# MSc Thesis - Benchmark of *de novo* Short Read Assembly Strategies for Metagenomics

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# Introduction

Metagenomics, the sequencing of environmental DNA, has demonstrated to be a promising approach for the discovery and investigation of microbes that cannot be cultured in the laboratory [9] as well as for the study of both free-living microbial communities [1] and microbial communities inside other organisms [26, 11].

In a typical shotgun metagenomics experiment the DNA of a community is isolated and high throughput sequencing is performed on a random sample of the isolated DNA [22]. The reads can either be analyzed as such, by e.g. blast searches against reference databases to obtain a functional profile of the microbial community [30], or they can be assembled to form longer stretches of DNA stemming from the same or closely related organisms that can subsequently be analyzed with regards to phylogenetic affiliation and functional properties. The output of the assembly process often includes scaffolds, contigs and unassembled reads [19]. One of the problems with assembling is that chimeric contigs or scaffolds may be formed. Closely related sequences are more likely to form chimeras and since closely related strains often occur in the same environment this is a challenge. Also, it is difficult to determine whether the formation of a chimera is natural due to homologous recombination or an error in the assembly process [31]. Another problem with assembly is variations in gene content among closely related strains, since a gene inserted in a subpopulation will cause conflicting assembly results [10]. After assembling the reads, a process called binning is performed, where the resulting scaffolds and contigs are assigned to phylogenetically related groups. Finally, gene calling and functional annotations are performed on the scaffolds.

In our studies several strategies for *de novo* assembly of metagenomic data have been evaluated. In an *in silico* performed comparison between Illumina, Sanger and 454 on cost of sequencing and resulting coverage of microbial communities, Illumina short read libraries were shown to be the best for communities of medium

complexity [20]. Therefore we have chosen to assess the assembly strategies for Illumina paired short reads specifically. In previous studies mostly in silico metagenomic data sets have been used [25, 19]. In contrast the community of our study is an in vitro simulated metagenome consisting of 59 species with completed or nearly completed genomes so the quality of our assessment is not dependent on the realisticness of read simulators. An even and uneven distribution of the 59 species were created in vitro. The community has been sequenced with different type of library preparations to be able to test the difference in library preparation as well. The following assembly programs have been tested: Velvet [36], Meta-Velvet [24], Newbler [27], Minimus2 [29], Ray Meta [4] and Bambus2 [13]. The quality of the assemblies have been evaluated by mapping the constructed contigs or scaffolds to the collection of reference genomes, hereafter referred to as the reference metagenome. In addition two pipelines have been constructed, one to perform the assemblies and another to perform the validation given there is a reference metagenome available.

# Related Work

In a study by Mavromatis et al. [19] three genome assemblers were evaluated: Phrap [6], Arachne [3] and Jazz [2]. For the evaluation three artificial communities were constructed of low, medium and high complexity by selecting Sanger reads from 113 isolate genomes. The low complexity community had one dominating population with several low-abundance ones, the medium more than one dominating population and the complex community had no dominating population at all. Resulting contigs were evaluated on chimericity and length distribution. Compared to using the original reads for gene annotation, assembly was demonstrated to give up to 20% increase in accurate gene prediction. An accurate gene prediction was defined as a predicted gene that was also annotated in the original genome. Missing genes and inaccurately predicted genes decreased with slightly more than 20% after assembly. The approach of using artificial communities has subsequently been used in adapted versions by several other assembly evaluation papers [25, 20]. In those cases simulated reads were used that imitated the then-current most used sequencing methods. In the benchmark by Pignatelli and Moya [25] the reads of the artificial communities were changed from Sanger to 454 and Illumina. For the Illumina reads, SSAKE [33] and Velvet were used to perform the assembly. No difference in chimericity between using the simulated 454 reads or the Illumina reads was spotted. The main cause of chimericity was sequence similarity of the organisms, no relation with genome coverage was found. At the functional level metagenomic assembly turned out to be counterproductive or equally good as using the original 454 reads for annotation. For the Illumina reads the assembly did improve the functional annotation. Mende et al. [20] used a metagenome of 10, 100 and 400 species with simulated reads of Illumina, 454 and Sanger where the number of reads for each technology was based on sequencing cost. The sequencing cost was kept constant. All of the technologies provided similar coverage for 10 species. Illumina was superior for 100 species due to the higher coverage one can get for a similar price. Sanger performed best for 400 species because of longer read length. Sanger reads were assembled with Arachne; 454 reads with Celera [23] and Illumina reads with SOAPdenovo [16]. In contrast to the study of Pignatelli and Moya [25] a year earlier, the authors concluded that assembly contigs improved functional annotation of the metagenome for all technologies. Furthermore using Illumina paired end data to determine contig links and construct scaffolds, although introducing more chimerism, resulted in an even better functional annotation. Beyond using simulated reads or real reads of in silico communities there has not been a comparison of assembly algorithms using an in vitro community yet. In vitro communities have been used previously with success to assess DNA extraction techniques for sequencing a low complexity community of nine bacterial genera [34], an oral community of seven bacterial genomes [7], a human gut mock community of 10 bacterial genomes [35] and a human microbiome mock community of 21 bacterial genomes [12]. The advantage of using an in vitro community for assembly evaluation is that one does not have to rely on the correctness of sequencing simulators, the assessment can thus be as good as the similarity of the *in vitro* community to a real community.

# Methods and Materials

To determine the quality of metagenomic assembly a mock community of species with known genomes was constructed *in vitro* and sequenced with Illumina. The resulting reads have been assembled using a combination of Velvet, Meta-Velvet, Ray, Minimus2, Newbler and Bambus2 resulting in nineteen different assembly strategies (see Figure 3.1 and Table 3.1). The strategies stem from current literature and our own ideas.

