

PMSGGA: A FAST DNA FRAGMENT ASSEMBLER

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Abstract: The DNA fragment assembly is an essential step in DNA sequencing projects. Since DNA sequencers output fragments, the original genome must be reconstructed from these small reads. In this paper, a new fragment assembly algorithm, Pattern Matching based String Graph Assembler (PMSGGA), is presented. The algorithm uses multipattern matching to detect overlaps and a minimum cost flow algorithm to detect repeats. Special care was taken to reduce the algorithm's run time without compromising the quality of the assembly. PMSGGA was compared with well-known fragment assemblers. The algorithm is faster than other assemblers. PMSGGA produced high quality assemblies with prokaryotic data sets. The results for eukaryotic data are comparable with other assemblers.

1 Introduction

DNA fragment assembly is a technique that attempts to reconstruct the original DNA sequence from a large number of fragments, each several hundred base-pairs long. The DNA fragment assembly is needed because current technology, such as gel electrophoresis, cannot directly and accurately sequence DNA molecules longer than 1000 bases. However, most genomes are much longer. For example, a human DNA is about $3.2 \cdot 10^9$ nucleotides in length and cannot be read at once. The shotgun sequencing technique was developed to deal with this limitation. First, the DNA molecule is amplified, that is, many copies of the molecule are created. The molecules are then cut at random sites to obtain fragments that are short enough to be sequenced directly. The overlapping fragments are then assembled into a target DNA sequence. Several algorithms have emerged to tackle the fragment assembly problem. Some of the most well-known assemblers are PHRAP (Green, 1999), CAP3 (Huang & Madan, 1999) and EULER (Pevzner & al., 2001).

In this paper, a new fragment assembly algorithm, Pattern Matching based String Graph Assembler (PMSGGA), is presented. The algorithm is based

on the overlap-layout-consensus paradigm. Pairwise overlaps are detected by finding common probes among fragments. Next, the layout is built using a minimum cost network flow algorithm. Finally, the consensus sequence is constructed using q -mers and position based hashing.

The paper is organized as follows. The details of the different phases of PMSGGA are described in Section 2. In Section 3, the experimental results and comparisons with other algorithms are presented. Finally, Section 4 concludes the article with suggestions for future directions and improvements.

2 The Algorithm

PMSGGA is an assembler based on the overlap-layout-consensus paradigm. In the overlap phase, the task is to find all possible pairwise overlaps between fragments and their reverse complements. In the layout phase, the order in which the fragments should be placed in the final assembly is determined. Finally, in the consensus phase, the aim is to construct the contigs by finding a consensus of the overlapping fragments. Next, the three phases of PMSGGA are described.

2.1 The Overlap Phase

The overlap phase is based on the *AMASS algorithm* (Kim, 1997). Kim used a multipattern matching algorithm to efficiently find possible pairwise overlaps between fragments.

PMSGGA begins by randomly selecting probes of constant lengths from fragments and their reverse complements. Then, a faster multipattern matching algorithm, called the *BG algorithm* (Salmela & al., 2006), is used to find the occurrences of these probes in the fragments.

The BG algorithm is based on the BNDM algorithm (Navarro & Raffinot, 2000) for a single pattern. The idea is to construct a generalized pattern that represents a group of patterns. For example, the group of patterns, *acgt*, *aacc*, and *gttt* can be represented by the generalized pattern: $[a,g][a,c,t][c,g,t][c,t]$. In the overlap phase, PMSGGA uses the BG algorithm to find the generalized pattern representing overlapping k -mers instead of single characters of the patterns. If $k = 2$ in the example above, the corresponding generalized pattern is given by $[ac,aa,gt][cg,ac,tt][gt,cc,tt]$. Each occurrence of the generalized pattern is a candidate for a real match. BNDM works as a filter and the candidate matches are checked by an exact method. In practice, the BG algorithm is very efficient (Salmela & al., 2006).

To deal with small repeats, the probes with too many occurrences are filtered out. However, if there are enough probes with about equal number of occurrences, they are left in. This is done to preserve large repeats, but to filter out small repeats that might cause false overlaps between fragments.

