

Genome analysis

SolidBin: improving metagenome binning with semi-supervised normalized cut

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Abstract

Motivation: Metagenomic contig binning is an important computational problem in metagenomic research, which aims to cluster contigs from the same genome into the same group. Unlike classical clustering problem, contig binning can utilize known relationships among some of the contigs or the taxonomic identity of some contigs. However, the current state-of-the-art contig binning methods do not make full use of the additional biological information except the coverage and sequence composition of the contigs.

Results: We developed a novel contig binning method, Semi-supervised Spectral Normalized Cut for Binning (SolidBin), based on semi-supervised spectral clustering. Using sequence feature similarity and/or additional biological information, such as the reliable taxonomy assignments of some contigs, SolidBin constructs two types of prior information: must-link and cannot-link constraints. Must-link constraints mean that the pair of contigs should be clustered into the same group, while cannot-link constraints mean that the pair of contigs should be clustered in different groups. These constraints are then integrated into a classical spectral clustering approach, normalized cut, for improved contig binning. The performance of SolidBin is compared with five state-of-the-art genome binners, CONCOCT, COCACOLA, MaxBin, MetaBAT and BMC3C on five next-generation sequencing benchmark datasets including simulated multi- and single-sample datasets and real multi-sample datasets. The experimental results show that, SolidBin has achieved the best performance in terms of *F*-score, Adjusted Rand Index and Normalized Mutual Information, especially while using the real datasets and the single-sample dataset.

Availability and implementation: <https://github.com/sufforest/SolidBin>.

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Supplementary information: [Supplementary data](#) are available at *Bioinformatics* online.

1 Introduction

Microbial communities consist of mixtures of large number of microorganisms. In metagenomic studies, genetic material is directly

extracted from the microbial communities and is sequenced using high-throughput sequencing technologies resulting in large quantity of sequencing reads of different read lengths depending on the

sequencing technologies and platforms. During sequencing, the information on what reads come from the same genome or closely related genomes is lost. Usually the first step in metagenomic data analysis is ‘reads assembly’ aiming to assemble the reads into relative longer genomic fragments called *contigs* based on the overlapping information of the reads. The second step is ‘contig binning’ aiming to cluster the contigs from the same genome or closely related genomes into the same bin. In the third step, reads are mapped to known genomic sequences or the contigs in the bins and the relative abundance profiles of genomes and genomic bins are estimated in each sample. Finally, statistical methods are used to associate the abundance profiles of genomes/bins with the environmental factors or phenotypes of the individuals (Bahram *et al.*, 2018; Huttenhower *et al.*, 2012; Qin *et al.*, 2010; Sunagawa *et al.*, 2015). Metagenomic studies have shown that microbial communities in human gut have been shown to be significantly associated with several diseases (Jostins *et al.*, 2012; Khor *et al.*, 2011) and the effect of immunotherapy (Chen *et al.*, 2017; Wilck *et al.*, 2017). In this paper, we concentrate on the usual second step of metagenomic data analysis, contig binning.

The available contig binning methods can be divided into two major categories: taxonomy dependent methods (also called supervised methods) that map the contigs to known reference databases, such as MEGAN (Huson *et al.*, 2007), and taxonomy independent methods (also called unsupervised methods), such as MaxBin (Wu *et al.*, 2014, 2016), CONCOCT (Alneberg *et al.*, 2014), MetaBAT (Kang *et al.*, 2015), COCACOLA (Lu *et al.*, 2017a) and BMC3C (Yu *et al.*, 2018) that cluster the contigs ‘*de novo*’ without mapping the contigs to known genomes. Both categories of methods have their own limitations. Since the known genomes are limited and a large number of genomes are still unknown, many contigs cannot be mapped to known genomes. In addition, alignment is computationally expensive and thus alignment based contig binning methods are generally slow. Without relying on reference genomes, unsupervised binning methods can partially overcome some of the above-mentioned shortcomings. On the other hand, existing unsupervised binning methods have not made full use of additional available biological information such as the alignment of some the contigs to known genomes, which can improve the binning performance. Therefore, it is imperative to develop effective contig binning approaches that can not only be used for the datasets that contain a large amount of contigs from unknown genomes, but also make use of the available biological information from alignment for better binning performance.

To tackle this issue, we develop a novel contig binning method, Semi-supervised Spectral Normalized Cut for Binning (SolidBin), which is inspired by various studies in semi-supervised clustering (Gu *et al.*, 2013; Wagstaff *et al.*, 2001) that can improve the clustering performance by incorporating pairwise constraints. There are two types of pairwise constraints: must-link (ML) constraints and cannot-link (CL) constraints. Specifically, SolidBin is based on semi-supervised spectral clustering. Compared with other clustering methods, spectral clustering algorithm can relatively easily incorporate additional information on pairwise relationships of contigs. SolidBin mainly utilizes two kinds of sequence features: contig coverage information and tetranucleotides frequencies. The pairwise constraints can be generated from both internal (contig features) and external information (additional biological information such as co-alignment). With respect to different kinds of information used for generating constraints, SolidBin has several different modes, SolidBin-sequence feature similarity (SFS), SolidBin-coalign, SolidBin-CL and SolidBin-SFS-CL. SolidBin-SFS generates ML

constraints between contig pairs with high similarity based on internal information of sequence features. On the other hand, SolidBin-coalign and SolidBin-CL generate constraints using external co-alignment information. By aligning contigs to known genomes, TAXAassign (<https://github.com/umerijaz/taxaassign>) can assign some contigs belonging to the known genomes to the corresponding genomes/species. The contig pairs with the same species assignment by TAXAassign have a ML constraint in SolidBin-coalign, while the contig pairs with different genus assignment have a CL constraint in SolidBin-CL. SolidBin-SFS-CL removes the CL pairs from the constraints set on the basis of SolidBin-SFS. The performance of SolidBin has been extensively investigated on five next-generation sequencing benchmark datasets including simulated multi- and single-sample datasets, as well as real multi-sample datasets. Compared with five state-of-the-art genome binner, CONCOCT, COCACOLA, MaxBin, MetaBAT and BMC3C, SolidBin has achieved the best performance in terms of *F*-score, Adjusted Rand Index (ARI) and Normalized Mutual Information (NMI), especially while using the real datasets and the single-sample dataset. Specifically, SolidBin-SFS achieves better performance when there are sufficient samples. SolidBin-coalign also works very well except the case that the dataset contains many genomes on the strain level.

