Sequence analysis

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

Andreas Bremges¹²*, Esther Singer², Tanja Woyke² and Alexander Sczyrba¹²

¹Center for Biotechnology and Faculty of Technology, Bielefeld University, Bielefeld 33615, Germany and
²U.S. Department of Energy Joint Genome Institute, Walnut Creek, CA 94598, USA

*To whom correspondence should be addressed.

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Abstract

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

Availability and implementation: https://github.com/metagenomics/MeCorS

Contact: abremges@cebitec.uni-bielefeld.de

Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

The vast majority of microbial species found in nature has yet to be grown in pure culture, turning metagenomics and—more recently—single cell genomics into indispensable methods to access the genetic makeup of microbial dark matter (Brown et al., 2015; Rinke et al., 2013). Frequently, single amplified genomes (SAGs) and shotgun metagenomes are generated from the same environmental sample, and are methodologically combined e.g. to validate metagenome bins with single cells or to improve the SAG’s assembly contiguity (Campbell et al., 2013; Hess et al., 2011). However, a single cell’s DNA needs to be amplified prior to sequencing, as usually accomplished by multiple displacement amplification (MDA; Lasken, 2007). This amplification is heavily biased, leading to uneven sequencing depth including ultra-low coverage regions with basically no informed error correction possible (Chitsaz et al., 2011; Supplementary Fig. S1). Moreover, chimera formation occurs roughly once per 10 kbp during MDA, further complicating SAG assembly (Nurk et al., 2013; Rodrigue et al., 2009).

While an array of error correction tools exist for a variety of use cases (Laehnemann et al., 2016), only one tool was specifically designed to correct SAG data: hammer (Medvedev et al., 2011), recently refined to BayesHammer (Nikolenko et al., 2013). We propose a metagenome-enabled error correction strategy for single cell sequencing reads. Our method takes advantage of largely unbiased metagenomic coverage, enabling it to correct positions with too low a coverage for SAG-only error correction, and to correct chimeric SAG reads through non-chimeric metagenome reads.

2 Methods

We correct potential errors using an algorithm similar to solving the spectral alignment problem (Pevzner et al., 2001). Given a set of trusted k-mers, we use a heuristic method to find a sequence with minimal corrections such that each k-mer on the corrected sequence is trusted. Using a k-mer size of 31, we consider a k-mer trusted if it occurs at least twice in the accompanying metagenome. This coverage threshold was determined empirically to work with most datasets (Supplementary Fig. S2).

Our correction algorithm was inspired by fermi (Li, 2012) and BFC (Li, 2015), but we do not act on the assumption of uniform
sequencing coverage, thereby accounting for the tremendous variation of coverage across the SAG. Instead, we exploit metagenomic sequence information to correct errors resulting from amplification and sequencing, as well as chimeras, even in ultra-low coverage regions of the SAG. The non-chimeric nature of the metagenome reads enables an implicit and thorough write-through correction of chimeric SAG reads.

MeCorS works in three phases:

1. MeCorS collects all 31-mers (and their reverse complements) occurring in the SAG reads. It uses this information to initialize a hash table with the 31-mers being valid keys.
2. MeCorS scans the accompanying metagenomic reads. For each stored 31-mer, it counts the occurrence of the next (i.e. the 32nd) base in the metagenome and stores the totals in the hash table. This step is largely I/O bound and dominates MeCorS’s runtime.
3. MeCorS processes each SAG read by using the 31-mer hash table to check if the 32nd base is sufficiently supported in the metagenome. Untrusted 32nd bases are replaced with the most frequent and trusted 32nd bases from the metagenome.

3 Results and discussion

As a realistic benchmark, we used eight Escherichia coli K12-MG1655 SAGs from Clingenpeel et al. (2014), a strain for which the complete genome sequence is available (Supplementary Table S1). A concomitant in vitro mock metagenome consisting of 26 microbial species, including E. coli K12-MG1655, was sequenced on Illumina’s HiSeq platform (Bowers et al., 2015). Based on metagenome read mapping, we estimate the relative abundance of E. coli to amount to 0.15%, corresponding to a mean per-base coverage of only 20.7x (Supplementary Table S2).

