## Comparative Studies of *de novo* Assembly Tools for Nextgeneration Sequencing Technologies

Yong Lin<sup>1, 3</sup>, Jian Li<sup>2</sup>, Hui Shen<sup>2</sup>, Lei Zhang<sup>1, 3</sup>, Christopher J Papasian<sup>3</sup>, Hong-Wen Deng<sup>1, 2, 3, \*</sup>

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#### **ABSTRACT**

**Motivation:** Several new *de novo* assembly tools have been developed recently to assemble short sequencing reads generated by next-generation sequencing platforms. However, the performance of these tools under various conditions has not been fully investigated, and sufficient information is not currently available for informed decisions to be made regarding the tool that would be most likely to produce the best performance under a specific set of conditions.

Results: We studied and compared the performance of commonly used *de novo* assembly tools specifically designed for next-generation sequencing data, including SSAKE, VCAKE, Euler-sr, Edena, Velvet, ABySS and SOAPdenovo. Tools were compared using several performance criteria, including N50 length, sequence coverage, and assembly accuracy. Various properties of read data, including single-end/paired-end, sequence GC content, depth of coverage and base calling error rates, were investigated for their effects on the performance of different assembly tools. We also compared the computation time and memory usage of these seven tools. Based on the results of our comparison, the relative performance of individual tools are summarized and tentative guidelines for optimal selection of different assembly tools, under different conditions, are provided.

Contact: hdeng2@tulane.edu.

Supplementary information: Supplementary data are available.

#### 1 INTRODUCTION

Recently developed next-generation sequencing platforms, such as the Roche 454 GS-FLX System, Illumina Genome Analyzer and HiSeq 2000 system, and ABI SOLiD™ System, have revolutionized the field of biology and medical research (Schuster, 2008). Compared to traditional Sanger sequencing technology (Bentley, 2006; Sanger, et al., 1977), these new sequencing platforms generate data much faster and produce much higher sequencing output, while decreasing costs by more than a thousand fold (Shendure and Ji, 2008). The ability to rapidly generate enormous numbers of sequence reads at markedly reduced prices has greatly extended the scope of economically feasible sequencing projects. The prospect of sequencing the entire human genome for a large number of samples has become a reality.

\*To whom correspondence should be addressed.

These new sequencing technologies also pose tremendous challenges to traditional *de novo* assembly tools designed for Sanger sequencing, as they are incapable of handling the millions to billions of short reads (35-400bp each) generated by next-generation sequencing platforms (Dohm, et al., 2007). Therefore, several novel *de novo* assembly tools have been developed, such as SSAKE (Warren, et al., 2007), VCAKE (Jeck, et al., 2007), SHARCGS (Dohm, et al., 2007), Euler-sr (Chaisson and Pevzner, 2008), Edena (Hernandez, et al., 2008), Velvet (Zerbino and Birney, 2008), Celera WGA Assembler (Miller, et al., 2008), ABySS (Simpson, et al., 2009) and SOAPdenovo (Li, et al., 2009).

With the recent introduction of multiple *de novo* assembly tools, it has become necessary to systematically analyze their relative performance under various conditions so that researchers can select a tool that would produce optimal results according to the read properties and their specific requirements. Zhang et al. (Zhang, et al., 2011) recently compared the performance of several of these tools for assembling sequences of different species. Although they evaluated multiple criteria such as run time, RAM usage, N50, and assembly accuracy, their results were based on simulation reads using only a single depth of coverage (100x) and a single base call error rate (1.0%). Further investigation is necessary to determine whether, and how, these assembly tools are differentially affected by varying depths of coverage, sequencing errors, read lengths, and extent of GC content of the sequence reads. Furthermore, the assembly performance of SOAPdenovo (v1.05) has dramatically improved for long read assembly. Consequently, sufficient information is not currently available for informed decisions to be made regarding the tool that would be most likely to produce the best results, based on variations in the practical conditions identified

Accordingly, in this study, we systematically studied and compared the performance of seven commonly used *de novo* assembly tools for next-generation sequencing technologies, using a number of metrics including N50 length (a standard measure of assembly connectivity, to be more specifically defined later), sequence coverage, assembly accuracy, computation time, and computer memory requirement and usage. To imitate different practical conditions, we selected a number of experimentally derived benchmark sequences with different lengths and extent of GC content, and

<sup>&</sup>lt;sup>1</sup> Center of System Biomedical Sciences, University of Shanghai for Science and Technology, Shanghai 200093, P. R. China.

