

Genome analysis

PlasBin-flow: A flow-based MILP algorithm for plasmid contigs binning

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Abstract

Motivation: The analysis of bacterial isolates to detect plasmids is important due to their role in the propagation of antimicrobial resistance. In short-read sequence assemblies, both plasmids and bacterial chromosomes are typically split into several contigs of various lengths, making identification of plasmids a challenging problem. In *plasmid contig binning*, the goal is to distinguish short-read assembly contigs based on their origin into plasmid and chromosomal contigs and subsequently sort plasmid contigs into bins, each bin corresponding to a single plasmid. Previous works on this problem consist of *de novo approaches* and *reference-based approaches*. De novo methods rely on contig features such as length, circularity, read coverage, or GC content. Reference-based approaches compare contigs to databases of known plasmids or plasmid markers from finished bacterial genomes.

Results: Recent developments suggest that leveraging information contained in the assembly graph improves the accuracy of plasmid binning. We present PlasBin-flow, a hybrid method that defines contig bins as subgraphs of the assembly graph. PlasBin-flow identifies such plasmid subgraphs through a mixed integer linear programming model that relies on the concept of network flow to account for sequencing coverage, while also accounting for the presence of plasmid genes and the GC content that often distinguishes plasmids from chromosomes. We demonstrate the performance of PlasBin-flow on a real data set of bacterial samples.

Availability: <https://github.com/acme92/PlasBin-flow>

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1 Introduction

Antimicrobial resistance (AMR) has emerged to become a major threat to public health. AMR is developed in bacteria through the propagation of antimicrobial resistance genes, often facilitated by mobile genetic elements (MGEs), which can be exchanged between bacteria through horizontal gene transfer (HGT) (16; 8). Plasmids are short circular extra-chromosomal MGEs known to be a major vector for the spread of AMR through HGT (7). Thus, the detection of plasmids in bacterial genomes is an important problem in microbial genomics, with applications in environmental ecology and public-health surveillance.

With the advent of DNA sequencing technologies, it is now possible to obtain whole-genome sequencing (WGS) data at a low cost, and the analysis of WGS data for public-health surveillance is now common (21; 10). Despite the increasing availability of long-reads sequencing data (22), most surveillance approaches in microbial genomics rely on the analysis of short-reads WGS datasets. Such datasets are generally assembled using assemblers such as Unicycler (24), SKESA (20) or SPAdes (5). It is typically impossible to assemble full sequences of chromosomes and plasmids using short-read WGS data, and consequently a short-read assembly contains a mixture of contigs coming from the chromosome and individual plasmids present in the sample.

The problem of detecting plasmids from a short-read assembly can be considered at three different levels: classification, binning and assembly. At the *classification* level, the aim is to classify the origin of a contig, which

is either a plasmid or the chromosome. Current state-of-the-art tools rely on machine-learning approaches and include mlplasmids (3), PlasClass (17), Deepplasmid (1) and RFPlasmid (23). However, a bacterial genome can contain several plasmids, and contigs classification does not provide a precise view of the plasmid content of a genome. This is addressed by the *plasmid binning problem*, where the aim is to group plasmid contigs into bins, such that the contigs in a bin are likely to originate from the same plasmid. Lastly, *plasmid assembly* aims to order and orient contigs in a plasmid bin into a fully assembled plasmid sequence.

In this work, we focus on the binning problem, motivated by the observation that, for downstream analysis, the most useful information are groups of genes belonging to a given plasmid. While the order and orientation of these genes along the plasmid sequence can yield interesting insights, important tasks such as plasmid typing can be done from the gene content and does not consider contigs or gene order (11; 9). Two main methodological avenues have been used in developing plasmid binning methods: reference-based and de-novo binning. Reference-based methods, such as MOB-recon (18), map contigs to a reference database of plasmids or plasmid gene sequences, clustered into families of plasmids. Contigs are then binned together if they match with plasmids belonging to the same reference cluster. The reliance on a reference database can potentially hinder the ability of reference-based methods to identify novel plasmids. De-novo binning methods, such as Recycler (19), PlasmidSPAdes (2) and gplas (4), bypass the requirement of reference sequences and rely instead on contig features assumed to be specific to different plasmids present in a bacterial cell. A feature central to many plasmid binning methods is contig read coverage. Different plasmids occur in the genome isolate in varying number (sometimes hundreds) of copies. Thus, the coverage of plasmid contigs is expected to be significantly different between the plasmids and chromosomes, as well as between different plasmids. Another feature used to detect plasmids in a bacterial assembly is the GC content, which is often slightly different in plasmids, especially short ones, as compared to the chromosome (15). Recent plasmid binning methods HyAsP (14) and PlasBin (12) combine ideas from both reference-based and de-novo binning.