2 Related work

Unsupervised contig binning methods can be roughly divided into three categories based on the sequence characteristics used for binning: (i) Nucleotide Composition-based, (ii) Differential Abundance-based and (iii) both Nucleotide Composition and Abundance-based (Sangwan *et al.*, 2016). Nucleotide Composition-based methods (Dick *et al.*, 2009; Laczny *et al.*, 2015) take oligonucleotide frequency as features, based on the assumption that the oligonucleotide composition of fragments from the same genome are much more similar than fragments from different genomes. Differential Abundance-based methods [Abundancebin (Wu and Ye, 2011)] cluster contigs mainly according to the differential coverage of average nucleotide coverage of each contig, based on the assumption that the contigs that belong to the same genome should have similar abundance in the same sample. Nucleotide Composition and Abundance-based methods (MaxBin, MetaBAT, CONCOCT and COCACOLA) combine the two kinds of information and have been shown to outperform methods using only one kind of information (Sedlar *et al.*, 2017).

The MaxBin algorithm utilizes the single-copy marker genes to determine the number of bins and expectation-maximization algorithm is performed while binning. MaxBin 2.0 can be applied to the datasets that contain multiple samples based on the assumption that all samples are independently sequenced. The MetaBAT algorithm utilizes two different genomic features to calculate the distance of contig pairs, and then converts the distance to the probability of them being in the same genome. Finally, the modified *K*-medoids algorithm is used for binning. CONCOCT takes sequence composition and coverage across multiple samples as sequence feature vectors, and then uses principal component analysis to reduce the dimension of the feature vectors. Finally, it obtains the binning results using the Gaussian mixture model. It determines the number of bins by the variational Bayesian approach. COCACOLA employs the similar sequence feature vectors as CONCOCT and it takes advantage of both the soft and hard clustering by employing non-negative matrix factorization with sparse regularization. The performances

of COCACOLA and CONCOCT depend largely on the number of samples, partially due to the way to construct feature vectors.

Several investigators realized the importance of incorporating additional biological information for binning, such as co-alignment and pair-end read linkage. However, the co-alignment information was not considered by CONCOCT, MaxBin and MetaBAT, while the pair-end reads linkage information was not considered in MaxBin and MetaBAT. CONCOCT utilizes the pair-end reads linkage information only in the post-processing stage of binning. It merges the clusters with sufficient pair-end links and similar coverage to perform a hierarchical clustering (<https://concoct.readthedocs.io/en/latest/>), which may improve the recall of the binning result but reduce the precision. On the other hand, COCACOLA uses non-negative matrix factorization to factorize the feature matrix of contigs, and integrates co-alignment information and paired-end read linkage across multiple samples into a cost function. It was shown that the additional information can improve the binning performance, especially when the number of samples is small. Hetero-RP (Lu et al., 2017b) automatically adjusts the weights of contig features according to the additional co-alignment information before using other binning methods, and the method is integrated into the COCACOLA-python version. However, the performance of Hetero-RP depends heavily on the accuracy of the additional information and the fraction of contig pairs having such information. These studies have inspired us to develop a new binning method with a semi-supervised spectral clustering approach, SolidBin, where pairwise constraints among contigs can be effectively integrated into the spectral clustering framework of contig graph partition.

Recently, several ensemble clustering methods have been developed for metagenomic contig binning. BMC3C (Yu et al., 2018) takes sequence composition, coverage information and codon usage as sequence feature vectors and repeatedly applying the *K*-means clustering with different initialization to form a weight graph of contigs for partitioning. Additionally, ensemble models combining results from multiple binning algorithms have been proposed. Binning refiner (Song and Thomas, 2017) performs a pairwise comparison of contigs between the output bins of two binners using blastn to obtain shared contigs between two sets of bins. Each set of shared contigs is treated as a refined bin if its total length is longer than a predefined threshold. DAS Tool (Sieber et al., 2018) proposes a scoring function based on single-copy genes and uses it to rank bins generated by different binners. It picks up high-scoring bins iteratively and updates the scores of the remaining candidate bins. MetaWRAP (Uritskiy et al., 2018) uses the output bin sets of MetaBAT2, MaxBin2 and CONCOCT to generate hybrid bin sets, in which every pair of contigs in different bins from any original sets are separated. MetaWRAP then selects the best bin based on the estimated completion and contamination scores. In contrast to these ensemble approaches, SolidBin is a stand-alone method that can be integrated into these ensemble approaches as a component for better binning performance.

3 Materials and methods

3.1 Semi-supervised clustering and spectral clustering

Semi-supervised clustering methods can use prior information to improve clustering performance. One important type of prior information is pairwise constraints that can be divided into two categories: ML constraints and CL constraints. A ML constraint means that two instances should be grouped in the same cluster, while a CL constraint indicates that two instances should appear in different

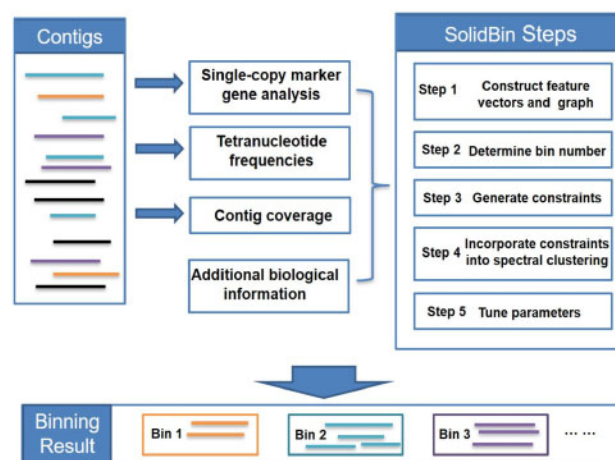


Fig. 1. The general workflow of SolidBin

clusters. For the contig binning problem, instances correspond to contigs and affinity between instances correspond to similarity between contigs.