The probe length (m) is obtained by solving $m = \ln F / \ln(1 - \epsilon)$, where F is the predefined average probability of probe occurrence and ϵ is the average error rate. In this work, the probe length varies between 10 and 30 base pairs, and $F = 0.4$ is used.

The overlap between a pair of fragments i and j is computed as follows. The common probes of these fragments are found using the BG algorithm. Let probes a and b occur in i and j at positions $(pos_{a,i}, pos_{a,j})$ and $(pos_{b,i}, pos_{b,j})$. Now, a and b are said to be a *consistent pair*, if

$$|(pos_{b,i} - pos_{a,i}) - (pos_{b,j} - pos_{a,j})| < \sigma$$

for a small threshold value σ . This threshold is needed to deal with insertion and deletion errors in the fragments. A set of common probe occurrences is a *consistent set*, if each consecutive pair of probe occurrences in the set is a consistent pair.

For each consistent set S_i that can be constructed from the probe hits in the overlapping area, a score is calculated as $Score(S_i) = N_d W + N_m$, where N_d is

the number of disjoint probe sets within the consistent set, W is the distance from the first probe in the set to the last one and N_m is the total number of probes in the set. The consistent set with the largest score is chosen to represent the overlap, and the length of the overlap is calculated by using the selected set.

The quality of the detected overlaps is checked as follows. Let a k -mer represent a sequence of k contiguous base pairs. Given a fragment a , occurrences of all possible k -mers in overlapping areas are determined. A vector V^a of length 4^k is constructed, where its elements represent the number of occurrences of a k -mer. Similarly, a vector V^b for fragment b is constructed.

The average error probabilities for the overlapping area of fragments a and b , denoted by p_a and p_b , are also calculated. To compute these probabilities, the PHRED quality scores (Ewing & Green, 1998) for each fragment are used.

The number of k -mers that occur only in one fragment is calculated as follows

$$N_{miss} = \sum_{i=1}^{4^k} |(V^a - V^b)_i| \quad (1)$$

The total number of k -mers, N_{tot} , is given by $N_{tot} = 2(L - k)$, where L is the length of the overlap being considered. The number of common k -mers is given by $N_{hit} = N_{tot} - N_{miss}$.

The probability of a sequencing error in a k -mer is given by $p_m = 1 - ((1 - p_a)(1 - p_b))^k$. The probability of observing N_{miss} with given error probability, is given by

$$P(X \geq N_{miss}) = \sum_{i=N_{miss}}^{N_{tot}} \binom{N_{tot}}{i} p_m^i (1 - p_m)^{N_{tot}-i} \quad (2)$$

A threshold ξ is set to discard all overlaps, where

$$P(X \geq N_{miss}) \leq \xi \quad (3)$$

Note that $P(X \geq N_{miss})$ is the incomplete beta function, $I_{p_m}(N_{miss}, N_{hit} + 1)$, which can be efficiently approximated.

To validate overlaps, PMSGGA counts the distinct k -mers, instead of counting errors, as was done by Keccioglu and Myers (1995). Note that by increasing the value of k , a greater emphasis is placed on the correct order of the matching substrings within the overlap at the expense of a smaller number of common k -mers. Fragments that are entirely contained in other fragments are removed and are used in the final consensus phase.

2.2 The Layout Phase

The layout phase of PMSGGA is based on string graphs (Myers, 2005). The idea is to construct a bidirected

overlap graph describing the overlaps and orientations between fragments. Then, the fast transitive reduction algorithm (Myers, 2005) is used to perform the transitive reduction of the graph.

As the overlap detection based on probe matching produces somewhat imperfect results, it is important to deal with possible false overlaps in the graph. Upon performing the transitive reduction, the count of edges that could be reduced using the given edge is stored for each remaining edge. A vertex in the graph is defined to be inconsistent if it has more than one in-edge or out-edge. Every inconsistent vertex is checked to see if some edges causing the inconsistency can be removed. The edge is removed if it was not used to reduce any other edge in the transitive reduction.

Missing overlaps usually cause the transitive reduction to produce two separate chains of vertices with the same direction between the two vertices. A chain represents a path of vertices where each vertex has exactly one in-edge and one out-edge. If the second chain contains either no vertices or only one intermediate vertex, it is likely to be caused by a missing overlap. In such cases, the shorter chain is removed from the graph.