<sup>&</sup>lt;sup>2</sup> School of Public Health and Tropical Medicine, Tulane University, New Orleans, LA 70112, USA

<sup>&</sup>lt;sup>3</sup> School of Medicine, University of Missouri-Kansas City, Kansas City, MO 64108, USA.

simulated single-end and paired-end reads with varying depths of coverage, base calling error rates, and individual read lengths. Based on the results of our analyses, we have developed guidelines for optimal selection of different assembly tools under different practical conditions. Identifying and recognizing the various limitations of specific tools under different practical conditions may also provide useful guidance and direction for improving current tools and/or designing new high-performance tools.

## 2 METHODS AND MATERIALS

## 2.1 De novo sequencing tools

Seven tools, SSAKE (v3.7), VCAKE(vcakec\_2.0), Euler-sr (v1.1.2), Edena (2.1.1), Velvet (v1.0.18), ABySS (v1.2.6) and SOAPdenovo (v1.05 for 64bit Linux), were selected for studies and comparative analyses. These tools are all publicly available, and most of these tools are currently often used to assemble short reads generated by next-generation sequencing platforms, such as Illumina Genome Analyzer (read length=35-150 bp) and ABI SOLID (read length=35-75 bp). Of these seven tools, all are capable of assembling single-end reads, but only SSAKE, Euler-sr, Velvet, ABySS and SOAPdenovo support paired-end reads assembly.

### 2.2 Benchmark sequences

Eight experimentally determined sequences (Table 1) were obtained from the NCBI database (http://www.ncbi.nlm.nih.gov/) and used as benchmark sequences to test the performance of the seven assembly tools. These sequences range from ~99 kbp (base pair) to ~100 Mbp, each with a different extent of GC content.

Table 1. Information for the eight benchmark sequences used in this study

Species	GenBank	Chr.	Seq.len(bp)	GC(%)
D.mel	AC018485	2L	99,441	36.90
H.inf	NC_007146	_	1,914,490	38.16
T.bru	AE017150	2	1,193,948	44.38
H.sap	NT_037622	4	1,413,146	49.81
E.coli	NC_009800	_	4,643,538	50.82
C.ele	NC_003283	V	20,919,568	35.43
H.sap	NT_007819	7	50,360,631	41.03
H.sap	NT_005612	3	100,537,107	38.96

D.mel: Drosophila melanogaster, H.inf: Haemophilus influenza, T.bru: Trypanosoma brucei, H.sap: Homo sapiens, E.coli: Escherichia coli, C.ele: Caenorhabditis elegans; GenBank: GenBank accession number; GC: percentage of GC contents reported by Tandem repeats finder (v4.40, http://tandem.bu.edu/trf/trf.html). H.inf and E.coli are the complete genomes. For clarity, H.sap-1 was used to refer to NT\_037622, H.sap-2 was NT\_007819 and H.sap-3 was NT\_005612.

## 2.3 Sequencing read simulations

Simulated single-end and paired-end reads were generated from benchmark sequences with several variable parameters, including depth of coverage, base calling error rate (BCER), and individual read length. Depth of coverage is the average number of reads by which any position of an assembly is independently determined (Taudien, et al., 2006). BCER is the estimated probability of error for each base-call (Ewing and Green, 1998).

Single-end reads simulation method was the same as that used previously (Dohm, et al., 2007), that is, each read was generated as a DNA fragment of the preset read length from any position in the benchmark sequence with

equal probability. Each base of the read was then randomly and independently changed into another base with probability of BCER. In paired-end read simulation, a fragment with length of fragment size was randomly obtained from the benchmark sequence, then two reads of the preset read length were generated simultaneously from the two ends of this fragment, which were considered as one pair. We applied the fragment size distribution based on the empirical distribution of the experimental read data set of the *E. coli* library (Genbank accession no. SRX000429) (Supplemental Fig 1). The simulation of base calling errors was the same as that of single-end read errors.