Spectral clustering methods are based on the spectral graph theory (Ng et al., 2002). A given dataset can be mapped to a graph whose nodes represent the instances of the dataset and the edges reflect the affinity of the instances. The essence of spectral clustering methods is to transform the clustering problem into an optimal partition problem of the graph. Spectral clustering has several variants according to different criteria for graph partitioning and the normalized cut (Ncut) criterion is a reliable one that measures both the total inter-cluster dissimilarity and the total intra-cluster similarity (Shi and Malik, 2000). Compared with other clustering methods, the combination of spectral clustering algorithm and the semi-supervised method has obvious advantages. Spectral clustering algorithm utilizes the similarity among instances, and thus the pairwise constraints among instances can be easily incorporated into the algorithm to improve clustering performance.

3.2 The SolidBin algorithm

In this paper, we developed a contig binning method, SolidBin, based on the semi-supervised spectral clustering method that can utilize additional biological information and make full use of both sequence composition and coverage information. The method has five major steps. First, the contigs are mapped into a graph after obtaining the feature representations of contigs. Then, the bin number *K* is initialized according to the single-copy marker genes, and the final bin number will be determined by the iterative binning results of *K*-means. Next, constraints are generated differently according to the modes of SolidBin, and then the constraints will be used for semi-supervised Ncut with the final bin number. Finally, the method will evaluate the clustering performance with different parameter values without labels and keep the optimal one as the final result. The general pipeline of SolidBin is shown in Figure 1 and more details of the steps are as follows.

3.2.1 Construct feature vectors and graph

Similar to CONCOCT and COCACOLA, we employ the combination of an *M* dimensional coverage vector and a *T* dimensional composition vector to represent each contig, where *M* is the number of the samples and *T* is the number of distinct tetramers. *T*=136 in our method considering one tetramer and its reverse-complement

are combined. After adding a pseudo-count, the composition vectors are normalized over contigs to account for different contig lengths. The coverage vectors are normalized over contigs and over samples, so that different read numbers from a sample are accounted for Alneberg et al. (2014). Therefore, the feature matrix of the contigs is denoted as $X \in \mathbb{R}^{N \times (M+T)}$, where N denotes the number of contigs.

For graph construction, we use L1-distance to measure the dissimilarity between each pair of contigs (Lu et al., 2017a) and convert it to the affinity using the following formula, where A denotes the affinity matrix, Dist denotes the dissimilarity matrix of the contigs and $\max(\cdot)$ and $\min(\cdot)$ denotes the maximum and minimum of all the elements of the matrix, respectively.

$$A_{ij} = 1 - \frac{\text{Dist}_{ij} - \min(\text{Dist})}{\max(\text{Dist}) - \min(\text{Dist})}. \quad (1)$$

3.2.2 Bin number determination

Similar to the method used in COCACOLA, we utilize single-copy marker genes to estimate the initial bin number k_0 , and then, we try a list of numbers larger than k_0 as the bin numbers sequentially and use the K -means algorithm for binning. Then we calculate the silhouette coefficient (Rousseeuw, 1987) of the binning result, which is an index to evaluate the clustering performance without labels by measuring the cohesion and the separation of the clusters. Our methods will find two local maxima of the silhouette coefficient values, and the larger one corresponds to a bin number, which is our final bin number.

3.2.3 Generate constraints

• Generate ML constraints

ML constraints are mainly generated in three ways as three modes of SolidBin:

- i. SolidBin-SFS mode: constraints are generated according to SFS, which is a taxonomy-independent method. In this mode, we underline the reliable information from sequence features themselves without using reference genomes. The contig pairs with high similarity are chosen as ML constraints. Because the affinity graphs of different datasets have different connection structures, the ratio of the pairwise constraints we selected from whole similarity matrix may vary widely. Therefore, we determine the ratio of the pairwise constraints adaptively as follows. We assume that the connection relationship obtained from the binning results is close to the real connection relationship and varies among different datasets. The assumption will be examined in the experiments. We run the NCut method and obtain the initial binning results for the connection graphs. The connection ratio is calculated as follows, where K represents the number of the bins used by NCut method and Num_k represents the number of contigs contained in the k -th bin.

$$\text{linkSum} = 2 \times \sum_{k=1}^K \binom{\text{Num}_k}{2} \quad (2)$$

$$\text{ratio} = \frac{\text{linkSum}}{N \times (N-1)} \times 100\%. \quad (3)$$

- ii. SolidBin-coalign mode: we take the contig pairs with the same assignment on species level using NCBI taxonomy by TAXAassign as ML constraints. In this mode, we take advantage of some contigs that belong to the known reference genomes although the reference genomes are incomplete.

- iii. SolidBin-SFS-CL: constraints are first generated according to the SFS as in SolidBin-SFS. We next remove the contig pairs assigned to different genera by TAXAassign.

• Generate CL constraints

SolidBin-CL: to obtain CL constraints with high accuracy, the contigs assigned to different genus by TAXAassign are regarded as CL pairs.

3.2.4 Incorporate constraints into spectral clustering

• Spectral clustering with NCut

A popular criterion in spectral clustering is the NCut, for which the cost function of given K clusters $\{P_1, \dots, P_K\}$ is defined as follows (Gu et al., 2013; Ji et al., 2006):

$$F_{\text{Ncut}} = \sum_{k=1}^K \frac{\text{Cut}(P_k, \bar{P}_k)}{\text{Cut}(P_k, V)} \quad (4)$$

where V denotes the vertex set, $\text{Cut}(P_k, \bar{P}_k) = \sum_{i \in P_k, j \in \bar{P}_k} W_{ij}$ and W denotes the affinity matrix. Let $H_i = [h_{1i}, h_{2i}, \dots, h_{Ni}]^T$ be the binary indicator vector of cluster P_i , and $h_{ni} = 1$ means the n -th contig belongs to P_i . Let D be a diagonal matrix with $d_{nn} = \sum_{i=1}^N W_{ni}$.

