

corsage: Metagenome-enabled error correction of single cell sequencing reads

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Received on XXXXX; revised on XXXXX; accepted on XXXXX

Associate Editor: XXXXXXXX

ABSTRACT

Summary: We present a new tool, corsage, to correct sequencing errors in Illumina data produced from single amplified genomes (SAGs). It uses sequence information derived from accompanying metagenome sequencing to accurately correct errors even in ultra-low coverage regions of the single cells. In evaluations on real data, we show that corsage outperforms BayesHammer, the current (and widely used) state-of-the-art. corsage particularly does well in correcting chimeric reads, which helps to improve both accuracy and contiguity of *de novo* assemblies.

Availability and implementation: corsage is implemented in C and is freely available under the open source MIT license from <https://github.com/metagenomics/corsage>

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1 INTRODUCTION

The vast majority of microbial species observed in nature cannot be grown in pure culture (Rappe and Giovannoni, 2003), turning metagenomics and – more recently – single cell genomics into indispensable methods to study the microbial dark matter (Rinke et al., 2013). Frequently, single amplified genomes (SAGs) and accompanying shotgun metagenomes are generated from the same environmental sample, and were methodologically combined to e.g. validate metagenome bins with single cells (Hess et al., 2011) or to improve the SAG's assembly contiguity (Campbell et al., 2013). But so far, no one explored the idea to combine both approaches to tackle SAG-specific error correction challenges.

A single cell's DNA needs to be amplified prior to sequencing, as usually done by multiple displacement amplification (MDA; Lasken, 2007). This introduces a tremendous coverage bias resulting in uneven sequencing depth (Chitsaz et al., 2011), including ultra-low coverage regions with basically no informed error correction possible (Table S1; Figure S1). Chimera formations during MDA occur roughly once per 10 kbp (Lasken and Stockwell, 2007; Rodrigue et al., 2009), further complicating SAG assembly (Nurk et al., 2013). While there are plenty of error correction tools for a variety of use cases available (Laehnemann et al., 2015), only one tool was specifically designed to correct SAG data: hammer (Medvedev et al., 2011), refined to BayesHammer (Nikolenko et al., 2013).

We propose a metagenome-enabled error correction strategy for single cell sequencing reads, taking advantage of largely unbiased metagenomic coverage at sites with too low a coverage for SAG-only error correction.

2 METHODS

We correct potential sequencing errors using an algorithm similar to solving the *spectral alignment problem* (Pevzner et al., 2001). Given a set of trusted *k*-mers, we attempt to find a sequence with minimal corrections such that each *k*-mer on the corrected sequence is trusted. We consider a *k*-mer trusted if it occurs at least $2\times$ in the accompanying metagenome.

In principle, corsage is similar to the error correction in fermi (Li, 2012). It differs in that it removes fermi's assumption of even sequencing coverage to account for the extreme coverage bias in single cell data. Instead, it exploits metagenomic sequence information to correct errors even in ultra-low coverage regions of the single cell.

corsage corrects errors in three phases:

1. corsage collects all 31-mers occurring in the SAG reads. It uses this information to initialize a hash table with the 31-mers being valid keys.
2. corsage scans the accompanying metagenomic reads. For each stored 31-mer, it counts the occurrence of the next (i.e. the 32-nd) base in the metagenome and stores this information in the hash table. This step is largely I/O bound and dominates corsage's runtime (Table S2).
3. corsage processes each SAG read by using the 31-mer hash table to check if the 32-nd base has sufficient support in the metagenome. Roughly speaking, corsage tries to correct an unsupported base if by looking up its 31-mer prefix we know the base is different from an overwhelmingly frequent 32-nd base in the metagenome.

3 RESULTS AND DISCUSSIONS

As a realistic benchmark, we used eight *Escherichia coli* K12-MG1655 SAGs from Clingenpeel et al., 2014, for which the complete genome sequence is available. We generated an accompanying *in vitro* mock metagenome consisting of 26 microbial species and sequenced it on Illumina's HiSeq (Table S2). Based on measured molarity (mapped metagenome reads), we estimate an abundance of 0.3% (0.24%) for *E. coli*, corresponding to a mean per-base coverage of $20.7\times$.