The probability that a consistent chain of fragments in the graph is traversed once in the reconstruction is given by

$$\binom{N}{k} \binom{\Delta}{G} \left(\frac{G-\Delta}{G} \right)^{N-k} \approx \frac{\left(\frac{\Delta N}{G} \right)^k}{k!} e^{-\frac{\Delta N}{G}} \quad (4)$$

where k is the number of fragments in the chain, G is the size of the genome in base pairs, Δ is the length of the fragment chain in base pairs, and N is the total number of fragments (Myers, 2005). The probability that the chain is traversed twice can be constructed in a similar fashion. An A-statistic for the chain is computed by taking the logarithm of the ratio of these probabilities (Myers, 2005): $A(\Delta, k) = \Delta N / G - k \ln 2$.

Upper and lower bounds for the number of traversals for each edge in the graph can be determined by setting a threshold on the A-statistic. In this work, the best results were obtained by using a threshold value of 5.

The actual number of traversals is obtained by solving this problem as a minimum cost flow problem (Myers, 2005). In this work, the cycle canceling algorithm was used (Bang-Jensen & Gutin, 2001). The cost for each edge was calculated from the amount of probe hit coverage detected in the overlap phase. Higher probe hit coverage results in lower cost, and lower hit coverage increases the cost.

In this work, repeats were not assembled. Consequently, all vertices with traversal count higher than one were removed.

Algorithm 1 Algorithm for consensus graph construction. C is a list of fragments in a given contig, ordered by their approximate left positions. The output of the algorithm is the consensus graph G . T is the maximum distance for node combination threshold and q is the length of the q -mer.

```

1:  $G \leftarrow \{\}$ 
2:  $c \leftarrow 0$ 
3: for each  $F \in C$  do
4:    $cpos \leftarrow \text{leftPosition}[F]$ 
5:    $last \leftarrow \text{NULL}$ 
6:    $c \leftarrow c + 1$ 
7:   while  $cpos < \text{length}[F] + \text{leftPosition}[F] - q$ 
   do
8:      $g \leftarrow F[cpos:cpos+q-1]$ 
9:     find  $n$  where  $|n.pos-cpos| < T$ ,  $n.mark \neq c$ 
       and  $n.mer = g$ 
10:    if  $n=\text{NULL}$  then
11:       $n \leftarrow \text{new node}(g,cpos)$ 
12:       $G.addVertex(n)$ 
13:    end if
14:    if  $last \neq \text{NULL}$  then
15:       $e \leftarrow G.getEdge(last,n)$ 
16:      if  $e = \text{NULL}$  then  $e \leftarrow G.addEdge(last,n)$ 
      endif
17:       $e.score \leftarrow e.score + \text{quality}[F][cpos+q-1]$ 
18:    end if
19:     $n.mark \leftarrow c$ 
20:     $last \leftarrow n$ 
21:     $cpos \leftarrow cpos + 1$ 
22:  end while
23: end for

```

2.3 The Consensus Phase

The order of the fragments was determined in the layout phase and the relative overlap length between fragments was computed in the overlap phase. The approximate position for each fragment within its contig can now be determined.

A structure called a *consensus graph* is used to efficiently produce the final consensus. The construction of the consensus graph for a given contig begins by dividing each fragment into a sequence of $n - q + 1$ q -mers, where n is the length of the fragment. An edge from q -mer A to B is added only if A and B come from the same fragment, and B is right next to A in the fragment. The q -mers are placed into a hash table and are merged into the same node in the graph only if their approximate positions are close enough in the contig.

The pseudocode of the consensus graph construction algorithm is given in Algorithm 1. The algorithm

efficiently calculates consecutive q -mers (line 8) using bit shifts. The `bf` find operation on line 9 can be efficiently implemented by placing all nodes in a hash table, where the corresponding q -mer is used as a key. If no node with required characteristics is found, a NULL is returned. In computing the score (line 17), PHRED quality scores (Ewing & Green, 1998), are taken into consideration.