F_{Ncut} can be rewritten as follows:

$$\begin{aligned} F_{\text{Ncut}} &= \sum_{i=1}^K \frac{H_i^T (D - W) H_i}{H_i^T D H_i} = K - \sum_{i=1}^K \frac{H_i^T W H_i}{H_i^T D H_i} \\ &= K - \sum_{i=1}^K \frac{H_i^T D^{\frac{1}{2}} D^{-\frac{1}{2}} W D^{-\frac{1}{2}} D^{\frac{1}{2}} H_i}{H_i^T D^{\frac{1}{2}} D^{\frac{1}{2}} H_i} \\ &= K - \sum_{i=1}^K Y_i^T D^{-\frac{1}{2}} W D^{-\frac{1}{2}} Y_i \end{aligned} \quad (5)$$

where $Y = [Y_1, Y_2, \dots, Y_K]$, $Y_i = D^{\frac{1}{2}} H_i / \|D^{\frac{1}{2}} H_i\|$, and $Y^T Y = I$.

$$F_{\text{Ncut}} \geq K - (\lambda_1 + \lambda_2 + \dots + \lambda_K) \quad (6)$$

where $\lambda_1, \dots, \lambda_K$ are the largest K eigenvalues of matrix $D^{-\frac{1}{2}} W D^{-\frac{1}{2}}$, and when Y_1, Y_2, \dots, Y_K are their corresponding eigenvectors, the equality sign of in Equation (6) holds.

• Integrate ML constraints

If the i -th contig and the j -th contig belong to the same species, the i -th and j -th row of the indicator matrix H should be the same. Therefore, If there are C ML constraints, the c -th ML constraints can be represented as a constraint row vector $U_c = [u_{1c}, u_{2c}, \dots, u_{Nc}]$, where $u_{ic} = 1$, $u_{jc} = -1$, and the rest of all elements are zero. We can encode all ML constraints by matrix $U_C = [U_1, U_2, \dots, U_C]^T$ such that:

$$\begin{aligned} UH = 0 &\Rightarrow \|UH\|^2 = 0 \Rightarrow \text{Tr}(H^T U^T UH) = 0 \\ &\Rightarrow \text{Tr}(Y^T D^{-\frac{1}{2}} U^T U D^{-\frac{1}{2}} Y) = 0. \end{aligned} \quad (7)$$

• Integrate CL constraints

If the i -th contig and the j -th contig belong to different genomes, the i -th and j -th rows of the indicator matrix H should be orthogonal to each other. So, we can use an $N \times N$ non-negative matrix to represent the CL information, where $Z_{ij} = 1$ means that the i -th contig and the j -th contig belong to different genomes, while $Z_{ij} = 0$ means that the relationship between the i -th contig and the j -th contig is unknown. The CL constraints can be represented as follows:

Table 1. Summary of different SolidBin modes

SolidBin mode	Constraints	Type	Parameter	Description	Performance profiles
SolidBin-naive	None	Taxonomy-independent	None	The NCut mode	Comparable performance to other binners
SolidBin-SFS	ML	Taxonomy-independent	α	Take the contig pairs with high similarity as ML constraints	Good performance when the number of samples is large. It can obtain good results with high similarity quality, and more samples could bring more useful coverage information
SolidBin-coalign	ML	Taxonomy-dependent	α	Take the contig pairs with the same assignment by TAXAassign as ML constraints	Good performance when the genomes contained in the datasets are on the species level, while using the results from TAXAassign. It can obtain good results if the pairwise ML constraints have high accuracy
SolidBin-CL	CL	Taxonomy-dependent	β	Take the contig pairs assigned to different genera by TAXAassign as CL constraints	Good performance when the genomes contained in the datasets are on the species level, often worse than SolidBin-coalign, while using the results from TAXAassign. It can obtain good results if the pairwise CL constraints have high accuracy
SolidBin-SFS-CL	ML	Taxonomy-dependent	α	Remove the contig pairs assigned to different genera by TAXAassign from the constraints set on the basis of SolidBin-SFS	Often a little better than SolidBin-SFS at a cost of spending much time on taxonomy alignment

Table 2. Datasets used in the experiments

Datasets	Data type	Sample num	Contig num	Labeled contig num
SpeciesMock	Simulated	96	37 628	37 628
StrainMock	Simulated	64	9417	9417
SimHC	Simulated	1	29 535	13 918
Sharon	Real	18	5579	2614
MetaHIT	Real	264	192 673	17 136

$$\text{Tr}(H^T ZH) = 0. \quad (8)$$

ML and CL constraints can be incorporated into the framework of the NCut as follows (Gu *et al.*, 2013):

$$\begin{aligned}
 F_{\text{SolidBin}} &= F_{\text{Ncut}} + \alpha \|UH\|^2 + \beta \text{Tr}(H^T ZH) \\
 &= K - \sum_{i=1}^K Y_i^T D^{-\frac{1}{2}} W D^{-\frac{1}{2}} Y_i \\
 &\quad + \alpha \text{Tr}(Y^T D^{-\frac{1}{2}} W D^{-\frac{1}{2}} Y) + \beta \text{Tr}(Y^T D^{-\frac{1}{2}} Z D^{-\frac{1}{2}} Y) \\
 &= K - \text{Tr}(Y^T D^{-\frac{1}{2}} (W - \alpha U^T U - \beta Z) D^{-\frac{1}{2}} Y).
 \end{aligned} \quad (9)$$

3.2.5 Tune parameters (α or β)

For parameter tuning, we use the Calinski–Harabasz index (Caliński and Harabasz, 1974) to evaluate the clustering performance without labels by measuring the cohesion and the separation of the clusters. There is only one parameter to be tuned in all the modes, and a one-dimensional search is conducted for the optimal parameter in a range of candidate α or β .

3.2.6 Different SolidBin modes

There are several SolidBin modes according to the prior adopted information. The summary of different modes and the summary of performance of different modes compared in the experiments are shown in Table 1.