# 3.1 Mock community

The mock community consists of 49 bacterial and 10 archaeal species with finished or nearly completed genomes (Table 3). The species have been chosen such that there are a number of closely related organisms and more distant ones. The number of species is about equal to the number of species one would find in the human gut of an individual [26]. The abundances of DNA from each species have been fixed in two types of configurations before sequencing. In the first configuration, the even configuration, all species have approximately equal genome copy numbers. In the second configuration, the uneven configuration, the phyla are mixed in proportions similar to log-normal distributions of phyla in soil [8]. The samples have been prepared with the Nextera 50ng sample preparation kit. The entire reference metagenome's size is about 195Mb. Mock community preparations and sequencing were performed by our collaborators, Christopher Quince at University of Glasgow and Linda D'Amore and Neil Hall at Liverpool's Centre for Genomics Research.

### 3.2 Quality trimming

Before assembling the reads one often starts with pre-processing them by quality trimming and/or removing PCR duplicates. Mende et al. [20] demonstrated that quality trimming could drastically improve the assembly. Before each assembly the same quality trimming procedure has been performed. For quality trimming the program sickle was used (see Table 1). Reads were trimmed from the 3' end if the average quality score was below 20 in a window of 10 bases. If the resulting read is shorter than 20 it is discarded. Only pairs are used in the subsequent assembly, not the single reads.

## 3.3 Assembly

In the assembly procedure reads are combined into contiguous sequences called contigs. Contigs can afterwards be joined using paired read information into longer scaffolds. In the scaffolding process contigs might be extended and repeats might be solved so scaffolding is not restricted to just the ordering of contigs.

There are a plethora of different assemblers available and by pre-processing reads and combining different assemblers an even larger amount of assembly strategies is possible. Velvet is one of the most used assembly programs and was therefore included in this assessment. Velvet's metagenomic counterpart, Meta-Velvet, is performed after executing Velvet so it is possible to determine how the metagenomic specific parameters improve the assembly. Another popular assembler that has recently received an update for metagenomics is Ray [4]. Ray is based on MPI and is runnable over multiple nodes distributing both memory and processor load, which makes it an ideal candidate for large metagenomic projects.

### 3.3.1 Contiging

Velvet, Ray and Meta-Velvet all use a de Bruijn graph to determine overlaps between reads. This involves cutting up the reads in sizes of a specified kmer size and let edges represent overlaps between kmers i.e. (k+1)mers. This way the graph, or the computational requirements, grow with the number of unique kmers in the library instead of the number of reads. For a more elaborate description of de Bruijn Graphs for sequence assembly see [21]. The resulting contigs are constructed by following paths in the graph. The paths that can be unambiguously followed are called unitigs. Ambiguous paths can be solved by using coverage information or paired-end information. Contigs thus consist of one or multiple unitigs. Choosing

the right kmer size is important. A shorter k gives more connectivity within the graph and hence requires lower sequencing coverage of the genomes, but at the same time the risk increases that a kmer occurs multiple times within a genome, or in multiple genomes (hence ambiguous paths will exist). A larger k can overcome this problem if it is larger than the multiply occurring region. But a larger k also requires higher sequence coverage.

#### How the assemblers differ

Velvet, Ray and Meta-Velvet differ in the way the graph is traversed. Velvet, meant for single genomes, looks for one coverage peak in the coverage distribution and tries to follow that, where the main idea is that the genome is approximately uniformly covered. Nodes in the graph below a certain coverage threshold are considered errors and ones with high coverage repeats. Meta-Velvet looks for multiple peaks in the coverage distribution. The contigs of each genome should have a distinct coverage peak due to the genome copy number of the corresponding genome being different from the other genomes in the metagenome. Meta-Velvet makes use of that property. Ray looks for 'seeds' in the graph and extends those seeds iteratively weighting choices by the number of reads supporting a certain path. The seeds are unitigs in the graph with a specific coverage. The metagenomic update to Ray changes the seed selection by looking at the coverage peak in the graph locally instead of globally.

#### 3.3.2 Merging

A way to get the advantage from both short and long kmers is by merging contigs generated in multiple assemblies with different kmer lengths. This is possible with Newbler, as done by Luo et al. [17], or with Minimus2, as done by for instance the Rnnotator pipeline [18]. Both Newbler and Minimus2 use an Overlap-Layout-Consensus method to merge contigs [29, 21].

### 3.3.3 Scaffolding

For the scaffolding procedure Bambus2 was chosen since it was one of the better scaffolders for single genomes in the GAGE assessment paper [28] and is suitable for metagenomes as well [13]. For a flow diagram of previously mentioned approaches see Figure 3.1. A total of twelve assembly strategies from the flow diagram have been tested. See Table 3.1 for an overview of the assembly strategies, Table 1 for versions of each program and Table 2 for the parameters of each strategy.

Assembly strategy name	Contiging	Merging	Scaffolding
velvetnoscaf	Velvet	-	-
velvetscaf	Velvet	-	Velvet
minimus2velvetnoscaf	Velvet	Minimus2	-
newblervelvetnoscaf	Velvet	Newbler	-
bambus2velvetnoscaf	Velvet	-	Bambus2
bambus2minimus2velvetnoscaf	Velvet	Minimus2	Bambus2
bambus2newblervelvetnoscaf	Velvet	Newbler	Bambus2
metavelvetnoscaf	Meta-Velvet	_	-
metavelvetscaf	Meta-Velvet	-	Meta-Velvet
minimus2metavelvetnoscaf	Meta-Velvet	Minimus2	-
newblermetavelvetnoscaf	Meta-Velvet	Newbler	-
bambus2metavelvetnoscaf	Meta-Velvet	-	Bambus2
bambus2minimus2metavelvetnoscaf	Meta-Velvet	Minimus2	Bambus2
bambus2newblermetavelvetnoscaf	Meta-Velvet	Newbler	Bambus2
raynoscaf	Ray	_	-
rayscaf	Ray	_	Ray
minimus2raynoscaf	Ray	Minimus2	-
newblerraynoscaf	Ray	Newbler	_

