochloris sp.	Table I: Absolute coverages for each organism per data set		
Chapt 38881221	Action 55759973	Coverage Ratio C. grayi : Asterochloris sp. 10:0 9:1 8:2 7:3 6:4 5:5 4:6 3:7 2:8 1:9 0:10	Coverage <i>C. grayi</i> 182x 157x 134x 112x 92x 74x 56x 40x 26x 13x 0x	Coverage Asterochloris sp. 0x 17x 33x 48x 61x 74x 86x 97x 107x 116x 125x

Figure 2: Dotplot of the pseudo-chromosomes of Cladonia grayi and Asterochloris sp.

2. Assembler Selection & Parameter Selection



For Omega, sga, Velvet & MetaVelvet we explored the parameter space (overlap size and k-mer size respectively) and used the maximization of the N50 size as the acceptance objective.

To address these questions, we performed an in silico study, simulating twin sets of genome skimming experiments of a lichen. We show that the quality of genome reconstructions from such data depends on assembler choice, but more importantly also on the parameter optimisation strategy and the ratio of the taxa in the metagenome. Optimising for assembly metrics such as N50 can in extreme cases lead to the exclusing of complete genomes. The outcome of a real-world metagenome skimming experiment of the lichen Lasallia pustulata not only shows a larger species diversity, but also hints to biased sequencing coverage for the algal genome.

3. Assembly Results

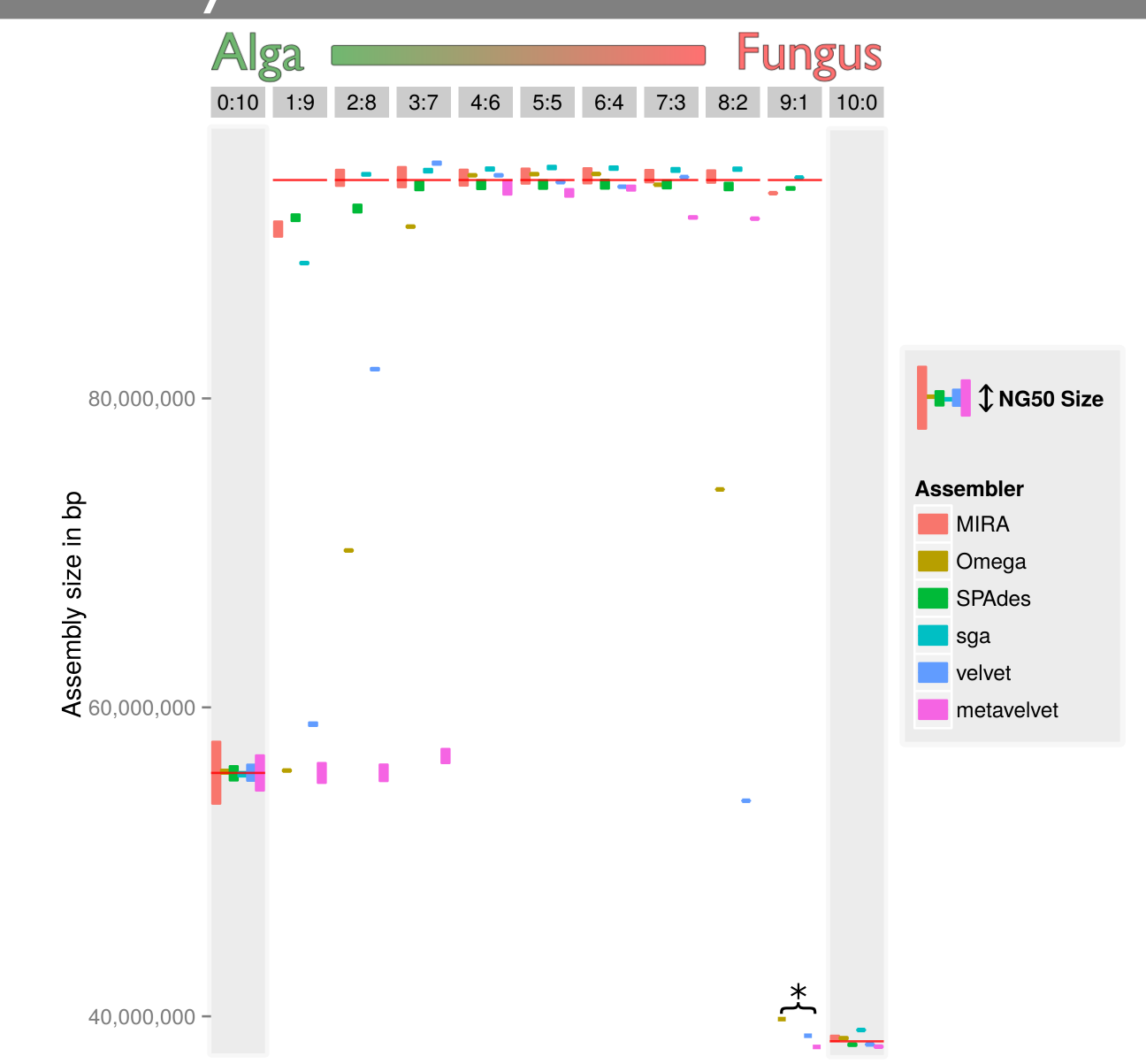


Figure 3: Assembly results for the 11 data sets and the diffferent assemblers. Bars are centered at total assembly length, red lines are reference lengths. Height of the bars shows the NG50 size. For the assemblies with the asterisk the total assembly length was less than 50% of the reference length. A default height was used in those instances.