4 Experiments

4.1 Data

As shown in Table 2, we compared SolidBin with the-state-of-art binners on five benchmark datasets.

- Multi-sample dataset on the species level: ‘SpeciesMock’ (Alneberg *et al.*, 2014)

The dataset was constructed based on the analysis of 16S rRNA samples originated from the Human Microbiome Project (Huttenhower *et al.*, 2012) and consisted of 101 different species across 96 samples. A total of 37 628 contigs remain for binning after co-assembly and filtering. The dataset is used for evaluating the influence of sample numbers on different SolidBin modes.

- Multi-sample dataset on the strain level: ‘StrainMock’ (Alneberg *et al.*, 2014)

The dataset was constructed to test the performance of the binners at different levels (Huttenhower *et al.*, 2012). It consisted of 20 different species or strains across 64 samples. A total of 9417 contigs remain for binning after co-assembly and filtering.

- Single-sample dataset: ‘SimHC’ (Wu *et al.*, 2014)

The single-sample dataset was simulated by Wu *et al.* (2014) and 29 535 contigs remain for binning after assembling and filtering. SimHC simulated high-complexity communities lacking dominant populations and it contains 100 genomes. A total of 13 918 out of 29 535 co-assembled contigs are unambiguously labeled on the species level for evaluation by Wang (Wang *et al.*, 2017) and we binned all the contigs. The dataset is used for evaluating the performance of different binners on single-sample datasets.

- Real dataset: ‘MetaHIT’

The dataset from MetaHIT consortium (Qin *et al.*, 2010) contains 264 samples, is the same dataset used in COCACOLA (Lu *et al.*, 2017a) and MetaBAT (Kang *et al.*, 2015). A total of 17 136 out of 192 673 co-assembled contigs are unambiguously labeled on

the species level for evaluation (Lu et al., 2017a). We binned the contigs with unambiguously labels for evaluation as in Lu et al. (2017a) and Yu et al. (2018). The dataset is used for evaluating the performance of different binners on high-complexity real datasets.

- Real dataset: ‘Sharon’ (Sharon et al., 2013)

The dataset from a time-series study of microbiome samples from a premature infant contains 18 samples. All the contigs are binned, and 2614 of 5579 contigs are unambiguously labeled on the species level for evaluation (Lu et al., 2017a). The dataset is used for evaluating the performance of different binners on low complexity real datasets.

4.2 Evaluation metric

For the unambiguously labeled contigs belonging to the datasets, the measures including precision, recall, *F*-score, Normalized Mutual Information and Adjusted Rand Index are used to evaluate the binning results and their definitions are shown in Supplementary Material.

4.3 Experimental procedures

We compared SolidBin with different modes to five advanced binners: CONCOCT-0.4.0, COCACOLA-python, MaxBin 2.2.4, MetaBAT 2.12.1 and BMC3C, respectively.

First, we conducted some preliminary experiments to show the necessity and effectiveness of determining the ratio of the pairwise constraints adaptively in SolidBin-SFS, and the accuracy of different constraints. We then compared the performance SolidBin with different modes with other binners. Finally, we investigated the effect of the number of samples on the performance of SolidBin-SFS and SolidBin-coalign based on sub-samples of the speciesmock dataset.

Table 3. The adjacency relationship between contigs in different datasets

Dataset	Estimated link ratio (%)	Real link ratio (%)
SpeciesMock	1.54	1.55
StrainMock	9.30	9.67
SimHC	1.58	1.74
Sharon	34.65	19.19
MetaHIT	3.71	3.44

Note: For Sharon and MetaHIT datasets, only unambiguously labeled contigs are considered.

Table 4. The accuracy of the constraints generated by different SolidBin modes

Mode		SpeciesMock		StrainMock		SimHC		Sharon_partial		MetaHIT_partial	
		Acc (%)	Num	Acc (%)	Num	Acc (%)	Num	Acc (%)	Num	Acc (%)	Num
SolidBin-SFS	ratio×0.2	100.00	4 410 072	100.00	1 658 091	72.38	1 824 430	99.24	363 286	99.80	2 190 826
	ratio×0.4	100.00	8 782 498	99.98	3 306 241	55.59	2 390 242	98.24	550 808	99.61	4 356 334
	ratio×0.6	100.00	13 154 614	99.89	4 950 081	45.13	2 633 940	93.86	769 282	98.83	6 474 940
	ratio×0.8	99.98	17 523 650	99.44	6 567 453	38.06	2 767 326	86.67	986 612	83.17	7 260 682
SoliBin-coalign	—	91.30	20 293 038	51.60	8 179 998	99.71	3 065 364	99.19	1 199 672	99.90	9 171 166
SolidBin-CL	—	99.99	1 370 250 256	99.62	67 069 672	100.00	188 819 244	99.96	3 955 520	99.95	237 793 612
SolidBin-SFS-CL	ratio×0.2	100.00	4 369 002	100.00	1 646 968	94.36	1 810 484	99.24	360 192	99.81	2 172 750
	ratio×0.4	100.00	8 727 240	99.98	3 249 328	89.83	2 376 178	98.60	547 706	99.71	4 336 152
	ratio×0.6	100.00	13 071 540	99.89	4 834 258	87.06	2 619 628	97.43	766 180	99.14	6 448 034
	ratio×0.8	99.98	17 416 512	99.44	6 382 828	85.28	2 752 684	95.74	983 508	92.36	7 228 632

4.4 Experimental results

To make a fair comparison, the input bin numbers are determined by the respective binners. Tables 5–9 show the results of SolidBin-SFS and SolidBin-SFS-CL when the ratio of pairwise constraints =40% of the estimated ratio. The values of ‘ α^* ’ and ‘ β^* ’ are the estimated parameters chosen by the Calinski–Harabasz index. COCACOLA-coalign means incorporating co-alignment information into COCACOLA.