We evaluated corsage along with BayesHammer (Nikolenko et al., 2013), the current state-of-the-art. corsage corrects more errors than BayesHammer, producing a higher fraction of better and

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Table 1. Performance of error correction

Program	% perfect	% chimeric	% better	% worse
raw	22.52 ± 1.07	0.73 ± 0.15	–	–
hammer	80.35 ± 8.77	0.77 ± 0.17	71.66 ± 2.12	0.33 ± 0.06
corsage	95.52 ± 0.43	0.06 ± 0.02	75.45 ± 1.11	0.26 ± 0.03

Refer to Table S3. Give all versions and the non-default command-line arguments: corsage option '-c 2', BayesHammer-v3.6.0 quality trimming disabled via config file. State that we use BWA-MEM (Li, 2013) and samtools (Li et al., 2009). Given is the mean and std.dev for all 8 *E. coli* SAGs, best in bold.

perfect reads after correction, while maintaining a high accuracy. Notably, corsage also suppresses chimeric reads, potentially due to the metagenome not being MDA-amplified. BayesHammer surprisingly introduces additional chimeric reads.

corsage also works well with single cell assemblers (Table 2). *Discuss results obtained with SPAdes-3.6.0 (Bankevich et al., 2012) and IDBA-UD-1.1.1 (Peng et al., 2012). Less misassembly events, probably because corsage corrects chimeric reads, too. Also slightly increased contiguity while maintaining or surpassing the original accuracy.*

Discuss outliers (6 & 7), better MDA = less coverage bias = better assembly. Here, hybrid error correction shows the least effect. Point to the Table S1. SAGs with highest MDA bias (3 & 4) are harder to correct (more ultra-low coverage regions), and also perform poorest in assembly (Tables S4 and S5). These benefit the most from metagenome-enabled error correction. Incorporating a second source of information (metagenome) has the potential to expand the current state-of-the-art, and produces excellent results.

Discuss, that metagenome-enabled error correction of SAG data might result in miscorrections of rare variants (low-abundant strain captured, reads might get corrected towards the most abundant strain). Argue that these do not matter much, and could be polished in a postprocessing step (e.g. pilon, sequel).

ACKNOWLEDGEMENT

Funding: AB is supported by a fellowship from the CLIB Graduate Cluster Industrial Biotechnology and is partially funded by the

International DFG Research Training Group GRK 1906/1. The work conducted by the U.S. Department of Energy Joint Genome Institute, a DOE Office of Science User Facility, is supported under Contract No. DE-AC02-05CH11231.

REFERENCES

- Bankevich, A. et al. (2012). SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. *J. Comput. Biol.*, 19(5):455–477.
- Campbell, J. H. et al. (2013). UGA is an additional glycine codon in uncultured SR1 bacteria from the human microbiota. *Proc. Natl. Acad. Sci. U.S.A.*
- Chitsaz, H. et al. (2011). Efficient de novo assembly of single-cell bacterial genomes from short-read data sets. *Nat. Biotechnol.*, 29(10):915–921.
- Clingenpeel, S. et al. (2014). Reconstructing each cell's genome within complex microbial communities—dream or reality? *Front Microbiol.*, 5:771.
- Gurevich, A. et al. (2013). QUASt: quality assessment tool for genome assemblies. *Bioinformatics*, 29(8):1072–1075.
- Hess, M. et al. (2011). Metagenomic discovery of biomass-degrading genes and genomes from cow rumen. *Science*, 331(6016):463–467.
- Laehnemann, D. et al. (2015). Denoising DNA deep sequencing data-high-throughput sequencing errors and their correction. *Brief. Bioinformatics*.
- Lasken, R. S. (2007). Single-cell genomic sequencing using Multiple Displacement Amplification. *Curr. Opin. Microbiol.*, 10(5):510–516.
- Lasken, R. S. and Stockwell, T. B. (2007). Mechanism of chimera formation during the Multiple Displacement Amplification reaction. *BMC Biotechnol.*, 7:19.
- Li, H. (2012). Exploring single-sample SNP and INDEL calling with whole-genome de novo assembly. *Bioinformatics*, 28(14):1838–1844.
- Li, H. (2013). Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM. *arXiv:1303.3997*.
- Li, H. et al. (2009). The Sequence Alignment/Map format and SAMtools. *Bioinformatics*, 25(16):2078–2079.
- Medvedev, P. et al. (2011). Error correction of high-throughput sequencing datasets with non-uniform coverage. *Bioinformatics*, 27(13):1137–141.
- Nikolenko, S. I. et al. (2013). BayesHammer: Bayesian clustering for error correction in single-cell sequencing. *BMC Genomics*, 14 Suppl 1:S7.
- Nurk, S. et al. (2013). Assembling single-cell genomes and mini-metagenomes from chimeric MDA products. *J. Comput. Biol.*, 20(10):714–737.
- Peng, Y. et al. (2012). IDBA-UD: a de novo assembler for single-cell and metagenomic sequencing data with highly uneven depth. *Bioinformatics*, 28(11):1420–1428.
- Pevzner, P. A. et al. (2001). An Eulerian path approach to DNA fragment assembly. *Proc. Natl. Acad. Sci. U.S.A.*, 98(17):9748–9753.
- Rappe, M. S. and Giovannoni, S. J. (2003). The uncultured microbial majority. *Annu. Rev. Microbiol.*, 57:369–394.
- Rinke, C. et al. (2013). Insights into the phylogeny and coding potential of microbial dark matter. *Nature*, 499(7459):431–437.
- Rodrigue, S. et al. (2009). Whole genome amplification and de novo assembly of single bacterial cells. *PLoS ONE*, 4(9):e6864.