The total number of reads was determined by the following formula:

$$Num_{Read} = \frac{Benchmark \ Sequence \ Length \times Depth \ of \ Coverage}{Individual \ Read \ Length}$$

To study and compare the seven selected *de novo* assembly algorithms, sequencing reads were simulated as follows.

- To determine how assembly performance was affected by different depths of coverage and GC contents, single-end reads (BCER = 0.6%, read length=35, 50 and 75bp) and paired-end reads (BCER = 0.6%, read length=35bp\*2, 75bp\*2, 125bp\*2) were generated from four benchmark sequences (sequences 1-4 in Table 1), in which GC content was ~36-50%.
- 2. To determine how the assembly performance was affected by different BCER, sequencing reads were generated with BCER set to 0.0%, 0.2%, 0.4%, 0.6%, 0.8% and 1.0%. Three benchmark sequences (sequences 1-3 in Table 1) were selected for the simulation. In single-end reads assembly, read length was 35bp, and depth of coverage was set to 30x and 70x. In paired-end reads assembly, read length was 35bp\*2 and depth of coverage was 30x and 70x.
- 3. To compare required computational demand (runtime and computer memory usage) of the seven tools, four benchmark sequences with gradually increasing lengths ranging from ~5 million bp to ~100 million bp, (sequences 5-8 in Table 1), were selected for simulation. BCER was set to 0.6%, individual read lengths were set to 35 bp for single-end and 35bp\*2 for paired-end reads, and depth of coverage was set to 70x.

## 2.4 Runtime settings

Runtime parameters for the seven assembly tools were generally set to the default or recommended values of each method with a few exceptions: for VCAKE, the runtime parameter c was set as 0.7 in order to make it consistent with SSAKE. (Each base call in VCAKE was dependent on a voting result; when the votes were totaled and the base proportion exceeded a threshold, c, that base was added to the output contig (Jeck, et al., 2007)). Parameter k for Velvet, ABySS, SOAPdenovo and parameter m for Edena should vary with read length in order to get good N50 lengths. Since no clear default settings for these parameters were presented in the manuals for the corresponding tool, we established values for k and m that produced relatively optimal N50 lengths, based on our own preliminary empirical testing of conditions for each tool. Specific values of the parameters k and m are provided in supplemental Table 1.

Most of the assembly was carried out on a cluster with 8 computer nodes, with each node consisting of dual Quad-Core (2.40 GHz) processors and 12GB RAM. Comparison tests of required computational demand were performed on a server with dual Quad-Core (2.40 GHz) Processors and 32GB RAM.

## 2.5 Performance evaluation

The seven selected *de novo* assembly tools were applied to assemble the simulated sequencing reads into contigs. In paired-end assembly, tools that

support paired-end reads performed an additional step of scaffold construction to get the final output contigs. Contigs with lengths greater than 100bp were used to evaluate the performance of each tool. Each simulation and assembly was conducted five times, and the assembly results were set as the average values.

The performance of each tool was measured by a number of metrics, including N50 length, sequence coverage, assembly error rate, computation time, and computer memory usage. N50 length is the longest length such that at least 50% of all base-pairs are contained in contigs of this length or larger (Lander, et al., 2001). N50 length provides a standard measure of assembly connectivity, reflecting the nature of the bulk of the assembly rather than the cutoff which defines the smallest reportable assembly unit (Jaffe, et al., 2003). Higher N50 length indicate better performance of the assembly tool. Sequence coverage refers to the percentage of the benchmark sequence covered by output contigs. In the calculation of assembly error rates, we aligned the output contigs to the benchmark sequence, and calculated the number of mismatched bases from alignment results. The assembly error rate was the percentage of these mismatched bases in the total bases of aligned contigs in the reference sequence. Sequence coverage and assembly error rates were analyzed by blastz (Schwartz, et al., 2003).