Table 3.1: Assembly strategies

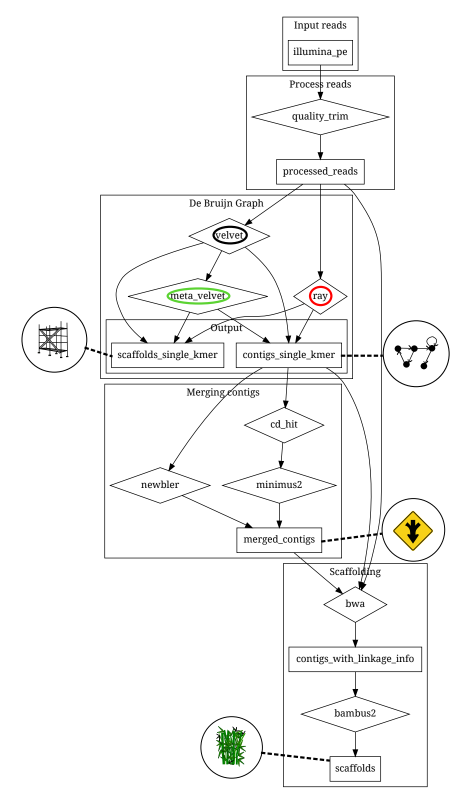


Figure 3.1: Assembly strategies using a combination of Velvet, Meta-Velvet, Ray, Minimus2, Newbler and Bambus2.

#### 3.4 Validation

The validation of a metagenomic assembly in case a reference metagenome is available often focuses on one or more of the following points:

- contig or scaffold length distribution
- contig/scaffold coverage of the reference metagenome
- chimericity of the contigs/scaffolds
- functional annotation accuracy
- phylogenetic classification accuracy

This study focusses on the first three points, since those are expected to improve the functional annotation and the phylogenetic classification. Mende et al. [20] showed that this was indeed the case for functional annotation.

#### 3.4.1 Length distribution

A variety of metrics are used in this validation. For the contig/scaffold length distribution of an assembly a common metric called the L50 value is used. L50 is a median value weighted by contig length. Half of all the bases in the assembly is in contigs equal to or larger than L50.

### 3.4.2 Coverage of the reference metagenome

For determining how well the assemblies matched the reference metagenome the assemblies were mapped against the reference metagenome using MUMmer 3.1 [14]. MUMmer finds maximal exact matches longer than l and clusters them if they are no more than g nucleotides apart. The alignments are afterwards extended for each cluster if the combined length of its matches is at least c. The alignments are extended in between the matches of the cluster and on the ends using a Smith-Waterman dynamic programming algorithm. The MUMmer package contains multiple scripts that make use of this approach. NUCmer (NUCleotide MUMmer) is a script included in the MUMmer package for DNA sequence alignment of a set of query contigs against a set of reference contigs. The command for NUCmer used was: nucmer --maxmatch -c65 -g90 -120. The maxmatch parameter makes sure all exact matches are used, whether they are unique or not, so contigs that consist only of a shared region or a repetitive element will be included in the alignments as well. Afterwards the script show-coords was used on the

resulting alignment file to extract information about each alignment such as its location in both the query and the reference, percent identity, percent similarity and percent of the reference and query covered.

To indicate how well the metagenome is covered by the assembled contigs we use a metric called genome contig coverage and metagenome contig coverage. For each genome in the reference metagenome, genome contig coverage is determined by dividing the number of bases that are covered by contigs mapping to it by the total number of bases in the genome. Metagenome contig coverage is the number of bases covered by all contigs mapping to it divided by the total number of bases in the metagenome. In both cases only the best alignment for each contig is used. The best alignment is the one with the highest purity score (for an explanation of the purity score see Section 3.4.3. In case a contig aligns equally well to multiple locations, only the first alignment is considered. This could for instance be a contig that represents a repeat or a shared region between genomes. By only counting one such alignment not resolving a repeat or shared region is penalized. Note that in general there's little difference between counting all such alignments or only one in bacterial and archaeal genomes.

#### 3.4.3 Chimericity

In previous research by Mavromatis et al. [19] chimericity of each contig was determined by the reads mapping to it. This could be done because reads were selected from finished genome projects, thus the origin of each read is known. At each taxonomic level from strain to domain the contigs were assigned to the phylogenetic group where the majority of reads stem from. The chimericity was then defined as the number of reads that belong to another phylogenetic group divided by the total number of reads mapping to the contig. Mende et al. [20] noted that certain reads might actually belong to multiple species since there are shared regions between species. Therefore they proposed a metric called contig score, which is determined by mapping each contig with blastn to the original genomes, taking the Highest Scoring Segment Pair (HSP) and multiplying the percent identity times the percent contig coverage. We use a similar metric, determined with MUMmer and referred to as purity instead to indicate its relation to chimericity. For each alignment the purity p is determined by multiplying percent conting coverage by percent identity divided by 10,000 to get the purity as a ratio between 0 and 1. The purity of a contig is the maximum purity of all its alignments. Purity of an entire assembly is computed as well. If n is the number of contigs,  $l_1, ..., l_n$  the length of each contig in nucleotides and  $p_1, ..., p_n$  the purity of each contig then the global purity P of an entire assembly is calculated as:

$$P = \frac{\sum_{i=1}^{n} l_i * p_i}{\sum_{i=1}^{n} l_i} \tag{3.1}$$

For both scaffolds and contigs, only nucleotides are counted when determining l. Thus unknown parts often represented as  $\mathbb{N}$  or – do not contribute to l.