Single Species Data Almost all assemblers reconstruct the two genomes over their full length

Asterochloris sp. (Figure 3, column 0:10 & 10:0), however with varying contiguity. For the alga many assemblers exceeded the NG50 size of the original draft assembly. For the fungus, it appears that the repetitive nature of the genome (c.f. Fig. 2) hinders the generation of longer contigs with the present WGS library layout (Figure 4).

Mixed Species Data Completeness of the genome reconstructions depends heavily on assembler choice and coverage ratios. MIRA and SPAdes perform best across all data sets. In contrast Velvet, Omega and MetaVelvet (the latter two being metagenome assemblers) fail to assemble large parts of the low coverage genome once coverage ratios become extreme (Fig. 3, 1:9 - 3:7, 9:1).

Parameter optimization for N50 impairs metagenome assembly Increasing the value of k reduces the frequency of all k-mers (Fig. 6), causing kmers from the low coverage genome to overlap with those introduced by the sequencing error. This prevents the formation of typically short contigs, thus increasing the N50 size by not assembling the low coverage genome.

L. pustulata Assembly Done with MIRA, the contigs were taxonomically assigned using MEGAN [13]. The algal assembly is much more fragmented than expected given the in silico study. (Box III). We think that this is a result of a biased library preparation, yielding a highly uneven read coverage for the algal genome.

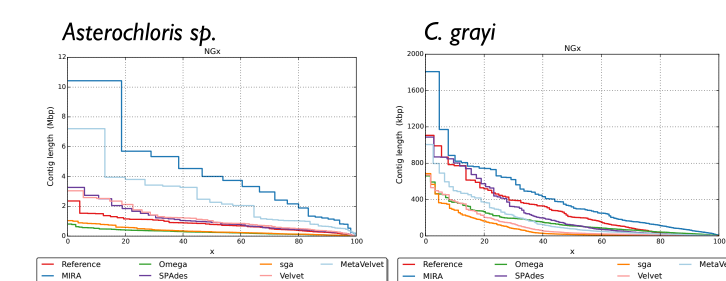


Figure 4: NGx distributions for Asterochloris sp. & C. grayi

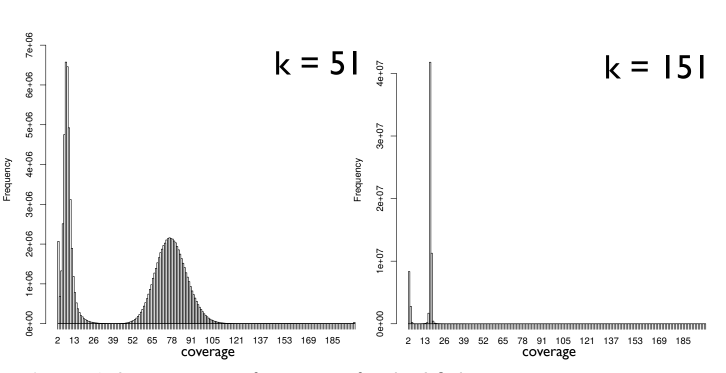
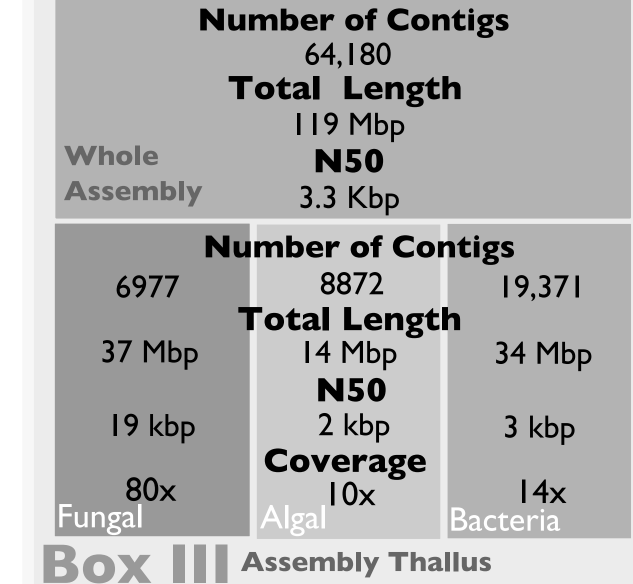


Figure 6: kmer-coverage frequencies for the 1:9 data set.



Summary

- Twin sets are valuable for guiding strategic decisions during planning of metagenome sequencing and assembly.
- For metagenomes optimising the assembly parameters using the N50 size can lead to the preclusion of entire genomes
- Assembler performance already varies substantially for single species data.
- Despite a 10x coverage, the algal genome assembly is highly fragmented.This may be due to an uneven algal genome

coverage, resulting from a biased library preparation.

Mixing data from different species inflates the assembler performance differences, with MIRA & SPAdes yielding the most contiguous sequences

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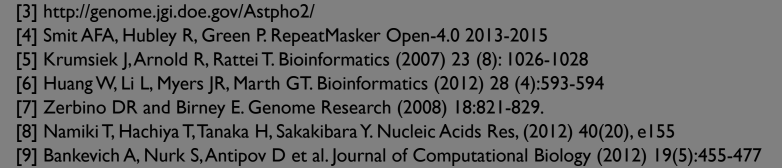
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