4.4.1 Preliminary experiments

- Determine the ratio of the pairwise constraints adaptively.
Table 3 shows the results of our method for determining the ratio of the pairwise constraints adaptively. The ‘real link ratio’ and ‘estimated link ratio’ are computed using Equation (2) according to the real labels and the binning result of SolidBin-naive, respectively. The ‘real link ratio’ varies greatly among different datasets and the estimated link ratios are close to their corresponding real link ratio on most datasets. For the Sharon dataset, the cluster numbers are underestimated by SolidBin using the contigs unambiguously labeled on species level. Overall, the consistency between real and estimated link ratios shows the validity of our estimation method.

- Accuracy of constraints.

The accuracy of constraints defined as the fraction of true relationships among all the constrained contig pairs on different datasets is shown in Table 4. In order to obtain adequate constraints with high accuracy on the SolidBin-SFS mode, we take the contig pairs with high similarity according to the estimated ratio as constraints. We test the accuracy of SolidBin-SFS mode constraints with the proportion ranging from 20 to 80%, with 20% as an incremental interval, and the proportion is set to 40% in our experiments. The results show that all the multi-sample datasets can obtain constraints with high accuracy on the SolidBin-SFS mode with the proportion 20 and 40%. Single-sample dataset ‘SimHC’ cannot obtain constraints with high accuracy on the SolidBin-SFS mode, which is reasonable given that the effect of coverage information is limited in the case of a single sample. In contrast, the SolidBin-SFS-CL mode can generate constraints with higher accuracy than the corresponding SolidBin-SFS mode in the case of a single sample, for instance, from 0.7238 to 0.9436 on the SimHC dataset. Moreover, the SolidBin-coalign mode can generate relatively accurate constraints except for the strain level dataset ‘StrainMock’, because the co-alignment information is reliable on the species or higher levels other than the strain level. For example, the accuracy of SolidBin-coalign constraints on the SpeciesMock is 0.9130, while the one on the StrainMock dataset is 0.5160.

Table 5. The performance of COCACOLA, CONCOCT, MaxBin, MetaBAT, BMC3C and SolidBin on the StrainMock dataset

StrainMock	Recall (%)	Precision (%)	F-score (%)	NMI (%)	ARI (%)
COCACOLA	99.20	99.26	99.23	98.63	99.03
COCACOLA-coalign	95.87	93.53	94.69	95.46	94.57
MaxBin	96.35	91.00	93.60	98.36	86.69
CONCOCT	98.21	93.85	95.98	96.24	93.99
MetaBAT	81.24	59.87	68.94	69.76	33.50
BMC3C(9401/9417)	99.01	99.01	99.01	98.42	98.73
SolidBin-naive	98.92	98.92	98.92	98.49	98.91
SolidBin-SFS($\alpha^*=20$)	99.29	99.29	99.29	98.75	99.13
SolidBin-coalign($\alpha^*=10$)	99.46	92.59	95.90	95.93	87.74
SolidBin-CL($\beta^*=0.1$)	99.29	99.29	99.29	98.74	99.12
SolidBin-SFS-CL($\alpha^*=30$)	99.29	99.29	99.29	98.75	99.12

Note: The optimal values of the results are in bold.

Table 6. The performance of COCACOLA, CONCOCT, MaxBin, MetaBAT, BMC3C and SolidBin on the 'StrainMock' dataset (evaluated on species level)

StrainMock	Recall (%)	Precision (%)	F-score (%)	NMI (%)	ARI (%)
COCACOLA	91.81	99.96	95.71	95.90	86.67
COCACOLA-coalign	91.27	99.94	95.41	96.89	88.01
MaxBin	91.01	91.01	91.01	91.87	78.50
CONCOCT	91.11	99.63	95.18	96.82	89.50
MetaBAT	81.24	67.32	73.63	72.46	34.10
BMC3C(9401/9417)	91.58	99.97	95.59	95.94	86.51
SolidBin-naive	91.94	99.99	95.80	96.06	86.89
SolidBin-SFS($\alpha^*=20$)	92.00	99.99	95.83	96.06	86.92
SolidBin-coalign($\alpha^*=10$)	99.46	99.97	99.71	99.39	98.83
SolidBin-CL($\beta^*=0.1$)	92.01	99.98	95.83	96.05	86.92
SolidBin-SFS-CL($\alpha^*=30$)	92.00	99.99	95.83	96.06	86.92

Note: The optimal values of the results are in bold.

Table 7. The performance of COCACOLA, CONCOCT, MaxBin, MetaBAT, BMC3C and SolidBin on the 'SimHC' dataset

SimHC	Recall (%)	Precision (%)	F-score (%)	NMI (%)	ARI (%)
COCACOLA	92.51	77.09	84.09	90.17	76.15
COCACOLA-coalign	94.41	80.43	86.86	91.66	80.73
MaxBin	85.84	83.28	84.54	89.34	76.90
CONCOCT	96.92	82.70	89.25	93.71	82.17
MetaBAT	90.38	72.18	80.26	86.42	57.07
BMC3C(13 918/13 918)	69.18	93.03	79.35	90.23	68.58
SolidBin-naive	90.20	78.88	84.16	89.95	75.78
SolidBin-SFS($\alpha^*=0$)	90.20	78.88	84.16	89.95	75.78
SolidBin-coalign($\alpha^*=10$)	98.85	90.94	94.73	97.32	93.19
SolidBin-CL($\beta^*=0.2$)	97.15	87.22	91.92	96.30	87.22
SolidBin-SFS-CL($\alpha^*=0$)	90.20	78.88	84.16	89.95	75.78

Note: The optimal values of the results are in bold.

4.4.2 The binning results on the simulated datasets

For the 'StrainMock' dataset, COCACOLA, SolidBin and BMC3C have better performance compared with other methods as shown in Table 5, and SolidBin-SFS performs best in general. Just considering the binning results of the different SolidBin modes, SolidBin-SFS and SolidBin-CL can generate constraints with high accuracy on this dataset as shown in Table 4, and have better performance in terms of all metrics compared with SolidBin-naive. On the other hand, since the co-alignment information is not reliable on the strain level, both the COCACOLA-coalign and SolidBin-coalign have worse

performance than COCACOLA and SolidBin-SFS on this strain level dataset.