### 3.5 Pipelines

The vast amount of assemblies and validations that had to be performed on a variety of libraries gave rise to a need for automation. This resulted in the development of two pipelines: MetAssemble and MASMVALI (Metagenomic ASseMbly VALIdation). The former for performing the assemblies, the latter for validating them given there is a reference metagenome available. The location of the repositories can be found in Table 3.2.

#### 3.5.1 MetAssemble

MetAssemble implements the strategies in Table 3.1 and Figure 3.1. The MetAssemble pipeline was written as a set of rules in GNU make. GNU make was chosen for a number of reasons:

- It integrates skipping computation of previously computed files. Recomputation is unfortunately only based on modification date, not on the actual file contents, but it is good enough for our purposes.
- GNU make is widely available on Unix and Unix-like systems.
- After the validation one would want to share only the commands to do a specific assembly i.e. not the entire pipeline. Doing a dry-run in GNU make gives you the exact commands that generated a certain file.

One of the biggest problems in metagenomic assembly are the computational demands [37]. The size of the reads and the number of reads as well as the complexity and the structure of the sequenced community affect the computational demands. Therefore it is difficult to determine the resource usage in advance. In our benchmark several strategies consist of parts with different computational requirements. Ray for instance runs over multiple nodes whereas merging with Newbler or Minimus2 requires one node with a lot of memory. Claiming all these computational resources for the entire run of the pipeline would be a waste. To circumvent this issue an approach where each step in the pipeline is scheduled as a separate job

using a job scheduler has been chosen. A library was developed for GNU make to schedule rules as jobs using either sbatch or qsub. The jobs are scheduled with dependencies on other jobs and the resource usage set for the rule.

#### 3.5.2 MASMVALI

After the assembly has been completed, the assembly can be validated by alignment against the reference metagenome. MASMVALI (Metagenomic ASseMbly VALIdation) is a pipeline written in Python that calculates the statistics described in Section 3.4. As minimum input it takes an assembly, a reference metagenome and alignments found by MUMmer of the assembly against the reference metagenome. MASMVALI as a pipeline can generate a HTML report that shows a collection of the plots that are used in this thesis. It is also possible to use MASMVALI as a package to do more analyses on generated assemblies using the available statistics. See for an example the ipython notebook in Table 3.2.

Name	URL
MetAssemble	https://github.com/inodb/metassemble
MASMVALI	https://github.com/inodb/masmvali
MASMVALI ipyhton notebook example	http://nbviewer.ipython.org/urls/raw.
	github.com/inodb/msc-thesis/master/
	notebooks/chris-mock-masmvali-length.
	ipynb
GNU Make job scheduler	https://github.com/inodb/
	gnu-make-job-scheduler

Table 3.2: URLs for MASMVALI and MetAssemble

# Results

## 4.1 Mock community libraries

All assemblies in Table 3.1 have been performed on the following libraries:



#### 50ng even

Community prepared with 50 ng Nextera library prepartion kit and equal genome copy numbers. The library has been sequenced with Illumina and contains  $\sim$ 10M paired reads, each single read has a length of 100.



#### 50ng uneven

Community prepared with 50 ng Nextera library preparation kit and phyla mixed similar to log-normal distributions of phyla in soil. The library has been sequenced with Illumina and contains  $\sim 10$ M paired reads, each single read has a length of 100.

The original reads of each library were mapped against the reference genome using BWA [15] and default values. Afterwards duplicates were removed with MarkDuplicates from Picard 1.77 (see also Table 1). In Figure 4.1 for the even library, the even spread of the coverage over all the genomes in the metagenome can be observed. The median coverage over all bases is 10. The uneven library covers a couple of genomes very well 4.2. The median coverage over all bases is 3.

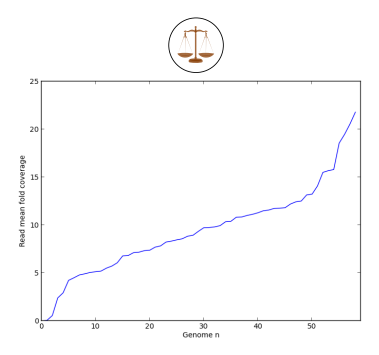


Figure 4.1: Read mean coverage of  $50 \, \mathrm{ng}$  even per genome. Median coverage over all bases is 10.

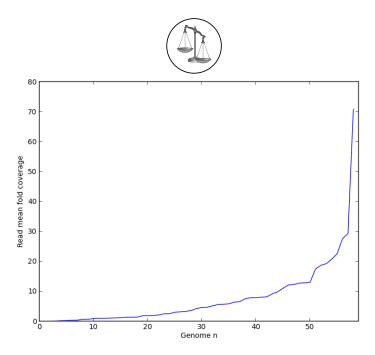


Figure 4.2: Read mean coverage of  $50 \, \mathrm{ng}$  uneven per genome. Median coverage over all bases is 3.

### 4.2 Length distribution

In Figure 4.3 all assembly strategies are compared over length distributions. The x-axis indicates the threshold of contig length used in calculating the sum of bases on the y-axis. The x-axis is logarithmically scaled and reversed in order to give a better indication of what the longer contigs contribute to the sum. The dots indicate L50 values on the x-axis. For each assembly strategy the assembly with a kmer that resulted in the highest L50 has been chosen. As an example, for the contiging strategies on the reads from the even library (top left in Figure 4.3), the assembly with the highest L50 value was for velvetnoscaf and metavelvetnoscaf k=25 and for Ray k=21. In this case Meta-Velvet scores the highest on L50, but not only by combining contigs that are not combined by Velvet. It actually also removes a lot of short contigs from the original Velvet assembly, resulting in a lower total sum of bases. Depending on what size of contigs are required for the post-processing the winner differs. Longer than 100, Velvet and to a lesser extent Ray give more total bases, but longer than 1000 Meta-Velvet would seem a better candidate. When comparing length distributions between different assemblers, only looking at L50 is thus not enough.