Several methods do leverage information from the *assembly graph*, a graph containing contigs as nodes and possible connections between them supported by sequencing data as edges; widely used bacterial genomes assemblers such as SPAdes, Unicycler and SKESA generate such a graph as a complement to the contigs assembly. The rationale for using the assembly graph is that individual molecules, such as chromosomes or plasmids, ideally correspond to walks in this graph. However, although due to sequencing and assembly artifacts, some edges from such walks can be missing, and spurious edges and repeated contigs may form unwieldy tangled structures that are difficult to analyze. Nonetheless, adjacent nodes often share the same molecule of origin and thus edges from the assembly graph provide an important information. For example, Recycler peels off cycles from the assembly graph assuming uniform coverage of sequenced plasmids; PlasmidSPAdes estimates the chromosomal coverage from the whole assembly, removes contigs with similar coverage as that of the chromosome and then computes putative plasmid bins from the connected components of the remaining graph. Gplas constructs a plasmidome network by defining a set of walks in the assembly graph. Each walk is initialized with a contig that is classified as plasmidic using mlplasmids (3). It uses a heuristic that extends a walk one contig at a time, based on a score that accounts for similarity between the contig read coverage and the mean coverage of the currently assembled walk. Putative plasmid bins are selected from the network using graph partitioning algorithms. HyAsP (14) uses a greedy walk-building heuristic that aims to iteratively extract walks from the assembly graph. The algorithm relies on a walk-extension objective function defined in terms of coverage consistency, GC content consistency, and high plasmid gene density (from a database of reference

plasmid genes) of contigs included in the walk. PlasBin (12) extends the approach used in HyAsP into an Mixed-Integer Linear Programming (MILP) formulation that defines plasmid bins as connected subgraphs of the assembly graph that optimize an objective function similar to the HyAsP one.

In the present work, we introduce a new optimization algorithm for the plasmid binning problem. The main novelty of our work is to state the problem as a network flow problem, where the flow accounts for the expected uniform coverage of the sequenced plasmid, while also considering the GC content and the density of reference plasmid genes in a plasmid bin. Our work improves upon PlasBin in several ways, including a method for scoring GC content based on a probabilistic model. We compare our method PlasBin-flow against PlasBin, HyAsP, plasmidSPAdes, MOB-recon, and gplas on a data set of 133 bacterial samples.

2 Method

Here we present PlasBin-flow, a network flow-based MILP formulation for the plasmid binning problem. It extracts putative plasmid bins as connected components of uniform coverage from an assembly graph. PlasBin-flow is a hybrid method that relies on a reference database of closed genome assemblies including plasmids. Our reference database is described in detail in section 3.1.

2.1 Input: Contigs and the assembly graph

The input of the PlasBin-flow consists of the contigs and the assembly graph from a short-read assembly of a bacterial isolate. In this work, we use Unicycler (24) to generate the assembly, but any other assembler that provides an assembly graph (such as PlasmidSPAdes (2) or SKESA (20)) can be used, as PlasBin-flow does not depend on a specific assembler.

The set of all contigs in the short-read assembly is denoted by C . Every contig $c \in C$ has two extremities: head c_h and tail c_t . Pairs of contigs potentially adjacent in the genome sequence are connected to each other via edges of the assembly graph, each edge linking two specific contig extremities. We denote the set of edges in the assembly graph by E . Each undirected edge $e \in E$ connects two extremities c_u and d_v (c_u and d_v being either a head or a tail of contigs c and d respectively) and is represented as an unordered pair $\{c_u, d_v\}$.

The PlasBin-flow input also contains a reference database of plasmid genes. In a practical application, the reference database would contain all known plasmid genes; however, since an exact match to the reference database would indicate a strong determination of the contig origin, we have taken care in our experiments to withdraw information from samples used for testing from the database used (see 3.1 for further description).