#### 3 RESULTS

# 3.1 Assembly performance affected by depth of coverage and GC content

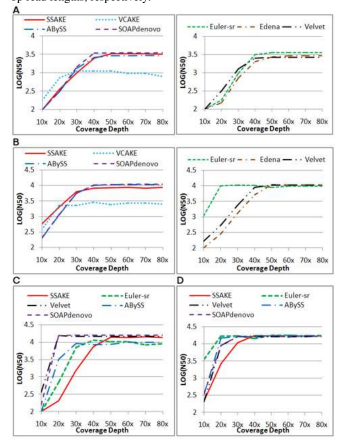
To determine whether, and how, the assembly performance of the seven tools was differentially affected by the depth of coverage and extent of GC content in the source sequences, these tools were used to assemble simulated sequence reads (BCER = 0.6%) generated from different benchmark sequences (GC content =  $\sim 36-50\%$ ) at different depths of coverage. Assembly performance of the seven tools is illustrated in Figure 1 and Tables 2-5. Figure 1 and Tables 4-5 present test results for part of a benchmark sequence as an example, but similar results were obtained for the other benchmark sequences tested (Supplementary Tables 2-9).

With increasing depths of coverage, the performance of these seven tools showed some interesting patterns (Figure 1) in assembly connectivity measured by N50 length. Although there was an initial increase in N50 lengths with increasing depth of coverage, N50 lengths reached a plateau when the depth of coverage reached a certain threshold. For simplicity, DCAP will be used here to refer to the depth of coverage at which the N50 length plateau was reached.

In single-end assembly, DCAP for SSAKE and Edena(~50x) was greater than that for VCAKE, Velvet, ABySS and SOAPdenovo(30-40x); DCAPs for Euler-sr varied with read length (~50x when read length was 35bp and ~20x when read length was 75bp). In paired-end assembly, DCAPs for most tools were lower than those observed in single-end assembly. DCAPs for SSAKE(~40x) was still greater than that for Velvet, ABySS and SOAPdenovo(20-30x); DCAPs for Euler-sr varied with read length (~40x when read length was 35bp\*2 and ~20x when read length was 75bp\*2).

To compare N50 values among the various tools, we chose N50 values at a depth of coverage of 70x, because this exceeded the DCAP for all tools (Table 2, 3). General observations for N50 values of these tools under these various conditions are described below. Comparison results varied with different read lengths and GC content. Sequences with a GC content of 36.90% and 38.16% are referred to as "Low GC content", whereas, those with a GC

content of 44.38% and 49.81% are referred to as "High GC content". Similarly, "short read" and "long read" refer to 35bp and 75 bp read lengths, respectively.



**Fig 1.** Comparison of the effect of various coverage depths on N50 length in *T.bru* assembly when BCER was 0.6%. A. Single-end reads assembly, read length (RL)=35bp; B. Single-end assembly, RL=75bp; C. Paired-end reads assembly, RL=35bp; D. Paired-end assembly, RL=75bp.

**Table 2.** Comparison of N50 lengths in assembly of single-end reads when depth of coverage was 70x and BCER was 0.6%

Seq	RL	SS	VC	Eu	Ed	Ve	AB	SO
D.mel	35bp	6717	2215	9064	4917	4085	4087	4145
H.inf		25558	2669	26491	19231	17988	18547	22036
T.bru		3264	963	3528	2934	2667	3014	3504
H.sap-1		1177	653	1393	1053	910	961	1202
D.mel		28646	3683	23676	22695	22679	22673	25115
H.inf	75bp	46069	3235	38667	38724	38715	38361	42778
T.bru		8205	2682	9733	10847	10682	10814	11108
H.sap-1		2706	691	2169	4315	3810	3358	5227

RL: read length; Seq: Benchmark Sequence. SS: SSAKE. VC: VCAKE. Eu: Euler-sr. Ed: Edena. Ve: Velvet. AB: ABySS. SO: SOAPdenovo.