Table 2. Effect on *de novo* assembly

Assembler	Corrector	NG50	# contigs	Largest contig	Total length	MA	MM	IND	GF (%)	# genes	NGA50
IDBA-UD	raw	49982	394	186696	4410124	15.00	7.38	0.55	93.474	3846	49714
	hammer	48672	400	168302	4411299	14.88	7.69	0.62	93.425	3837	48653
	corsage	64127	327	191229	4422372	6.88	5.72	0.40	93.877	3919	64074
SPAdes	raw	77752	473	212552	4531527	18.88	21.88	1.82	94.247	3889	76859
	hammer	86615	369	222585	4472703	14.50	19.16	1.89	94.041	3902	85054
	corsage	87774	347	228604	4484817	8.88	16.13	1.51	94.268	3934	86690

Refer to Table S4 and S5. Assembled with IDBA-UD-1.1.1 (Peng et al., 2012; default options) and SPAdes-3.6.0 (Bankevich et al., 2012; options '-sc --only-assembler --careful -k 21,33,55,77'). Statistics computed with QUASt-3.0 (Gurevich et al., 2013; default options). Given is the mean for all 8 *E. coli* SAGs, best in bold.

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1 SINGLE AMPLIFIED GENOMES

Describe why and how the SAGs were generated. Cite (Clingenpeel et al., 2014). State average sequencing depth (constant), but different MDA quality levels. Highlight two exceptionally well amplified SAGs (6 & 7), that are closer to isolate-grade genomes. Also highlight the problematic ones mentioned in main text, where the hybrid error correction clearly outperforms SAG-only correction. Table S1.

Table S1. Per-base coverage for the eight *E. coli* SAGs.

SAG	mean	std.dev	min	max	Q1	Q2	Q3
0	289.749	486.983	0	6560	36	112	331
1	297.379	386.285	0	4774	70	169	355
2	272.085	708.577	0	12254	18	57	188
3	260.303	762.898	0	16120	11	49	198
4	284.087	783.65	0	13996	7	43	174
6	266.821	275.48	3	3861	105	179	321
7	278.141	241.48	4	2124	124	195	334
8	299.661	399.265	0	4816	58	168	384

Describe methods: mapping of raw reads with bwa mem (Li, 2013) then samtools to view, sort and depth (Li et al., 2009) options of depth: -q0 -Q0 (with source code modified to remove the hardcoded coverage max). bla bla

2 MOCK METAGENOME

Generated at the JGI, initially for internal benchmarking. NCBI accession number or some way to get it via IMG or GOLD (alternatively, might be okay to provide upon request, contact Tanja?). Describe what's in there. Refer to table with molecular weights and mapping statistics (genome coverage), Table S2.