Contig features. PlasBin-flow considers several features associated to a contig c , namely its GC content gc_c , length ℓ_c , sequencing coverage rc_c and plasmid gene density gd_c . The sequencing coverage is the normalized coverage provided by Unicycler for each contig, i.e. the average base coverage of the contig normalized by the median coverage of the contigs in the whole assembly; chromosomal contigs are thus expected to have a normalized coverage close to 1. To obtain the plasmid gene density, we map each contig against the reference database of plasmid genes using blastn (version 2.6.0) (6), discarding any match with identity below 95% or a match that covers less than 95% of a gene. The plasmid gene density gd_c is the fraction of the sequence length of contig c covered by these matches.

Contigs passing certain thresholds related to length and gene density are referred to as *seed* contigs. These contigs are more likely to be part of a plasmid than others. We describe how to determine corresponding

thresholds using the reference database in Section 2.4. The set of all seed contigs is denoted as C_s , and we also use a boolean feature $s_c = [c \in C_s]$.

2.2 PlasBin-flow overview

PlasBin-flow works iteratively, identifying one plasmid bin in each iteration by solving a mixed integer linear program (MILP), described in Section 2.3. In each iteration, the identified bin is required to contain at least one seed contig. After each iteration, PlasBin-flow updates the assembly graph (contigs, edges and contigs read coverage) to account for the removal of the plasmid bin. The modified assembly graph is then used as input for the next iteration. This process is repeated until the modified assembly graph does not contain any seed contig.

PlasBin-flow defines a plasmid bin as a connected subgraph of a network defined from the assembly graph, while enforcing that this subgraph contains at least one seed contig and assigning to this subgraph (1) a *flow value* defined in terms of the coverage of contigs it contains, that serves as a proxy for the copy number of the plasmid (bin) defined by the subgraph, and (2) a GC content value. This subgraph is chosen as the one optimizing an objective function defined as a linear combination of several terms: (1) the flow value, (2) a GC content term penalizing contigs whose GC content does not agree with the GC content assigned to the subgraph, and (3) a plasmid gene density term penalizing contigs with a low plasmid gene density. An important and novel feature of PlasBin-flow is that it assigns a *multiplicity* to each contig in a plasmid bin, with repeated contigs having multiplicity greater than 1.

The GC content assigned to a solution is defined as a discrete category, each of the k categories corresponding to a GC content interval. This set K of k intervals is pre-defined through a preliminary analysis of the reference database. For each contig c and each GC content interval $b \in K$, we pre-compute a *GC content penalty* $gc_pen_{c,b}$ defined in terms of the likelihood that a sequence of length ℓ_c and GC content gc_c would be observed in a plasmid whose GC content is in interval b . We provide a detailed description of $gc_pen_{c,b}$ in Section 2.5.

2.3 MILP formulation

Network and flow: The MILP implemented in PlasBin-flow takes as an input network N obtained by modifying the original assembly graph. Using contigs C and edges E in the undirected assembly graph, we define a directed network $N = (C \cup \{s, t\}, E')$ including two new nodes: the source s and the sink t . New set of edges E' is obtained by (1) doubling each edge $\{c_u, d_v\} \in E$ into two directed edges $(c_u, d_v), (d_v, c_u)$, (2) adding edges (s, c_t) and (s, c_h) for every seed contig c , and (3) adding edge (d_h, t) and (d_t, t) for every contig d (regardless of its seed status). Edge $e = (c_u, d_v)$ has *capacity* $cap_e = \min\{rc_c, rc_d\}$ that upper-bounds the flow through e ; if e links a contig extremity to s or t , its capacity is $M = \max_{c \in C}\{rc_c\}$, the maximum coverage of the whole assembly.

The choice of the contigs and edges in the optimal subgraph is guided by a flow through the edges of the network. We define a flow as a map $f : E' \rightarrow \mathbb{R}$ that satisfies the following constraints:

- *Capacity constraints:* The flow through an edge cannot exceed its capacity; the cumulative flow through a contig extremity can not exceed the read coverage of the contig.
- *Conservation constraints:* The cumulative flow into a contig extremity should be equal to the cumulative flow out of the other extremity of the same contig.
- *Source and sink constraints.* Exactly one edge outgoing from s and incoming to t has a non-zero flow value.

The value of flow F is thus equal to the quantity flowing out of s , which, by the conservation constraints, is also equal to the quantity flowing into t .