Next, the path with the largest sum of scores yields a contig. These contigs are used as new fragments, and the algorithm is executed once more. The program terminates when two consecutive runs produce similar output results.

3 Experimental Results

PMSGGA described in the previous section was tested under several settings. The data sets used to test the algorithm were obtained from the NCBI assembly archive (NCBI, 2008). Additionally, artificial data sets were created using the ReadSim simulator (Schmid & al., 2008). All results were compared to PHRAP (Green, 1999) and CAP3 (Huang & Madan, 1999) assemblies. Only data sets with ID 1, 2, and 3 were tested with EULER (Pevzner & al., 2001). The EULER assembler could not be tested with the larger data sets, with ID 4 to 8, due to high memory requirements (see Tables 2 and 3).

The measuring criteria given in Table 1 were used to evaluate the performance of the algorithms.

Table 1: The measuring criteria.

Runtime	The time to perform a full assembly.
Contigs	Number of noncontiguous sequences at the end of the assembly.
Errors	Number of misassembled fragments.
Identity	Percentage of identical nucleotides in the pairwise alignment of the output and the original sequence.
Sequence covered	Percentage of the original sequence covered by the contigs in the output.
N50	The N50 length is the length x such that 50% of the sequence is contained in contigs of length x or greater (Waterston & al., 2003).

The first five data sets obtained from the NCBI assembly archive are described in Table 2. This set contains three viruses and two bacteria.

The various assemblers were tested with the data sets of Table 2. The results are reported in Table 3. As can be seen in Table 3, PMSGGA was considerably faster. PMSGGA also produced fewer contigs than the

Table 2: Prokaryotic data sets retrieved from NCBI assembly archive (NCBI, 2008).

ID	Name	Length	Seq. Covered	Avg. Error
1	<i>Antelope coronavirus US/OH1/2003</i>	31 kbp	15.8	3.1%
2	<i>Bacteriophage KVP40</i>	240 kbp	10.7	4.1%
3	<i>Chlorella virus MT325</i>	310 kbp	11.1	1.4%
4	<i>Campylobacter fetus subsp. fetus 82-40</i>	1.8 Mbp	10.3	1.2%
5	<i>Streptococcus agalactiae A909</i>	2.1 Mbp	9.0	2.6%

other assemblers. In 3 out of the 5 cases, PMSGGA yielded one contig. In test cases with ID 4 and 5, the output coverage of PMSGGA was slightly lower when compared to the results of CAP3 and PHRAP. This is probably due to the fact that some fragments that contain repeats were not considered in the final layout.

Next, the assemblers were tested with the eukaryotic genomes described in Table 5. To create data sets with ID 6, 7, and 8, the DNA sequences were first downloaded from the NCBI genome database. Then, the fragments were created from the sequences, by using the ReadSim simulator. In these data sets, the fragment length was uniformly distributed between 600 and 1000 base pairs. Note that PHRED quality scores are not available for data sets generated with ReadSim.

Table 5: Eukaryotic data sets generated using ReadSim simulator (Schmid & al., 2008).

ID	Name	Length	Seq. Covered	Avg. Error
6	<i>Theileria parva strain muguga</i> Chromosome 1	2.5 Mbp	8.0	3.0%
7	<i>Pichia stipitis</i> Chromosome 1	3.5 Mbp	8.0	2.0%
8	<i>Schizosaccharomyces pombe</i> Chromosome 1	5.6 Mbp	7.0	2.0%

The results are reported in Table 4. Once again, PMSGGA outperformed the other assemblers in terms of speed. PMSGGA produced fewer contigs than CAP3. The number of contigs obtained by PMSGGA is comparable to those obtained by PHRAP. As for coverage, PMSGGA's results vary between 97.5% and 99.2%.

Next, the percentage of time PMSGGA spent on the

Table 3: Results for assemblies of data sets described in Table 2.

