

Genome analysis

Binning_refiner: improving genome bins through the combination of different binning programs

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Abstract

Summary: Microbial genomes have recently been reconstructed from metagenomic datasets using binning approaches. Inconsistent binning results are however often observed between different binning programs, likely due to the different algorithms or statistical models used. We present Binning_refiner, a pipeline that merges the results of different binning programs. Our results demonstrated that Binning_refiner can significantly reduce the contamination level of genome bins and increase the total size of contamination-free and 'good-quality' genome bins. Binning_refiner is thus an useful tool to improve the quality of genome bins derived from metagenomic data.

Availability and Implementation: Binning_refiner is implemented in Python3 and is freely available at: https://github.com/songweizhi/Binning_refiner.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

High-throughput shotgun sequencing provides a powerful way to study the 'unexplored' and uncultured diversity of microbial communities (Eloe-Fadrosh et al., 2016). A number of approaches have been recently developed to reconstruct complete or partial microbial genomes from metagenomics shotgun sequences, a process called genome binning (Alneberg et al., 2014; Imelfort et al., 2014; Kang et al., 2015; Lin and Liao, 2016). These approaches cluster contigs assembled from metagenomics sequences based on compositional properties (e.g. GC content, tetra-nucleotide frequency) or sequence coverage profiles across multiple samples (or combinations thereof). Inconsistent results have however often been observed between binning approaches, which is likely due to the differences in algorithm or statistical models employed (Kang et al., 2015; Lin and Liao, 2016). It is important that all sequences of a bin are specific to a given organism (i.e. bins are free of contamination from any other organism) as this would otherwise lead to erroneous functional and metabolic inferences (Albertsen et al., 2013; Rinke et al., 2013). To assess this, a number of algorithms have been established that measure the level of contamination and genome completeness using sets of marker sequences (Parks *et al.*, 2015; Sangwan *et al.*, 2016; Seah and Gruber-Vodicka, 2015).

Here we have developed a pipeline called Binning_refiner that reconciles the outputs of different binning programs with the aim to improve the quality of genome bins, in particular with respect to contamination levels.

2 Materials and methods

The main steps of Binning_refiner are summarized in Figure 1A. The output bins of two binning programs are used as inputs for Binning_refiner. In the first step, the bin names are added to contig names. The bins from each binning program are then combined into one file and a pairwise blastn (Altschul *et al.*, 1990) is performed between the two combined files. The blastn results are filtered and only matches passing the user-defined criteria are kept (here we used query length = subject length = alignment length and identity = 100%). Each set of shared contigs between the two sets of

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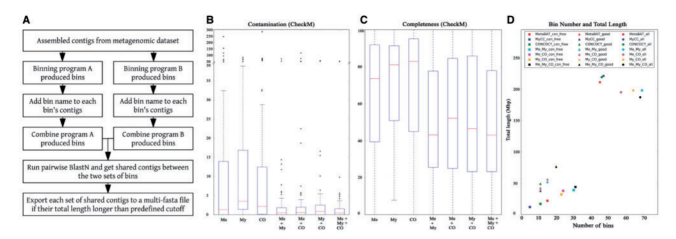


Fig. 1 (A) Binning_refiner workflow (B–D). Qualities of genome bins from a real metagenomic dataset. The boxes in the whisker plot are bound by the lower to upper quartile values with the red line being the median. Whiskers represent the range of the corresponding data. 'con_free' refers to 'contamination-free bins', 'good' refers to bins with a completeness higher than 70% and contamination lower than 5%. 'all' refers to all bins produced. 'Me', 'My' and 'CO' refer to 'MetaBAT', 'MyCC' and 'CONCOCT', respectively. The black and yellow triangles in panel D are overlapped

bins are treated as a refined bin (see Supplementary Fig. S1) and exported in multi-fasta format, if their total length is longer than a defined cutoff (e.g. 0.5 Mbp). CheckM (Parks *et al.*, 2015) is used after Binning_refiner to assess the quality (contamination and completeness) of input and refined bins.

3 Results and discussion

Binning_refiner's performance was first assessed on the MBARC-26 mock dataset, which consists of shotgun sequences for a defined mixture of 23 bacterial and three archaeal strains with publicly available complete genomes (Singer et al., 2016). Sequence assembly using IDBA-UD (Peng et al., 2012) generated 92.4 Mbp of contiguous sequences ≥5 Kbp, of which 52.3 Mbp were assigned to 24 complete genomes with an identity cutoff of 99% (contigs assigned to more than one genomes were excluded). These contigs unique genome assignments were used as input for the binning programs MetaBAT (Kang et al., 2015), MyCC (Lin and Liao, 2016) and CONCOCT (Alneberg et al., 2014) using default parameters. Both MetaBAT and MyCC resulted in 23 bins ≥ 0.5 Mbp, while CONCOCT gave 13 bins ≥ 0.5 Mbp (Table 1). All bins produced by the three programs were merged pairwise with Binning_refiner and also sequential by merging two outputs and combining this with the bins from the third program. Given the availability of reference genomes for this mock dataset, overall precision (defined as how pure a bin is) and recall (defined as how complete a bin is) for each set of bins were calculated with Evaluate.py from the MyCC package (Lin and Liao, 2016). The results show that the performance of CONCOCT was much worse than MetaBAT and MyCC in terms of precision and bin number on this mock dataset. As a consequence, merging CONCOCT bins with bins from the other two programs did not result in improved refined bins (Table 1). However, a substantial improvement was observed by merging MetaBAT and MyCC bins (Supplementary Fig. S1). The precision was increased from 87% (MetaBAT) and 91% (MyCC) to 96% (Binning_refiner), while the recall was decreased from 91% (MetaBAT) and 94% (MyCC) to 88% (Binning_refiner) (Table 1, Supplementary Fig. S2).