Decision variables: The MILP decision variables define a *solution* to the optimal plasmid bin finding problem, composed of a subgraph of N , GC content value associated to it, and a flow value. To each contig c and edge e in N , we associate a decision binary variable (denoted respectively by x_c and y_e) indicating whether it is a part of the solution subgraph; we also call edges and contigs included in the solution *active*.

We associate to every edge $e \in N$ a continuous decision variable f_e that encodes the quantity $f(e)$ flowing through the edge. The overall flow value (flowing out the source s and into the sink t) is encoded by a continuous decision variable F . The flow F also helps in determining the multiplicity for each contig in the plasmid bin defined by a solution. Let c_u^+ be the set of edges incoming into extremity c_u , and c_u^- be the set of edges outgoing from c_u . For any contig extremity c_u , we denote by c_u^+ the set of edges into c_u and c_u^- the set of edges out of c_u . The multiplicity of a contig m_c in the plasmid bin is related to the flow value as $m_c = \sum_{e \in c_h^+ \cup c_t^+} f_e / F$.

The GC content of the solution subgraph is chosen as one of the k pre-defined intervals and is encoded by binary decision variables: for every pre-defined GC content interval $b \in K$, the decision variable GC_b is assigned value 1 if b is the chosen GC content interval, and 0 otherwise.

The total number of decision variables defining a solution to the plasmid bin finding problem (subgraph and associated flow and GC content) is thus $O(|C| + |E| + |K|)$. Note that the subgraph of N defined as above may not be connected. However, we aim to define a plasmid bin by a connected subgraph. We describe later how we handle this problem through delayed constraints generation.

Objective function: We formulate the MILP as a maximization problem with objective function $F + GC + GD$. Here, F represents the overall flow value from s to t , GC is a penalty for active contigs whose GC content differs significantly from the overall bin GC content, and GD is a term penalizing inclusion of contigs with low plasmid gene density. In particular,

$$GC = \sum_{c \in C, b \in K} -(gc_pen_{c,b}) \cdot x_c \cdot GC_b,$$

and

$$GD = \sum_{c \in C} (gd_c - 0.5) \cdot x_c.$$

Note that the term GC is not linear as it involves the product of two binary decision variables. Such a product can be linearized by addition of a single auxiliary binary decision variable and three associated linear constraints (see (13)). The gene density in the objective function is used in a similar manner as probability. Thus, the MILP is discouraged from choosing contigs with gene density lower than 0.5 as part of the solution.

The rationale for this objective function is as follows: by maximizing the flow value F , we aim to identify a high copy-number plasmid that is more likely to be reliable. The two penalty terms aim to assign to the plasmid bin a GC content that reflects the GC content of the selected contigs (we refer to Section 2.5 for a more detailed discussion) and prefer contigs that have a higher plasmid genes density and are thus more likely to originate from actual plasmids. Our approach that seeks an optimal plasmid bin through exact optimization improves upon greedy heuristics (such as HyAsP) in that it will include contigs that impact negatively the objective function if this inclusion allows to connect to other contigs whose positive contribution to the objective function will be beneficial overall.

Constraints: We now describe the constraints used in the formulation:

- Exactly one edge out of s is part of the solution:

$$\sum_{e=(s,c_u), c \in C_s, u \in \{h,t\}} y_e = 1.$$

This also ensures that every solution contains at least one seed contig.

- The flow through an edge e is non-zero only if e was selected in the solution subgraph and can not exceed its capacity:

$$f_e \leq \text{cap}_e \cdot y_e.$$

- For any contig c , the cumulative flow through c can not exceed its read coverage:

$$\sum_{e \in c_h^+ \cup c_t^+} f_e \leq r c_c.$$

- For any contig c , the cumulative flow into c_h should be equal to the cumulative flow out of c_t , and conversely:

$$\sum_{e \in c_h^+} f_e = \sum_{e \in c_t^-} f_e, \quad \sum_{e \in c_t^+} f_e = \sum_{e \in c_h^-} f_e.$$

- Flow value F should be equal to the flow out of s and into t :

$$F = \sum_{e=(s,v)} f_e, \quad F = \sum_{e=(v,t)} f_e,$$

- If an edge $e = (c_u, d_v)$, $c \in C$, $u, v \in \{h, t\}$ is in the solution ($y_e = 1$), then the contigs c and d should also be in the solution. Note that the converse need not be true. In other words, contigs c and d may be part of the solution due to other edges. In that case, y_e should be 0. To implement this, we use the following constraints:

$$y_e \leq \min\{x_c, x_d\}$$

- A contig c is active (selected in the solution) if at least one edge incident to one of its extremities is active.