To figure out whether the coalignment information is useful to the 'StrainMock' dataset, we evaluated the results of the StrainMock at the species level. As shown in Table 6, both the COCACOLA and SolidBin have better performance with coalignment information at the species level. SolidBin-coalign has larger improvement than COCACOLA-coalign. For example, the ARI of COCACOLA-coalign increases from 0.8667 (COCACOLA) to 0.8801, and the ARI of SolidBin-coalign increases from 0.8689

Table 8. The performance of COCACOLA, CONCOCT, MaxBin, MetaBAT, BMC3C and SolidBin on the 'MetaHIT' (partial) dataset

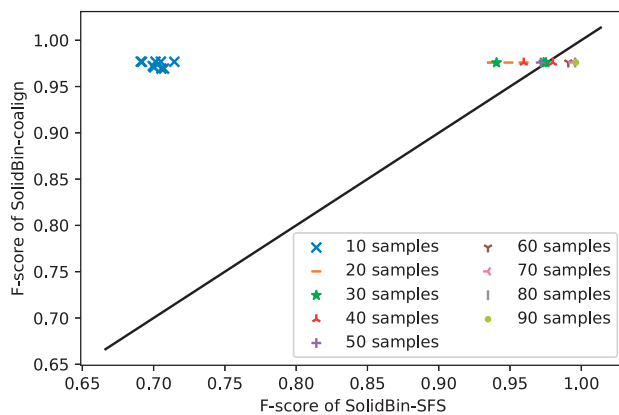
MetaHIT_partial	Recall (%)	Precision (%)	F-score (%)	NMI (%)	ARI (%)
COCACOLA	91.87	83.63	87.56	84.26	76.38
COCACOLA-coalign	91.92	85.39	88.54	86.04	80.91
MaxBin	84.75	73.58	78.77	77.84	69.02
CONCOCT	80.41	90.26	85.05	85.29	74.33
MetaBAT	74.34	52.59	61.60	61.37	30.19
BMC3C(17 122/17 136)	87.50	91.45	89.43	87.90	82.82
SolidBin-naive	85.85	87.15	86.49	85.76	77.30
SolidBin-SFS($\alpha^*=10$)	86.24	87.22	86.73	85.91	77.95
SolidBin-coalign($\alpha^*=40$)	97.01	97.18	97.10	96.01	93.49
SolidBin-CL($\beta^*=0.2$)	90.47	91.17	90.82	89.98	85.34
SolidBin-SFS-CL($\alpha^*=0$)	85.85	87.15	86.49	85.76	77.30

Note: The optimal values of the results are in bold.

Table 9. The performance of COCACOLA, CONCOCT, MaxBin, MetaBAT, BMC3C and SolidBin on the 'Sharon' dataset

Sharon	Recall (%)	Precision (%)	F-score (%)	NMI (%)	ARI (%)
COCACOLA	91.21	97.98	94.47	91.58	93.21
COCACOLA-coalign	92.65	97.49	95.00	91.43	92.02
MaxBin	96.02	90.57	93.21	89.18	87.74
CONCOCT	93.66	98.90	96.21	93.53	94.53
MetaBAT	90.91	68.74	78.28	71.49	58.97
BMC3C(2591/2614)	99.27	99.02	99.14	97.16	97.98
SolidBin-naive	98.11	97.58	97.84	93.90	94.38
SolidBin-SFS($\alpha^*=1$)	99.33	98.76	99.04	96.61	97.66
SolidBin-coalign($\alpha^*=10$)	99.64	99.09	99.37	97.71	98.46
SolidBin-CL($\beta^*=0.1$)	99.44	98.88	99.16	97.03	97.96
SolidBin-SFS-CL($\alpha^*=1$)	99.53	98.97	99.25	97.35	98.16

Note: The optimal values of the results are in bold.

**Fig. 2.** Evaluation of the result of SolidBin-coalign and SolidBin-SFS on subsamples of the 'SpeciesMock' dataset

(SolidBin-naive) to 0.9883. Therefore, SolidBin-coalign can better utilize coalignment information than COCACOLA-coalign, which highlights the effectiveness of spectral clustering in incorporating pairwise constraints.

For the only single-sample dataset 'SimHC', CONCOCT and SolidBin have better performance compared with other methods as shown in Table 7, and SolidBin performs better in two modes. The F-score, NMI and ARI of SolidBin-coalign reach 0.9473, 0.9732 and 0.9319, respectively. In comparison, CONCOCT achieves 0.8925 in terms of F-score, 0.9371 in terms of NMI, and 0.8217 in

terms of ARI, respectively. Taking different SolidBin modes into consideration, the SolidBin-SFS mode cannot obtain constraints with high accuracy and have better binning results than SolidBin-naive on the single dataset due to insufficient sample number. In contrast, sequence alignment information is quite useful in improving the binning results of the dataset. Both SolidBin-coalign and SolidBin-CL have better performance in terms of all metrics compared with other binners and SolidBin modes. Both COCACOLA-coalign and SolidBin-coalign have better performance than COCACOLA and SolidBin-naive on this dataset, but the improvement of SolidBin-coalign is more significant, from 0.8416 to 0.9473 in terms of F-score.

4.4.3 The binning results of the real datasets

For the 'MetaHIT' dataset, the results of the 17 136 contigs with unambiguous labels are shown in Table 8. SolidBin have better performance than other tools in general. Both SolidBin-SFS and SolidBin-SFS-CL have comparable performance with COCACOLA and CONCOCT, and achieve much better performance than Maxbin and MetaBAT, but worse performance than BMC3C. Considering the performance of different SolidBin modes, all the other modes have better performance than the SolidBin-naive mode, especially the modes that use the sequence alignment information. Both COCACOLA-coalign and SolidBin-coalign have better performance than COCACOLA and SolidBin-naive on this dataset, but the improvement of SolidBin-coalign is more significant, from 0.7730 to 0.9349 in terms of ARI and from 0.8649 to 0.9710 in terms of F-score.