In the first row that shows the contiging strategies, the uneven sample increases the gap in sum of bases between Meta-Velvet and the other assemblers even more. The merging strategies increase lengths quite significantly for both libraries. The difference between Newbler and Minimus2 being rather small. Minimus2 results in slightly larger contigs, only for Meta-Velvet the difference is substantial. The internal scaffolding options of the assemblers improves lengths, but the sum of bases decreases substantially. For the external scaffolder, Bambus2, the sum decreases even more.

Notice also how the internal scaffolder from Meta-Velvet and Velvet result in the same assembly for the even library. Meta-Velvet is based on following multiple peaks in the coverage distribution where each peak would theoretically represent an organism. Velvet can follow one peak with the <code>-exp\_cov</code> auto parameter. For the contiging approach however this parameter is not set (See Table 2), but for Meta-Velvet it is, which results in different assemblies for the even library. To enable internal scaffolding in Velvet one has to set the <code>-exp\_cov</code> auto parameter, making Velvet and Meta-Velvet follow the same peak in coverage for the even library. For the uneven library the results differ, since Meta-Velvet finds multiple peaks and Velvet only uses one.

# 4.3 Metagenome coverage

In Figure 4.4 the sum of bases axes from Figure 4.3 have been replaced by the metagenome coverage. The actual non-overlapping contig coverage of the reference metagenome after adding a particular contig (see Section 3.4.2). An addition of a contig that doesn't increase the metagenome coverage by its own length could be due to:

- 1. the contig not aligning anywhere
- 2. the contig aligns only partially because it is chimeric or has errors
- 3. the mapping location is already partially or fully covered by another contig.

Overall the plots look very similar to the original length distributions. The increase in coverage is less smooth than the increase in length indicating that one or more of the three previously mentioned points occur, but not in an extreme fashion. In case no reference metagenome is available looking at the length distributions in this manner is thus a good indication of the metagenome coverage. Only for the merging strategies significant differences can be spotted. The metagenome coverage for Minimus2 and Newbler follow more similar curves than is the case in Figure 4.3. Merging with Newbler results in a higher coverage of the metagenome for both Velvet and Ray than merging with Minimus2.

We found that nearly all contigs for each assembly are aligning. None of the assemblers thus creates bogus contigs. Whether point 2 or 3 is the case is shown in Section 4.4 for the different assembly strategies.

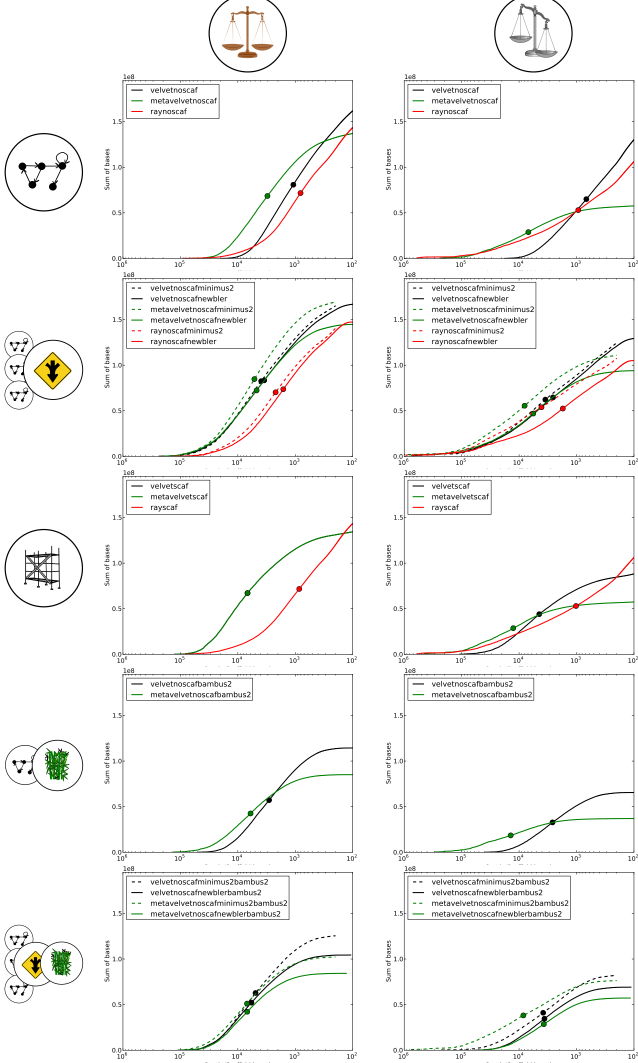


Figure 4.3: A comparison of assembly strategies on length distributions. On the y-axis the sum of bases and on the x-axis the contig length.