Probably also include a tree containing all 26 genomes, based on their 16S (or – simpler – extracted from iTOL, as all are known!). This might be of interest to see how closely related the stuff in there is. Discuss with others!

3 METRICS PER SAG

Read-based in Table S3, assembly-based in Tables S4 and S5.

Explain the parameter sweep for -c, the metagenomic coverage threshold. Setting this to 2 already produces much better results, with a higher threshold yielding slightly better results, but more

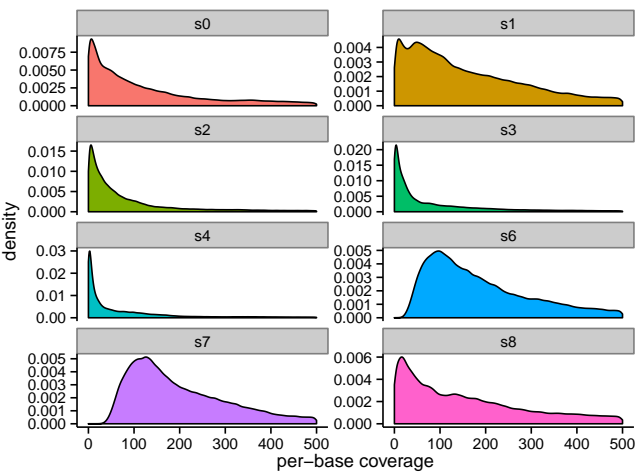


Fig. 1: Per-base coverage for the eight *E. coli* SAGs.

sample-specific. Setting it to 2 should work in all cases, and thus is the default setting. You might get better results if you play around with it, depends on target coverage in the metagenome.

Assembly parameters (k) were tuned to maximize the NG50 value of the BayesHammer + SPAdes assembly. Changing from the default -k 21,33,55 to -k 21,33,55,77 produced better assemblies, thus we picked this combination. --careful reduces the number of misassemblies by (1) less aggressive assembly and (2) polishing of the contigs in a postprocessing step involving read mapping.

REFERENCES

Clingenpeel, S. et al. (2014). Reconstructing each cell's genome within complex microbial communities—dream or reality? *Front Microbiol*, 5:771.
Gurevich, A. et al. (2013). QUAST: quality assessment tool for genome assemblies. *Bioinformatics*, 29(8):1072–1075.
Li, H. (2013). Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM. *arXiv:1303.3997*.
Li, H. et al. (2009). The Sequence Alignment/Map format and SAMtools. *Bioinformatics*, 25(16):2078–2079.

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Table S2. Profile of the mock metagenome.