Binning_refiner was also tested on metagenomic sequence data for three replicate bacterial communities from the surface of the marine alga *Caulerpa filiformis* (Roth-Schulze *et al.*, 2016). The

Table 1. Performance of Binning_refiner on the MBARC-26 mock dataset

	Me	My	СО	Me +	Me +	My +	Me +
				My	CO	CO	My + CO
Bin number	23	23	13	26	23	24	26
Precision (%)	87	91	60	96	87	93	96
Recall (%)	91	94	99	88	91	94	87

'Me', 'My' and 'CO' refer to 'MetaBAT', 'MyCC' and 'CONCOCT', respectively.

paired-end sequence data was generated on an Illumina HiSeq2000 platform and contains a total of 97.94 Gbp. Sequence assembly with IDBA-UD generated 237 Mbp of contiguous sequences ≥ 5 Kbp. MetaBAT, MyCC and CONCOCT gave 45, 47 and 46 bins ≥ 0.5 Mbp, respectively. Qualities of bins given by the three programs as well as refined bins generated by different combinations of merging were given in Figure 1(B-D). The contamination levels of the four combination of refined bin sets are much lower than the three input bin sets (Fig. 1B), but completeness was also reduced (Fig. 1C). We next looked at contamination-free and 'good-quality' bins, which are defined with completeness higher than 70% and contamination less than 5%. The best outcomes for both contamination-free bins and 'good-quality' bins were obtained by merging all three input bin sets (Supplementary Fig. S3). The total length of the refined contamination-free bins is 44 Mbp, which is 2 times higher than that of the MetaBAT bins (22 Mbp), 3.7 times higher than that of the MyCC bins (12 Mbp) and 2.6 times higher than that of the CONCOCT bins (17 Mbp) (Fig. 1D). Likewise, the total length of the 'good-quality' refined bins is 76 Mbp, which is 1.8 times higher than that of the MetaBAT bins (42 Mbp), 2 times higher than that of the MyCC bins (38 Mbp) and 1.6 times higher than that of the CONCOCT bins (49 Mbp) (Fig. 1D). The results of pairwise merging also show improvement compared to the input bins (Fig. 1D).

Our results demonstrate that Binning_refiner can significantly reduce contamination levels and increase the precisions of genome bins, which resulted in an improved total length of 'good-quality' bins and contamination-free bins.

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Conflict of Interest: none declared.

References

- Albertsen, M. et al. (2013) Genome sequences of rare, uncultured bacteria obtained by differential coverage binning of multiple metagenomes. Nat. Biotechnol., 31, 533–538.
- Altschul, SF. et al. (1990) Basic local alignment search tool. J. Mol. Biol., 215, 403–410.
- Alneberg, J. et al. (2014) Binning metagenomic contigs by coverage and composition. *Nat. Methods*, 11, 1144–1146.
- Eloe-Fadrosh, E.A. et al. (2016) Metagenomics uncovers gaps in amplicon-based detection of microbial diversity. Nat. Microbiol., 1, 15032.
- Imelfort, M. et al. (2014) GroopM: an automated tool for the recovery of population genomes from related metagenomes. PeerJ, 2, e603.

- Kang, D.D. et al. (2015) MetaBAT, an efficient tool for accurately reconstructing single genomes from complex microbial communities. PeerJ, 3, e1165.
- Lin,H.H. and Liao,Y.C. (2016) Accurate binning of metagenomic contigs via automated clustering sequences using information of genomic signatures and marker genes. Sci. Rep., 6, 24175.
- Parks,D.H. et al. (2015) CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. Genome Res., 25, 1043–1055.
- 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.
- Rinke, C. et al. (2013) Insights into the phylogeny and coding potential of microbial dark matter. Nature, 499, 431–437.
- Roth-Schulze, A.J. et al. (2016) Partitioning of functional and taxonomic diversity in surface-associated microbial communities. Environ. Microbiol., 18, 4391–4402.
- Sangwan, N. et al. (2016) Recovering complete and draft population genomes from metagenome datasets. Microbiome, 4, 1.
- Seah, B.K. and Gruber-Vodicka, H.R. (2015) gbtools: interactive visualization of metagenome bins in R. Front. Microbiol., 6, 1451.
- Singer, E. et al. (2016) Next generation sequencing data of a defined microbial mock community. Sci. Data, 3, 160081.