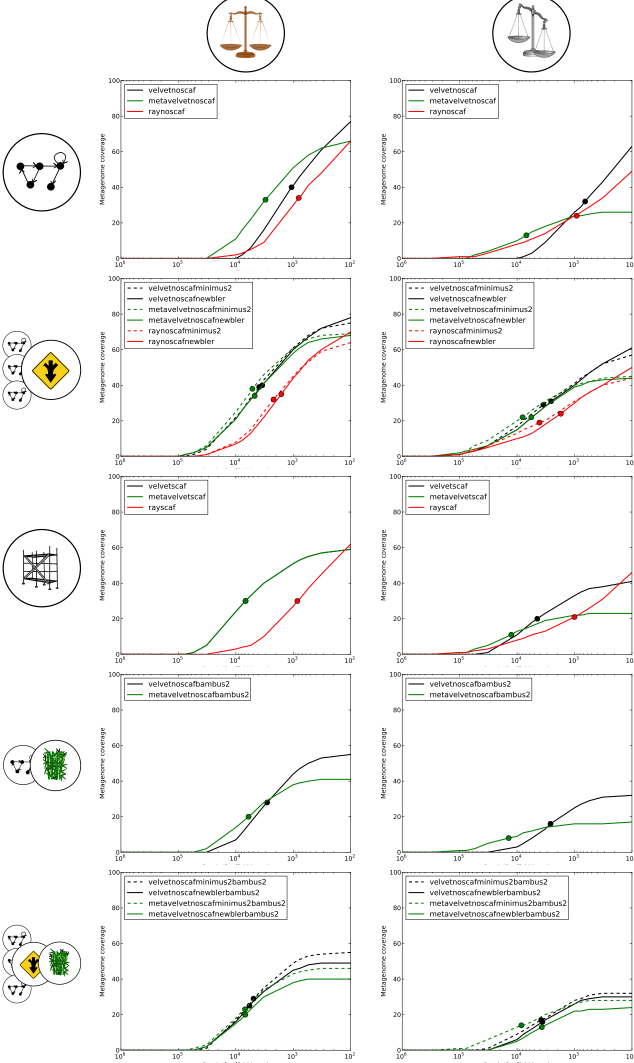


Figure 4.4: A comparison of assembly strategies on metagenome coverage. On the y-axis the metagenome coverage and on the x-axis the contig length.

# 4.4 Chimericity

To determine the chimericty of the contigs in an assembly the global purity metric described in Section 3.4 is used. In Figure 4.5 all assemblies are compared on purity where global purity is shown as a function of L50. In this plot every dot represents a single assembly. The size of the dot represents the kmer size from 19 to 75. Although a high L50 is not a perfect indication of a good length distribution, it does allow one to see how increases in length are related to the chimericity of the contigs in the assembly. The L50 values are all based on a cut-off length of 100.

The contiging strategies display a minor decrease in purity for an increase in L50 with shorter k. There is always some point visible for the Meta-Velvet assemblies where the L50 goes down again with a too short k. At that point the de Bruijn graph becomes saturated with too many connections. This point occurs between k=20 and k=30 for all three assemblers although for Velvet and Ray it is not clearly visible from the graph. The even community results in a more continuous curve with varying k than the uneven library for Meta-Velvet. For the uneven library some clusters are visible. No cause for these clusters has been researched, but a possible explanation is that it has to do with the way Meta-Velvet selects the coverage peaks, since that is a binary choice. At a certain k the coverage threshold is reached and a peak is detected, which could give rise to the clusters in the plot.

The merging strategies perform well, the L50 is increased and the purity is kept high. The Newbler assemblies are slightly purer than Minimus2. The larger contig lengths of Minimus2 compared to the metagenome coverage in Figure 4.4 are thus at least partly due to chimeric contigs.

The internal scaffolding introduces a lot of impure contigs. Notice also how Ray's internal scaffolder decreases in purity with decreasing k, whereas Velvet and Meta-Velvet have increased purity with decreasing k. At the lower values of k the purity is very similar. Bambus2 results in very pure assemblies. As demonstrated in Section 4.2 the larger L50 values compared to the other assembly stategies is in large part a result of losing many short contigs.

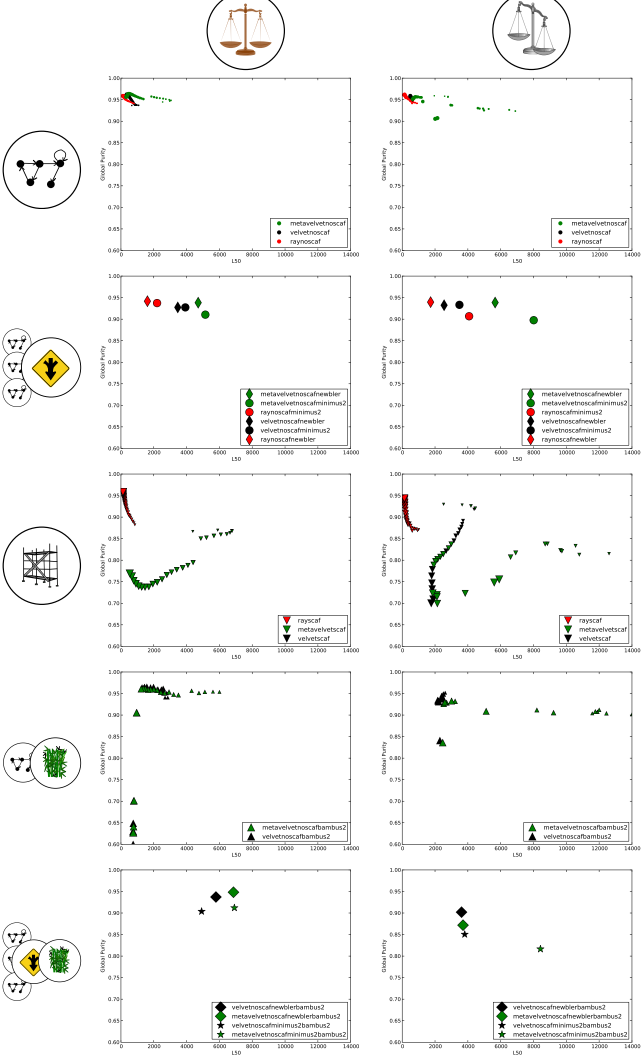


Figure 4.5: A comparison of assembly strategies on purity versus L50.

# Discussion

It is difficult to name one true winner. Depending on the post-processing and the computational resources available the most suitable assembly strategy differs. In case one is only interested in post-processing contigs larger than 1K, Meta-Velvet performs well. Larger than 100: Velvet is a suitable candidates. For large libraries that would require too much memory to run Velvet, Ray can be used. In general the merging strategies preform well on all three validated properties: length, metagenome coverage and purity. There are minor differences between using either Newbler or Minimus2 for merging. Minimus2 results in more chimerism in some cases. Even though both mergers do not make use of metagenomic heuristics they perform well. A merger that does implement the use of all the information available, such as the read coverage of the contigs and pair linkage between contigs seems like a worthwhile project.