ContigID	ContigLength	MappedReads	AvgCoverage
Desulfosporosinus acidophilus SJ4, DSM 22704 : Desaci_Contig240.3	3897	367804	13410.8450089813
Desulfosporosinus acidophilus SJ4, DSM 22704 : Desaci_Contig241.2	60447	2309615	5644.16513640048
Desulfosporosinus acidophilus SJ4, DSM 22704 : Desaci_Contig243.1	4926837	50525496	1520.41947886646
Desulfotomaculum gibsoniae Groll, DSM 7213 : Desgi_Contig329.1	4855529	24329020	741.835021889479
Desulfosporosinus meridiei S10, DSM 13257 : Desmer_Contig92.1	4873567	16066750	487.573607585573
Echinicola vietnamensis KMM 6221, DSM 17526 : Echvi_Contig97.1	5608040	2160987	57.2232760465332
Fervidobacterium pennivorans Ven5, DSM 9078 : Ferpe_Contig798.1	2166381	39566833	2708.24050709455
Frateuria aurantia Kondo 67, DSM 6220 : Fraau_Contig24.1	3603458	13922996	568.15908080516
Halovivax ruber XH-70, DSM 18193 : Halru_Contig_35	3223876	6060380	274.952728640928
NC_000913 Escherichia coli str. K-12 substr. MG1655 chromosome.	4639675	647555	20.6521661538793
NC_008261 Clostridium perfringens ATCC 13124, complete genome.	3256683	1461840	66.5510195496461
NC_009012 Clostridium thermocellum ATCC 27405, complete genome.	3843301	1542460	59.6166532363715
NC_012982 Hirschia baltica ATCC 49814, complete genome.	3455622	27502745	1182.5057080896
NC_012983 Hirschia baltica ATCC 49814 plasmid pHbal01	84492	641481	1126.13552762392
NC_014008 Coraliomargarita akajimensis DSM 45221 chromosome.	3750771	11810956	467.028614916773
NC_014168 Segniliparus rotundus DSM 44985 chromosome.	3157527	4886507	225.630138079579
NC_014210 Nocardioopsis dassonvillei subsp. dassonvillei DSM 43111 chromosome.	5767958	5761	0.0573058957780206
NC_014211 Nocardioopsis dassonvillei subsp. dassonvillei DSM 43111 plasmid pNDAS01.	775354	879	0.0682900971685192
NC_014212 Meiothermus silvanus DSM 9946 chromosome.	3249394	27813938	1261.46624232088
NC_014213 Meiothermus silvanus DSM 9946 plasmid pMESIL01.	347854	3461768	1466.53106188228
NC_014214 Meiothermus silvanus DSM 9946 plasmid pMESIL02.	124421	955914	1123.36214947637
NC_014363 Olsenella uli DSM 7084 chromosome.	2051896	7839526	552.985752201866
NC_014364 Spirochaeta smaragdinae DSM 11293 chromosome.	4653970	39431130	1255.9706035922
Natronobacterium gregoryi SP2, DSM 3393 : Natgr_Contig127.1	3788356	8676937	335.230170554193
Natronococcus occultus SP4, DSM 3396 : Natoc_Contig265	12939	243363	2564.16840559549
Natronococcus occultus SP4, DSM 3396 : Natoc_Contig266	287963	819298	396.405625028215
Natronococcus occultus SP4, DSM 3396 : Natoc_Contig267	4013216	11443075	397.25378947956
Pseudomonas stutzeri RCH2 : Psest_Contig40.4	2804	62801	3135.38908701855
Pseudomonas stutzeri RCH2 : Psest_Contig44.3	9865	4321	64.2942726811961
Pseudomonas stutzeri RCH2 : Psest_Contig45.2	12763	54681	630.421844393951
Pseudomonas stutzeri RCH2 : Psest_Contig47.1	4575057	5326365	171.651879965649
Terriglobus roseus KBS 63, DSM 18391 : Terro_Contig51.1	5227858	8464573	239.329327805002
Thermobacillus composti KWC4, DSM 18247 : Theco_Contig54.2	149182	1912248	1900.01226689547
Thermobacillus composti KWC4, DSM 18247 : Theco_Contig56.1	4206343	29042078	1016.11924491179
gi 14286347 gb AE004092.1 Streptococcus pyogenes M1 GAS, complete genome	1852441	1502208	120.443499145182
gi 161501984 ref NC_010067.1 Salmonella enterica subsp. arizonae serovar 62:z4,z23:- str. RSK2980 chromosome	4600800	1831145	58.7192305685968
gi 339998036 ref NC_015761.1 Salmonella bongori NCTC 12419	4460105	501312	16.60908140055
gi 58036263 ref NC_003450.3 Corynebacterium glutamicum ATCC 13032	3309401	1063668	47.6651768703762

Describe methods: mapping of raw reads with bwa mem (Li, 2013) then samtools (Li et al., 2009), input from Esther is needed. Total of 355875608 reads (53381341200bp) sequenced.