Table 10. The running time and memory usage of the binners on the different datasets

Binners	StrainMock		SimHC		Sharon		MetaHIT	
	Time	Memory (MiB)	Time	Memory (MiB)	Time	Memory (MiB)	Time	Memory (MiB)
COCACOLA	4 min 50 s	220	45 min 30 s	403	2 min 01 s	148	11 min 33 s	473
COCACOLA-coalign	6 min 54 s	2881	54 min 45 s	4144	3 min 33 s	1777	16 min 16 s	2024
MaxBin	22 min 08 s	49	63 min 10 s	160	4 min 31 s	27	600 min 43 s	47
CONCOCT	2 min 07 s	466	28 min 30 s	1151	2 min 14 s	359	88 min 05 s	1098
MetaBAT	39 s	112	6 min 16 s	412	13 s	69	1 min 51 s	128
BMC3C	9 min 58 s	1327	104 min 10 s	2681	1 min 47 s	477	25 min 23 s	1897
SolidBin-naive	7 min 10 s	4258	71 min 37 s	35 932	2 min 32 s	1549	26 min 31 s	11 425
SolidBin-SFS	22 min 45 s	5176	379 min 26 s	40 631	6 min 51 s	2022	95 min 37 s	13 944
SolidBin-coalign	21 min 18 s	4955	302 min 43 s	40 441	7 min 43 s	2639	73 min 43 s	14 033
SolidBin-CL	28 min 20 s	16 640	385 min 20 s	196 770	6 min 43 s	3056	99 min 28 s	47 369
SolidBin-SFS-CL	37 min 48 s	16 342	566 min 48 s	197 001	9 min 35 s	3155	143 min 02 s	47 519

For the ‘Sharon’ dataset, we binned all the contigs and the results of the 2 614 contigs with unambiguous labels are shown in [Table 9](#) and the estimated input bin number is 7 for SolidBin. All the modes of SolidBin and BMC3C have better performance than other tools. The *F*-score, NMI and ARI of the best performed SolidBin-coalign reach 0.9937, 0.9771 and 0.9846, respectively. In comparison, the *F*-score, NMI and ARI of CONCOCT, best one among other binners on the dataset, is 0.9621, 0.9353 and 0.9453, respectively. Taking different SolidBin modes into consideration, all the other modes have better performance than the SolidBin-naive mode, which shows that both the additional biological information and the internal information are useful for the binning. On the other hand, COCACOLA-coalign achieved similar performance with COCACOLA, which indicates the inefficiency of COCACOLA in incorporating coalignment information.

4.4.4 Evaluation of the impact of incorporating constraint information on sub-samples of the ‘SpeciesMock’ dataset
In order to evaluate the impact of co-alignment information and the ML constraints generated by SolidBin-SFS mode when the number of samples is small, we investigate the performance of SolidBin-SFS and SolidBin-coalign on sub-samples without overlapping of the ‘SpeciesMock’ dataset. The sample size ranges from 10 to 90, with 10 as increment. As shown in [Figure 2](#), the results of SolidBin-SFS vary greatly with the number of samples and SolidBin-SFS can obtain better performance when the number of samples is large. The results of ARI and NMI are in the [Supplementary Material](#). The results of SolidBin-coalign are relatively stable and can obtain good performance even when the number of samples is relatively low.

4.5 Running time and memory usage of the binners
All the experiments were done on a machine with 4-way 6-core 1.87 GHz Intel Xeon CPUs and 1 T memory. We ran all the binners with multiple threads. The running time and memory usage of the binners on different datasets are as shown in [Table 10](#). On the one hand, SolidBin has excellent performance on the binning, but on the other hand, it takes a lot of time and memory to compute the similarity matrix for singular value decomposition, especially for the datasets with large contig numbers. However, unlike other binners, the number of the samples almost has no impact on the running time and memory of SolidBin.

4.6 Analysis of the results
In summary, SolidBin with different modes have better performance than other binners in most cases. For the different SolidBin modes, the performance mainly depends on the accuracy of the constraints

of different modes, which will be affected by the properties of the datasets, such as the number of samples. SolidBin-SFS is a reliable mode without any taxonomy information when the number of samples is large, and our method can determine the ratio of the pairwise constraints adaptively according to the data distribution. Sequence alignment information can improve the binning results markedly such as the SolidBin-coalign mode and the SolidBin-CL mode, especially when there are insufficient samples and the genomes contained in the dataset are at the species level, not at the strain level. The performance profiles of different SolidBin modes are shown in [Table 1](#). Both COCACOLA and SolidBin can utilize co-alignment information, however, the improvements of SolidBin-coalign are always more significant, which shows that spectral clustering algorithm can integrate additional information better.

5 Discussion and conclusion

In this paper, we developed a binning method, SolidBin, based on semi-supervised spectral clustering. To our knowledge, this is the first method to apply semi-supervised spectral clustering for contig binning. It uses two types of prior information: ML constraints and CL constraints. We examined two ways to generate ML constraints, one is taxonomy independent, named as SolidBin-SFS and the other is based on the co-alignment information, which can take advantage of some contigs that belong to the known reference genomes, named as SolidBin-coalign. In this paper, our method is compared with five advanced binning tools, CONCOCT, COCACOLA, MaxBin, MetaBAT and BMC3C, using five different types of datasets. As shown in the Section 4.4, our method, compared with the-state-of-the-art binning methods, has best performance in terms of *F*-score, ARI and NMI in most cases.

However, our method has its own limitations. For example, SolidBin-SFS does not perform best when the number of samples is small. But SolidBin-coalign may perform well under these circumstances if the genomes contained in the dataset are at the species level. However, how to choose α , β and bin number K is still challenging. In addition, how to extend our method to handle large-scale datasets is still worthy further investigation.

In future research, we plan to incorporate more prior information into our method, such as linkage information used in [Lu et al. \(2017a\)](#), gene prediction information ([Sunagawa et al., 2015](#)) and DNA methylation ([Beaulaurier et al., 2018](#)).

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