The current results are only based on two libraries so future work of doing the same analysis on another library and the same or a different mock community would give more insight whether the conclusions hold in other circumstances. Ideally one could get an insight for each type of community what assembly strategy is preferred. Making the analysis in this paper available as a web service allowing users to run it on their own assembly, library, mock community combination and comparing them to the existing ones, could lead to a form of crowd sourced benchmarking.

The post-processing steps such as taxonomic classification and gene annotation have not been part of this study. It would be significant to know what size of contigs can be used in these post-processing steps and what purity level is required. Is there a balance between length increase and purity loss that is beneficial for a certain type of post-processing?

Our efforts have focussed on validating assemblies when a reference is available. Apart from the length distribution plot that is a decent indication of metagenome coverage, no reference-less validation approaches have been suggested. Comparing our results versus a reference-less validation technique, such as Feature Response Curve [32] or ALE [5] would be an important addition, especially since metagenomics rose out of the problem that the vast majority of microbes cannot be cultured in the lab.

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Program	Version	URL
sickle	commit 2013-4-3 e435598aa6bbb74b3254d2884a9dea1b0639d464	https://github.com/najoshi/sickle
Velvet	1.2.01	https://github.com/
		dzerbino/velvet/commit/
		5d5b636c52e94fde745cb8bf753ffa134126e04d
Meta-Velvet	1.2.01	
Minimus2	Amos commit 2012-3-6	http://amos.git.sourceforge.net/git/
		<pre>gitweb.cgi?p=amos/amos;a=commit;h=</pre>
		7c70d6c36dba244ae0b266fc52a5206ab202a36c
Newbler	RunAssembly $2.6 (20110517_{-1502})$	
MUMmer	3.23	http://sourceforge.net/projects/mummer/
		files/mummer/3.23/MUMmer3.23.tar.gz/
		download
CD-HIT	4.5.7	
Picard	1.77	
bwa	0.6.1-r104	
Bambus2	Amos commit 2012-3-6	http://amos.git.sourceforge.net/git/
		<pre>gitweb.cgi?p=amos/amos;a=commit;h=</pre>
		7c70d6c36dba244ae0b266fc52a5206ab202a36c
samtoafg	Amos commit 2012-3-6	http://amos.git.sourceforge.net/git/
		<pre>gitweb.cgi?p=amos/amos;a=commit;h=</pre>
		7c70d6c36dba244ae0b266fc52a5206ab202a36c
Ray	2.1.0	2.1.0 http://denovoassembler.sourceforge.net/

Table 1: Program versions used for the analysis

Assembly strategy name	Program Parameters
velvetnoscaf	Run velveth \$dir 31,84,2 -fastq -shortPaired \$pairs.fastq and run
	velvetg -scaffolding no on the resulting directories.
velvetscaf	Run velveth \$dir 31,84,2 -fastq -shortPaired \$pairs.fastq and run
	velvetg -scaffolding yes -exp_cov auto on the resulting directories.
metavelvetnoscaf	Run velveth \$dir 31,84,2 -fastq -shortPaired \$pairs.fastq and run
	velvetg -scaffolding no -exp_cov auto -read_trkg yes && -scaffolding no on
	the resulting directories.
metavelvetscaf	Run velveth \$dir 31,84,2 -fastq -shortPaired \$pairs.fastq and run
	velvetg -scaffolding no -exp_cov auto -read_trkg yes && -scaffolding yes on
	the resulting directories.
raynoscaf	Run Ray -k \$kmersize -i \$pairs.fastq -o output_dir.
minimus2*	Concatenate contigs larger than 200 and run
	cd-hit-est -c 0.99 -i \$concat.fasta -o \$derep.fasta to remove similar sequences.
	Run toAmos -s \$derep.fasta -o \$derep.afg followed by minimus2 \$derep. From http:
	//ged.msu.edu/angus/metag-assembly-2011/velvet-multik.html
newbler*	Cut all contigs up in chunks of 1999 bases with an overlap of 1900 bases and run
	runAssembly -o $tin cut-up-contigs.fasta. From [17].$
bambus2*	Map paired reads with bwa using default parameters to contigs and remove duplicate reads with
	MarkDuplicates. Run samtoafg to convert to afg, import to Amos bnk with bank-transact
	and finally run goBambus2.

Table 2: Program parameters per assembly strategy.