Table S3. Performance of error correction

Metric	Program	0	1	2	3	4	6	7	8
Reads	–	9365134	9604918	8811278	8396488	9257066	8609900	8990744	9682468
Perfect	raw	2120932	2179609	1937541	1954244	1872800	2049675	2063874	2183454
	hammer	7656274	8260510	6302861	5970186	6297298	7639715	8068555	8317006
	corsage -c1	8276184	8578469	7861670	7433499	8192894	7736330	8046250	8656282
	corsage -c2	8886436	9188854	8440502	7965810	8829995	8264229	8611867	9272559
	corsage -c3	8965114	9270617	8511335	8024365	8904161	8330927	8683349	9347321
	corsage -c4	8974388	9278745	8511843	8023150	8910290	8338543	8689922	9354172
	corsage -c5	8943102	9243789	8471748	7993790	8875735	8310857	8654540	9312765
Chimeric	raw	69568	67625	81938	75509	79443	52246	44983	59265
	hammer	72820	70590	87564	80336	84813	53875	46387	61948
	corsage -c1	15129	13376	18281	12859	13501	10913	9954	12303
	corsage -c2	5502	4593	10397	4648	5257	3941	3824	4889
	corsage -c3	5224	4245	10181	4349	4802	3642	3555	4614
	corsage -c4	5239	4247	10230	4881	4777	3656	3558	4646
	corsage -c5	5318	4350	10321	4703	4896	3698	3611	4731
Better	raw	–	–	–	–	–	–	–	–
	hammer	6743983	7026478	6156387	5669978	6574627	6244274	6645542	7095951
	corsage -c1	6899557	7131162	6610328	6169861	7102763	6311717	6653538	7200704
	corsage -c2	7008163	7236195	6707136	6260408	7206731	6403639	6749255	7306961
	corsage -c3	7017190	7247251	6715402	6258028	7215037	6412774	6759006	7315526
	corsage -c4	7011329	7242576	6709552	6238573	7211254	6408499	6753669	7310524
	corsage -c5	6987270	7218080	6681337	6211648	7184655	6388594	6729406	7280437
Worse	raw	–	–	–	–	–	–	–	–
	hammer	31644	26951	32315	29096	41584	25270	26959	28960
	corsage -c1	76694	76914	65596	75134	75213	62950	65427	73162
	corsage -c2	25990	25304	20560	27335	27390	20791	21074	24001
	corsage -c3	21353	20867	17696	23556	23768	17299	17238	20241
	corsage -c4	20967	20559	17040	24583	23721	17037	16932	20007
	corsage -c5	21386	21512	16922	25583	24349	17529	17774	20919
Time	hammer corsage								
RAM	hammer corsage								

A read is said to become *better* (or *worse*) if the best alignment of the corrected sequence has more (or fewer) identical bases to the reference genome than the best alignment of the original sequence. The table gives the number of reads mapped *perfectly*, number of *chimeric* reads (i.e. reads with parts mapped to different places), number of corrected reads becoming *better* and the number of corrected reads becoming *worse* than the original reads. Mapping of raw and corrected reads to reference with BWA-MEM-0.7.12 (Li, 2013), postprocessing with samtools-1.2 (Li et al., 2009) and some custom scripts, all available on GitHub eventually.

Table S4. IDBA-UD assembly statistics

Metric	Program	0	1	2	3	4	6	7	8
NG50	raw	45284	53863	31250	24834	17196	87102	80574	59754
	hammer	40924	51004	29256	24023	17246	95532	87102	44292
	corsage	59081	73496	54946	41749	31569	90184	80997	80997
# contigs	raw	330	342	514	607	654	201	200	303
	hammer	335	345	530	626	653	191	194	328
	corsage	277	272	409	497	512	200	198	249
Largest contig	raw	227106	203026	203098	141383	102074	221687	232585	162612
	hammer	203098	157125	197417	141494	107872	221687	178322	139398
	corsage	236473	203098	144213	141579	124628	221683	221683	236473
Total length	raw	4400079	4587934	4400469	4139052	3940625	4640153	4639167	4533515
	hammer	4402128	4591089	4400334	4144742	3934141	4641409	4638005	4538546
	corsage	4408926	4590112	4421966	4171137	3972787	4639453	4636457	4538141
# misassemblies	raw	16	11	15	32	36	0	0	10
	hammer	17	11	9	37	37	0	0	8
	corsage	9	4	2	13	20	0	0	7
# mismatches per 100 kbp	raw	4.17	3.64	10.16	17.02	20.41	0.31	0.13	3.16
	hammer	4.88	3.88	12.50	16.35	20.76	0.15	0.11	2.90
	corsage	4.38	3.80	7.93	8.86	12.82	2.30	2.39	3.27
# indels per 100 kbp	raw	0.32	0.24	0.69	1.47	1.09	0.13	0.09	0.36
	hammer	0.32	0.29	0.79	1.52	1.51	0.11	0.09	0.34
	corsage	0.25	0.31	0.53	0.90	0.85	0.09	0.09	0.18
Genome fraction (%)	raw	93.656	97.182	93.344	87.892	83.101	98.266	98.245	96.104
	hammer	93.631	97.165	93.304	87.924	82.958	98.211	98.175	96.028
	corsage	93.928	97.431	93.988	88.827	84.053	98.271	98.252	96.265
# genes	raw	3873	4049	3741	3477	3220	4186	4191	4027
	hammer	3859	4052	3709	3469	3208	4194	4199	4005
	corsage	3931	4126	3857	3593	3347	4204	4202	4088
NGA50	raw	44890	53863	31250	24224	16220	87102	80574	59589
	hammer	40924	51004	29207	24023	17227	95444	87102	44292
	corsage	59081	73496	54946	41749	31567	90184	80997	80574
Time	raw								
	hammer								
	corsage								
RAM	raw								
	hammer								
	corsage								