We evaluated MeCorS along with BayesHammer (Nikolenko et al., 2013), a widely used error correction tool for SAG data. Our method corrects more errors than BayesHammer, producing a significantly higher fraction of better and perfect reads after correction (Table 1; Supplementary Table S3). In contrast to BayesHammer, MeCorS reduces the amount of chimeric SAG reads by one order of magnitude, likely due to the non-chimeric nature of the metagenome reads. MeCorS works well with modern single cell assemblers, most notably reducing the misassembly rate of both IDBA-UD (Peng et al., 2012) and SPAdes (Bankevich et al., 2012) by half, while providing high sequence contiguity (Fig. 1). In particular poorly amplified SAGs benefit from metagenome-enabled error correction, yielding improved assembly accuracy and contiguity (Supplementary Tables S4 and S5).

We note that such a hybrid error correction of SAG data may result in miscorrection(s) of rare variants. If the captured cell contains a variant that is rare or absent in the corresponding metagenome, correction will be biased towards the most abundant variant in the metagenome sequence. If strain resolution is desired, we suggest pollya variant that is rare or absent in the corresponding metagenome, correction will be biased towards the most abundant variant in the metagenome sequence. If strain resolution is desired, we suggest pooling sequencing errors and their correction.

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Conflict of Interest: none declared.

References

Bankewich, A. et al. (2012) SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J. Comput. Biol., 19, 455–477.
Bowers, R.M. et al. (2015) Impact of library preparation protocols and template quantity on the metagenomic reconstruction of a mock microbial community. BMC Genomics, 16, 856.
Brown, C.T. et al. (2015) Unusual biology across a group comprising more than 15% of domain Bacteria. Nature, 523, 208–211.
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., 110, 5540–5545.
Chitsaz, H. et al. (2011) Efficient de novo assembly of single-cell bacterial genomes from short-read data sets. Nat. Biotechnol., 29, 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, 1072–1075.
Hess, M. et al. (2011) Metagenomic discovery of biomass-degrading genes and genomes from cow rumen. Science, 331, 463–467.
Laehnemann, D. et al. (2016) Denoising DNA deep sequencing data—high-throughput sequencing errors and their correction. Brief. Bioinf., 17, 154–179.

Table 1. Performance of SAG error correction

| Program   | % perfect | % chimeric | % better | % worse |
|-----------|-----------|------------|----------|---------|
| Raw       | 22.52 ± 1.07 | 0.73 ± 0.15 | –        | –       |
| BayesHammer | 80.35 ± 8.77 | 0.77 ± 0.17 | 71.66 ± 2.12 | 0.33 ± 0.06 |
| MeCorS    | 95.52 ± 0.43 | 0.06 ± 0.02 | 75.45 ± 1.11 | 0.26 ± 0.03 |

Mean percentage and standard deviation of perfect reads, chimeric reads (i.e. reads with parts mapped to different places), corrected reads becoming better and worse than the raw reads. Evaluation as described in Li (2015); please refer to Supplementary Table S3 for per-SAG metrics, including runtime and memory usage.
Lasken, R. S. (2007) Single-cell genomic sequencing using Multiple Displacement Amplification. *Curr. Opin. Microbiol.*, 10, 510–516.

Li, H. (2012) Exploring single-sample SNP and INDEL calling with whole-genome de novo assembly. *Bioinformatics*, 28, 1838–1844.

Li, H. (2015) BFC: correcting illumina sequencing errors. *Bioinformatics*, 31, 2885–2887.

Medvedev, P. et al. (2011) Error correction of high-throughput sequencing datasets with non-uniform coverage. *Bioinformatics*, 27, i137–i141.

Nikolenko, S. I. et al. (2013) BayesHammer: Bayesian clustering for error correction in single-cell sequencing. *BMC Genomics*, 14, S7.

Nurk, S. et al. (2013) Assembling single-cell genomes and mini-metagenomes from chimeric MDA products. *J. Comput. Biol.*, 20, 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, 1420–1428.

Pevzner, P. A. et al. (2001) An Eulerian path approach to DNA fragment assembly. *Proc. Natl. Acad. Sci. U. S. A.*, 98, 9748–9753.

Rinke, C. et al. (2013) Insights into the phylogeny and coding potential of microbial dark matter. *Nature*, 499, 431–437.

Rodrique, S. et al. (2009) Whole genome amplification and de novo assembly of single bacterial cells. *PLoS One*, 4, e6864.