ID	Algorithm	Runtime	Coverage	Identity	Contigs	N50	Errors
1	EULER	5 m	97.8%	98.9%	5	7,676	0
	CAP3	1 m 20 s	100%	99.9%	2	19,489	0
	PHRAP	11 s	100%	99.8%	5	30,994	0
	PMSGGA	4 s	100%	100%	1	30,994	0
2	EULER	38 m	61.6%	99.4%	87	6,517	0
	CAP3	8 m 20 s	100%	99.4%	118	3,679	0
	PHRAP	1 m 10 s	100%	99.8%	16	94,898	0
	PMSGGA	27 s	100%	100%	1	244,834	0
3	EULER	39 m	99.7%	99.6%	4	256,318	1
	CAP3	11 m	100%	99.9%	24	22,941	1
	PHRAP	1 m 30 s	100%	100%	1	314,327	0
	PMSGGA	20 s	99.5%	100%	1	312,631	0
4	CAP3	1 h 40 m	99.2%	99.9%	156	18,101	5
	PHRAP	13 m	99.5%	100%	19	218,453	8
	PMSGGA	2 m 30 s	96.3%	100%	12	619,222	4
5	CAP3	2 h 2 m	99.1%	99.0%	1879	1,868	6
	PHRAP	20 m	98.9%	99.9%	399	39,775	7
	PMSGGA	8 m 30 s	91.1%	99.9%	47	87,267	3

Table 4: Results for assemblies of data sets described in Table 5.

ID	Algorithm	Runtime	Coverage	Identity	Contigs	N50	Errors
6	CAP3	40 m	100%	100%	57	77,313	6
	PHRAP	7 m	100%	100%	14	281,034	2
	PMSGGA	2 m 50 s	99.2%	100%	15	187,017	2
7	CAP3	57 m	99.9%	100%	56	103,076	5
	PHRAP	11 m	99.8%	100%	27	178,492	3
	PMSGGA	4 m 20 s	97.9%	100%	34	147,755	3
8	CAP3	1 h 20 m	99.7%	100%	190	54,555	5
	PHRAP	15 m	100%	100%	79	117,652	7
	PMSGGA	8 m	97.5%	100%	95	97,404	7

Table 6: Distribution of time spent on the 4 tasks.

ID	Pattern matching	Overlap construction	Graph construction	Consensus
1	20%	46%	4%	23%
2	15%	51%	8%	23%
3	19%	44%	6%	26%
4	18%	39%	15%	33%
5	6%	44%	16%	33%

four different tasks was measured. The data sets from Table 2 were used, and the runtimes spent on pattern matching, overlap list construction, string graph construction and consensus building were recorded. The results of this experiment are presented in Table 6. Note that the sum of the percentages reported in Table 6 does not add to 100%. The time spent on tasks such as file I/O and probe selection is not reported. As can be seen in Table 6, the graph construction phase is generally the fastest one, followed by pattern matching. Overlap construction is the slowest phase. The

algorithm was implemented in C++. All tests were run under Linux on AMD Opteron 246/2 GHz dual-processor CPU with 6 gigabytes of memory. Only one processor was used in testing.

4 Conclusion

This article introduced a fast and efficient DNA fragment assembler, PMSGGA. PMSGGA is faster than other commonly used assemblers such as PHRAP, CAP3, and EULER. PMSGGA's speed is due to the BG algorithm for pattern matching used in the overlap phase and the algorithm for the consensus graph construction. In terms of assembly quality, the number of contigs and the coverage produced by PMSGGA are either better or comparable to the results obtained by PHRAP.

The experimental results of this work demonstrated that PMSGGA is successful in assembling prokaryotic genomes as well as eukaryotic genomes.

We are aware that next-generation sequencing methods are lurking on the horizon. We are aware that non-Sanger-based sequencing technologies, such as the ones being developed by Solexa/Illumina, Agencourt/ABI, and Helicos Biosciences, are delivering on their promise of sequencing DNA at unprecedented speed and that one day they will enable impressive scientific achievements and novel biological applications. But as long as Sanger-based sequencing methods are being used, the DNA fragment assembly problem addressed in this work, remains a current and challenging problem that requires efficient algorithms such as PSMGA.

PMSGGA can further be improved by taking steps to handle long repeats. Another option would be to let the researcher manually perform additional tests to decide the final path in the graph, as suggested by Myers (2005). A further improvement could be achieved by parallelizing the overlap phase.

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