IDBA-UD. Explain metric definitions as outlined by QUAST. State, that everything was computed with QUAST (Gurevich et al., 2013), default settings, contigs greater than 500bp, bla bla. Availability in GitHub.

Table S5. SPAdes assembly statistics

Metric	Program	0	1	2	3	4	6	7	8
NG50	raw	65444	95218	66287	48903	26823	114661	117715	86966
	hammer	86625	95218	67436	52817	31448	120770	132608	105995
	cor sage	86625	95517	72055	53947	36236	112350	132608	112853
# contigs	raw	447	400	606	718	813	245	233	324
	hammer	302	275	474	594	676	198	185	250
	cor sage	288	279	418	534	569	210	213	263
Largest contig	raw	203603	224667	218793	178300	113773	269308	268816	223154
	hammer	204882	203257	218793	167410	135551	312119	269348	269318
	cor sage	203394	224320	218793	178231	155221	268535	312008	268327
Total length	raw	4522153	4703061	4533214	4290463	4138591	4713277	4718163	4633297
	hammer	4443696	4633876	4464269	4233011	4046113	4686582	4689565	4584513
	cor sage	4452849	4645907	4471868	4240428	4065827	4698390	4702810	4600454
# misassemblies	raw	15	2	22	45	51	1	3	12
	hammer	11	7	19	31	38	1	2	7
	cor sage	6	3	10	23	22	1	0	6
# mismatches per 100 kbp	raw	15.30	11.57	34.70	48.21	50.53	2.84	2.14	9.72
	hammer	12.70	10.30	30.34	40.41	48.42	1.27	2.17	7.66
	cor sage	10.41	9.32	22.69	30.86	36.47	5.66	5.21	8.43
# indels per 100 kbp	raw	0.89	1.17	2.26	4.48	4.24	0.31	0.22	1.00
	hammer	1.17	1.19	3.16	3.48	4.58	0.24	0.35	0.94
	cor sage	0.64	0.95	2.24	3.30	3.28	0.55	0.31	0.83
Genome fraction (%)	raw	94.241	97.948	94.239	89.098	84.372	98.665	98.708	96.702
	hammer	94.050	97.527	93.984	89.133	84.220	98.505	98.446	96.459
	cor sage	94.223	97.629	94.223	89.543	84.798	98.580	98.541	96.603
# genes	raw	3876	4117	3782	3532	3281	4217	4224	4081
	hammer	3898	4124	3805	3562	3300	4211	4219	4093
	cor sage	3937	4133	3866	3608	3390	4218	4220	4097
NGA50	raw	65444	95218	66287	41757	26823	114661	117715	86966
	hammer	82361	95218	67436	51035	31446	120770	132608	99558
	cor sage	86625	95435	71474	52883	36219	112350	132608	105926
Time	raw								
	hammer								
	cor sage								
RAM	raw								
	hammer								
	cor sage								

SPAdes --sc --only-assembler -k 21,33,55,77 --careful. Explain metric definitions as outlined by QUAST. State, that everything was computed with QUAST, cite it, default settings, contigs greater than 500bp, bla bla. Availability in GitHub.