In single-end reads assembly, with:

- Low GC content and short read: N50euler-sr  $\geq$  N50ssake>N50soapdenovo  $\approx$  N50edena>N50volvet  $\approx$  N50abyss>N50voake.
- Low GC content and long read: N50ssake> N50soapdenovo> N50edena ≈ N50velvet ≈ N50abyss ≈ N50euler-sr > N50vcake.

- High GC content and short read:  $N50_{EULER-sr} \ge N50_{SOAPdenovo}$  $\approx N50_{SSAKE} > N50_{Edena} \approx N50_{Velvet} \approx N50_{ABvSS} > N50_{VCAKE}$ .
- High GC content and long read:  $N50_{SOAPdenovo} > N50_{Edena} > N50_{Velvet} \approx N50_{ABySS} > N50_{SSAKE} > N50_{EULER-sr} > N50_{VCAKE}$ .

**Table 3.** Comparison of N50 length in assembly of paired-end reads when depth of coverage was 70x and BCER was 0.6%

Seq. (GC %)	RL.	SS	Eu	Ve	AB	SO
D.mel (36.90)		29771	27326	28604	29892	30308
H.inf (38.16)	35bp	91821	90275	92349	93956	119805
T.bru (44.38)	ээор	14470	9498	14948	9998	15598
H.sap-1 (49.81)		3188	3116	4730	4281	14972
D.mel (36.90)		29963	29029	29676	30923	30863
H.inf (38.16)	75bp	122151	107232	120699	120175	120886
T.bru (44.38)	750p	16768	17051	16094	17566	16326
H.sap-1 (49.81)		7436	4041	34592	33429	34265

GC: GC content

In paired-end reads assembly:

- SOAPdenovo generated the greatest N50 lengths in almost all tests
- SSAKE generated relatively high N50 lengths when GC content was low.
- N50 lengths for Velvet and ABySS were comparable to one another for all tests.
- N50 lengths for Velvet and ABySS were comparable to SOAPdenovo when assembling long reads.
- N50 lengths for Euler-sr were the lowest for almost all tests.

## 3.2 Assembly performance with regard to sequence coverage and assembly error rate

Using benchmark sequences *D.mel* and *T.bru* as examples, we compared assembly performance of the seven tools with regard to sequence coverage and assembly error rate (Tables 4, 5). Generally, long reads resulted in high sequence coverage and assembly error rates.

In single-end reads assembly:

- SSAKE and VCAKE were comparable to one another, and provided higher sequence coverage than the other tools. Sequence coverage for SOAPdenovo was a little lower, but very close to SSAKE when assembling long reads (75 bp).
- Edena, Velvet and ABySS were clustered together, with slightly lower sequence coverage than SOAPdenovo.
- Euler generated the lowest sequence coverage for almost all tests.
- ABySS showed the lowest assembly error rates for almost all tests
- SSAKE, VCAKE, SOAPdenovo and Euler-sr generated higher assembly error rates than Edena, Velvet and ABySS.

In paired-end reads assembly:

- Sequence coverage comparisons had the following relationships: SC  $_{ABySS}$  >SC  $_{SOAPdenovo}$   $\approx$  SC  $_{SSAKE}$   $\approx$  SC  $_{Velvet}$  >SC  $_{Euler-}$
- ABySS showed the lowest assembly error rates for almost all tests.

**Table 4.** Comparison of sequence coverage and assembly error rates in assembly of single-end reads with various GC contents and depths of coverage (BCER = 0.6%)