Table 3: Mock community

$N_{\rm o}$	Genome Name	Genome Size (bp)	Domain	Phylum	Class
	Acidobacterium capsulatum ATCC 51196	4127356	Bacteria	Acidobacteria	Acidobacteriae
2	Akkermansia muciniphila ATCC BAA-835	2664102	Bacteria	Verrucomicrobia	Verrucomicrobiae
က	Anaerocellum thermophilum Z-1320, DSM 6725	2919718	Bacteria	Firmicutes	Clostridia
4	Bacteroides thetaiotaomicron VPI-5482	6293399	Bacteria	Bacteroidetes	Bacteroidia
ಬ	Bacteroides vulgatus ATCC 8482	5163189	Bacteria	Bacteroidetes	Bacteroidia
9	Bordetella bronchiseptica RB50	5339179	Bacteria	Proteobacteria	Betaproteobacteria
7	Burkholderia xenovorans LB400	973113	Bacteria	Proteobacteria	Betaproteobacteria
∞	Caldicellulosiruptor saccharolyticus DSM 8903	2970275	Bacteria	Firmicutes	Clostridia
6	Chlorobaculum tepidum TLS	2154946	Bacteria	Chlorobi	Chlorobia
10	Chlorobium limicola DSM 245	2763181	Bacteria	Chlorobi	Chlorobia
11	Chlorobium phaeobacteroides DSM 266	3133902	Bacteria	Chlorobi	Chlorobia
12	Chlorobium phaeovibrioides DSM 265	1966858	Bacteria	Chlorobi	Chlorobia
13	Chloroflexus aurantiacus J-10-fl	5258541	Bacteria	Chloroflexi	Chloroflexi
14	Clostridium thermocellum ATCC 27405	3843301	Bacteria	Firmicutes	Clostridia
15	Deinococcus radiodurans R1	3284156	Bacteria	Thermi	Deinococci
16	Desulfovibrio desulfuricans desulfuricans ATCC 27774	2873437	Bacteria	Proteobacteria	Deltaproteobacteria
17	Desulfovibrio piger ATCC 29098	2826240	Bacteria	Proteobacteria	Deltaproteobacteria
18	Dictyoglomus turgidum DSM 6724	1855560	Bacteria	Dictyoglomi	Dictyoglomia
19	Enterococcus faecalis V583	3359974	Bacteria	Firmicutes	Bacilli
20	Fusobacterium nucleatum nucleatum ATCC 25586	2174500	Bacteria	Fusobacteria	Fusobacteria
21	Gemmatimonas aurantiaca T-27T	4636964	Bacteria	Gemmatimonadetes	Gemmatimonadetes
22	Herpetosiphon aurantiacus ATCC 23779	6785430	Bacteria	Chloroflexi	Chloroflexi
23	Hydrogenobaculum sp. Y04AAS1	1559514	Bacteria	Aquificae	Aquificae
24	Leptothrix cholodnii SP-6	4909403	Bacteria	Proteobacteria	Betaproteobacteria
25	Nitrosomonas europaea ATCC 19718	2812094	Bacteria	Proteobacteria	Betaproteobacteria
26	Nostoc sp. PCC 7120	7211789	Bacteria	Cyanobacteria	unclassified
27	Pelodictyon phaeoclathratiforme BU-1	3018238	Bacteria	Chlorobi	Chlorobia
28	Persephonella marina EX-H1	2467104	Bacteria	Aquificae	Aquificae
29	Porphyromonas gingivalis ATCC 33277	2354886	Bacteria	Bacteroidetes	Bacteroidia
30	Rhodopirellula baltica SH 1	7145576	Bacteria	Planctomycetes	Planctomycetacia
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$\sim$	Genome Name	Genome Size (bp)	Domain	Phylum	Class
31	Rhodospirillum rubrum ATCC 11170	4406557	Bacteria	Proteobacteria	Alphaproteobacteria
32	Ruegeria pomeroyi DSS-3	4601053	Bacteria	Proteobacteria	Alphaproteobacteria
33	Salinispora arenicola CNS-205	5786361	Bacteria	Actinobacteria	Actinobacteria
34	Salinispora tropica CNB-440	5183331	Bacteria	Actinobacteria	Actinobacteria
35	Shewanella baltica OS185	5312910	Bacteria	Proteobacteria	Gammaproteobacteria
36	Shewanella baltica OS223	5358884	Bacteria	Proteobacteria	Gammaproteobacteria
37	Sulfitobacter sp. EE-36	3547243	Bacteria	Proteobacteria	Alphaproteobacteria
38	Sulfitobacter sp. NAS-14.1	4002069	Bacteria	Proteobacteria	Alphaproteobacteria
39	Sulfurihydrogenibium sp. YO3AOP1	1838442	Bacteria	Aquificae	Aquificae
40	Sulfurihydrogenibium yellowstonense SS-5	1534471	Bacteria	Aquificae	Aquificae
41	Thermoanaerobacter pseudethanolicus ATCC 33223	2362816	Bacteria	Firmicutes	Clostridia
42	Thermotoga neapolitana DSM 4359	1884562	Bacteria	Thermotogae	Thermotogae
43	Thermotoga petrophila RKU-1	1824357	Bacteria	Thermotogae	Thermotogae
44	Thermotoga sp. RQ2	1877693	Bacteria	Thermotogae	Thermotogae
45	Thermus thermophilus HB8	2116056	Bacteria	Thermi	Thermi
46	Treponema denticola ATCC 35405	2843201	Bacteria	Spirochaetes	Spirochaetes
47	Wolinella succinogenes DSM 1740	2110355	Bacteria	Proteobacteria	Epsilonproteobacteria
48	Zymomonas mobilis mobilis ZM4	2223497	Bacteria	Proteobacteria	Alphaproteobacteria
49	Archaeoglobus fulgidus DSM 4304	2178400	Archaea	Euryarchaeota	Archaeoglobi
50	Ignicoccus hospitalis KIN4/I	1297538	Archaea	Crenarchaeota	Thermoprotei
51	Methanocaldococcus jannaschii DSM 2661	1664970	$\operatorname{Archaea}$	Euryarchaeota	Methanococci
52	Methanococcus maripaludis C5	1 780 761	Archaea	Euryarchaeota	Methanococci
53	Methanococcus maripaludis S2	1661137	Archaea	Euryarchaeota	Methanococci
54	Nanoarchaeum equitans Kin4-M	490 885	Archaea	Nanoarchaeota	Nanoarchaea
55	Pyrobaculum aerophilum IM2	$2\ 222\ 430$	Archaea	Crenarchaeota	Thermoprotei
56	Pyrobaculum calidifontis JCM 11548	$2\ 009\ 313$	Archaea	Crenarchaeota	Thermoprotei
57	Pyrococcus horikoshii OT3	1738505	Archaea	Euryarchaeota	Thermococci
58	Sulfolobus tokodaii 7(S311)	2694756	Archaea	Crenarchaeota	Thermoprotei
59	Treponema vincentii I	2512734	Bacteria	Spirochaetes	Spirochaetia