	RL (bp)	Seq (GC%)	DC	SS	VC	Eu	Ed	Ve	AB	SO
		D1	30x	79.48	78.76	75.44	77.17	77.43	77.55	78.70
		D.mel (36.90)	50x	79.74	77.60	76.33	78.55	77.97	78.29	78.59
	35	(30.90)	70x	79.54	77.70	76.40	78.33	77.86	78.06	78.62
	33	T 1	30x	72.64	71.01	68.07	67.02	67.78	67.19	68.74
		<i>T.bru</i> (44.38)	50x	73.16	70.39	68.40	67.45	67.94	67.21	68.65
SC		(44.36)	70x	73.56	70.40	68.59	67.27	67.58	67.15	68.76
(%)		Danal	30x	80.93	79.44	78.94	78.41	79.93	79.92	80.09
	75	D.mel (36.90)	50x	80.13	79.45	78.69	80.58	79.83	79.82	80.49
			70x	80.99	79.83	79.40	80.02	79.99	79.83	80.82
		T.bru (44.38)	30x	77.92	77.12	71.02	75.67	74.84	74.83	76.99
			50x	77.57	78.43	70.60	76.20	74.91	74.66	76.59
			70x	78.68	78.38	71.48	76.99	74.86	75.02	76.70
		D.mel (36.90)	30x	0.31	0.27	0.34	0.23	0.28	0.26	0.32
			50x	0.39	0.29	0.36	0.33	0.29	0.23	0.32
	35		70x	0.32	0.24	0.38	0.26	0.23	0.26	0.39
		<i>T.bru</i> (44.38)	30x	0.27	0.17	0.25	0.08	0.07	0.04	0.16
			50x	0.32	0.14	0.26	0.07	0.05	0.04	0.14
AER			70x	0.33	0.16	0.26	0.09	0.06	0.04	0.10
(%)		Dmal	30x	0.42	0.75	0.42	0.53	0.28	0.23	0.39
		D.mel (36.90)	50x	0.42	0.76	0.45	0.49	0.37	0.29	0.41
	75	(30.90)	70x	0.45	0.79	0.49	0.53	0.35	0.31	0.43
	13	T.bru	30x	0.63	0.67	0.47	0.59	0.46	0.42	0.66
		(44.38)	50x	0.56	0.84	0.52	0.46	0.49	0.48	0.63
			70x	0.65	0.88	0.48	0.53	0.46	0.49	0.67

SC: Sequence Coverage; AER: Assembly Error Rate.

**Table 5.** Comparison of sequence coverage and assembly error rates in assembly of paired-end reads with various GC contents and depths of coverage (BCER = 0.6%)

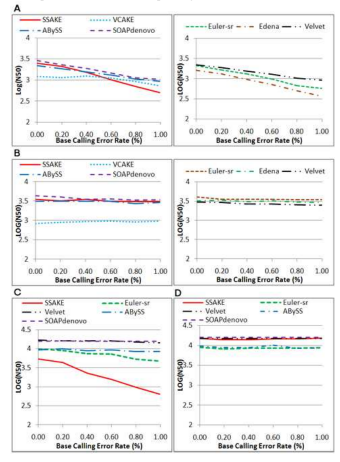
	RL (bp)	Seq (GC%)	DC	SS	Eu	Ve	AB	SO
		D.mel	30x	77.05	71.12	78.75	79.53	78.85
		(36.90)	50x	79.16	71.98	79.03	79.65	78.52
	35	(30.90)	70x	79.11	70.61	78.95	79.71	78.92
	33	T.bru	30x	72.69	71.07	71.37	73.46	70.07
		(44.38)	50x	72.50	71.67	71.50	73.11	70.34
SC		(44.50)	70x	73.59	69.08	71.32	73.27	70.78
(%)		D.mel	30x	79.77	71.31	79.18	81.00	80.20
		(36.90)	50x	79.88	70.17	78.79	80.82	80.36
	75		70x	79.73	69.54	78.43	81.59	79.69
		T.bru (44.38)	30x	77.19	71.56	76.52	81.97	76.25
			50x	78.29	70.12	76.66	81.97	76.54
			70x	78.49	70.86	78.17	81.10	76.55
		D.mel (36.90)	30x	0.33	0.54	0.17	0.14	0.28
			50x	0.32	0.52	0.19	0.16	0.29
	35	(30.90)	70x	0.34	0.44	0.21	0.19	0.33
	33	T.bru	30x	0.30	1.30	0.25	0.19	0.21
		(44.38)	50x	0.34	0.87	0.22	0.17	0.23
AER			70x	0.40	0.73	0.21	0.21	0.20
(%)		Danal	30x	0.47	0.37	0.27	0.13	0.35
		D.mel (36.90)	50x	0.41	0.48	0.21	0.14	0.42
	75	(30.90)	70x	0.62	0.38	0.23	0.16	0.35
	13	Thm	30x	0.52	0.79	0.55	0.45	0.39
		<i>T.bru</i> (44.38)	50x	0.52	0.61	0.57	0.39	0.43
			70x	0.61	0.72	0.61	0.42	0.42

- SOAPdenovo generated more assembly errors than Velvet in assembly of sequences with low GC content (e.g. *D.mel*) but fewer assembly errors than Velvet in assembly of high GC content sequence (e.g. *T.bru*). The assembly error rate for SOAPdenovo and Velvet were both lower than SSAKE.
- Euler-sr generated the highest assembly error rates for almost all tests.

## 3.3 Assembly performance affected by different BCER

To determine whether, and how, assembly performance of the seven tools was differentially affected by changes in BCER, these tools were applied to assemble sequencing reads simulated from three benchmark sequences (*D.mel*, *H.inf* and *T.bru*) with variable BCER (0.0%-1.0%, with a 0.2% incremental change at every step).

Since similar results were obtained with the three benchmark sequences (Supplementary Tables 10-15), we present the results for sequence *T.bru* as an example (Figure 2).



**Fig 2.** Comparison of the effects of various BCER on N50 length in *T.bru* assembly when read length was 35bp. A. Single-end reads assembly, Depth of Coverage(DC)=30x; B. Single-end assembly, DC=70x; C. Paired-end reads assembly, DC=30x; D. Paired-end assembly, DC=70x.

N50 lengths for all seven tools showed decreasing trends, with increases in BCER, but generated different patterns.

- When depth of coverage was below the DCAP of a tested tool, N50 lengths for the specific tool decreased exponentially with increases in BCER. When depth of coverage was below the DCAP (e.g. 30x), increases in BCER produced more significant decreases in N50 lengths for SSAKE, Edena and Euler-sr than for Velvet, ABySS and SOAPdenovo.
- When depth of coverage exceeded the corresponding DCAP, however, N50 lengths were essentially unaffected by changes in BCER
- For instance, in Figure 2A, N50 lengths decreased with increasing BCER when depth of coverage was at 30x for all tools, but were essentially unaffected by changes in BCER when depth of coverage exceeded their DCAP (e.g., 70x, Figure 2B and 2D).
- Similarly, for paired end assembly at a depth of coverage of 30x, N50 lengths for SSAKE and Euler-sr decreased exponentially with increases in BCER, but N50 lengths for Velvet, ABySS and SOAPdenovo remained stable as BCER increased (Figure 2C). Thus the pattern described above is sustained, because 30x is below DCAP of SSAKE and Euler-sr (~50x), but exceeded DCAP of Velvet, ABySS and SOAPdenovo (20-30x).

## 3.4 Computational demand

When selecting a tool for *de novo* sequence assembly, computational demand by the tool should also be considered. This is particularly important when analyzing large genome sequence data (e.g. human genomes) for large samples. The utility of a tool can be seriously limited if it takes up excessive memory space, consumes too much CPU time and exceeds reasonable execution time. Consequently, we compared the runtime (RT) and resident memory usage (RM) required for the seven tools to assemble large data sets. The test results are presented in Table 6.

- It was not feasible to use some of these tools to assemble large sequences because memory required for the assembly process was beyond our computer power. For instance, SSAKE could not assemble sequences greater than 20 mega bps (*C.ele*, *H.sap-2* and *H.sap-3*). VCAKE and Euler-sr could not assemble sequences greater than 50 mega bps (*H.sap-2*, *H.sap-3*). Edena could not assemble sequence greater than 100 mega bps (*H.sap-3*). Velvet could not assemble pairedend reads of the *H.sap-3* sequence.
- Runtime and RM usage varied dramatically in this test. For all tools, there was an approximately linear increase in memory consumption with increasing benchmark sequence lengths, with RM\_{SSAKE}> RM\_{VCAKE}> RM\_{Edena}> RM\_{Euler-sr}> RM\_{Velvet}> RM\_{ABySS} \geqslant RM\_{SOAPdenovo} in single-end reads assembly and RM\_{SSAKE}> RM\_{Euler-sr}> RM\_{SOAPdenovo}> RM\_{ABySS} in paired-end reads assembly.
- The runtime of these tools also increased approximately linearly with increasing benchmark sequence lengths, with  $RT_{SSAKE} > RT_{VCAKE} > RT_{Euler-sr} > RT_{Edena} > RT_{ABySS} > RT_{Velvet} > RT_{SOAPdenovo}$ .
- Runtime and RM usage for Velvet sometimes became abnormal in paired-end reads assembly of large genomes. For example, in paired-end reads assembly of *H.sap-2*, Velvet consumed much more memory and runtime than ABySS and

	Runtime (s)										
	Bench.Seq	E.coli	C.ele	H.sap-2	H.sap-3						
	(Length: bp)	(4.6M)	(20.9M)	(50.3M)	(100.5M)						
	SSAKE	2,776									
	VCAKE	1,672	16,742								
	Euler-sr	1,689	11,961	29,622							
SE	Edena	895	8,450	17,043							
	Velvet	205	1,003 2,786		6,098						
	ABySS	265	1,300	3,307	6,608						
	SOAPdenovo	62	253	560	1,029						
	SSAKE	9,163									
	Euler-sr	1,455	15,068								
PE	Velvet	229	1,351	55,581							
	ABySS	458	3,081	9,199	21,683						
	SOAPdenovo	78	374	889	2,257						
RAM (MB)											
	Bench.Seq	E.coli	C.ele	H.sap-2	H.sap-3						
	(Length: bp)	(4.6M)	(20.9M)	(50.3M)	(100.5M)						
	SSAKE	9,933									
	VCAKE	4,099	17,408								
	Euler-sr	1,536	7,065	13,312							
SE	Edena	1,741	7,557	30,720							
	Velvet	1,229	4,045	9,830	22528						
	ABySS	1,126	3,993	8,909	18432						
	SOAPdenovo	935	2,867	8,089	18227						
	SSAKE	16,384									
	Euler-sr	1,638	7,578								
PE	Velvet	1,331	5,324	30,720							
	ABySS	950	4,505	9,830	18,432						
	SOAPdenovo	1,638	5,939	10,342	19,456						

*piens*. ABySS produced fewer assembly errors, and consumed a little less memory and more runtime than Velvet. When assembling paired-end reads, ABySS produced the highest assembly coverage of all tools tested. When assembling larger genomes, Velvet sometimes used exceptionally high runtimes and memory.

Edena needs a high depth of coverage, comparable to SSAKE, to reach the DCAP. It produced similar, or greater, N50 values to Velvet in most single-end assemblies, and generated assembly error rates that were comparable to Velvet. The computation demands of Edena were intermediate, between SSAKE and ABySS.

SOAPdenovo was the fastest assembler. Its DCAP was as low as that of ABySS and it produced among the highest N50 values in paired-end read assembly, and relatively high N50 values in single-end assembly. SOAPdenovo generated higher assembly error rates and lower sequence coverage than ABySS. It also consumed more memory than ABySS when assembling paired-end reads. The appropriate setting for SOAPdenovo (SOAPdenovo31mer, SOAPdenovo63mer and SOAPdenovo127mer that support kmer ≤31, ≤63 and ≤127 respectively) must be selected based on read length. SOAPdenovo63mer/SOAPdenovo127mer consumed two/four times as much RAM as SOAPdenovo31mer.

In light of our results, investigators may choose the most appropriate assembly tool(s) to use based on their specific experimental setting and available computational resources. Our results may also serve as a reference, when designing sequencing projects, for selecting targeted depths of coverage, control levels of sequencing error rates, etc. Given the rapid increase in use of next generation sequencing technologies, our results should be of value to both empiricists, during experimental design, and to bio-informaticians who seek guidance for selecting appropriate assembly tool(s) for data analyses and who attempt improvement of the assembly tools.

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