$$x_c = \min\{1, \sum_{e \in c_h^+ \cup c_t^+} y_e\}$$

- An edge e is active only if the flow through the edge is non-zero. To implement this condition, we introduce an auxiliary variable $F_e = F \cdot y_e$. We further add the constraint:

$$F_e \leq f_e$$

Thus, if there is no flow through edge e , F_e is forced to be 0 and in turn $y_e = 0$. To handle the fact that F_e is the product of a binary variable and a continuous variable, we use the same approach as for the GC term of the objective function (13).

Ensuring connectivity: A solution that satisfies the above constraints may not be connected. It will contain a connected component with s and t , but may also contain other components disconnected from s and t . Such disconnected components have a closed flow circulating within the component without violating any of the initial constraints.

If the proposed solution contains disconnected components, we mute the edges in the disconnected component. For each edge e in the disconnected component, we add a constraint $y_e = 0$ and run the MILP again. This process is repeated until the MILP returns a solution with a single component connected to s and t . While in theory this delayed constraint generation process could involve a large number of iterations, in practice we observe that it requires a small number of iterations.

2.4 Determining seeds

Similarly to HyAsP and PlasBin, PlasBin-flow enforces that every plasmid bin contains at least one seed contig under the assumption that such contigs were identified as likely to belong to a plasmid due to their sequence features (length and plasmid gene density). We used the reference database for which closed assemblies with annotated plasmids were available, to obtain the thresholds determining seed contigs. Using Unicycler, we assembled the reads datasets for each sample in the reference database. We explored a grid of pairs of values (ℓ, gd) respectively for the contig length and plasmid gene density, with ℓ ranging from 50bp to 5000bp in increments of 50 and gd ranging from 0.01 to 1 in increments of 0.01. For each pair (ℓ, gd) we considered as a seed every contig c such that $\ell_c \geq \ell, gd_c \geq gd$; then mapping the contigs onto the closed assemblies of the reference dataset, for which plasmids are known, we counted (1) the number SP of plasmids onto which at least one seed contig was mapped to (seeded plasmids), and (2) the number NPS of seed contigs that mapped to the chromosome (non-plasmid seeds). We then determined the pair of thresholds that maximized the expression $SP - NPS$ and used them as thresholds defining seed contigs in our experiments. The pair of thresholds thus obtained was (2650, 0.58).

2.5 Determining GC content intervals and penalties

For each contig c , we associate a penalty relating the observed GC content of c to the GC content interval $b \in K$ associated to a solution to the MILP. This penalty is based on a probabilistic model in which we can compute the likelihood that a contig of a particular length and GC content will belong to a plasmid whose GC content is in a particular interval b . In this section we first define the model and describe how we define the GC content penalty term. Subsequently we describe the set of intervals K used in our experiments.

Probabilistic model. If a given contig c of length ℓ has GC content $x \in (0, 1)$, we model the observed number n of GC nucleotides within this contig by the binomial distribution with parameters x and ℓ . However, GC content within a plasmid may fluctuate for various biological reasons, and therefore we model contig-level GC content x as a random variable sampled from a beta distribution with parameters $\alpha = pm$ and $\beta = (1-p)m$, where $p \in (0, 1)$ is the true GC content of the whole plasmid and $m > 0$ is a fixed parameter. This results in n being sampled from the beta-binomial distribution with parameters ℓ , α and β ; namely, $\Pr(n|\ell) = \binom{\ell}{n} B(n+\alpha, \ell-n+\beta) / B(\alpha, \beta)$ where B is the beta function. Values α and β represent pseudocounts added to the observed counts of GC and non-GC nucleotides, respectively. We use $m = 10$ for the overall pseudocount.

In our scenario, p is an unknown value coming from GC content interval $b = [p_s, p_e]$, and we assume a uniform prior on p . The overall probability of b being the true source interval for observed count n is thus $\int_{p_s}^{p_e} \Pr(p|n, \ell) dp$, where $\Pr(p|n, \ell) \propto \Pr(n|p, \ell) \cdot \Pr(p)$ is obtained by the Bayes theorem. We normalize these probabilities over all intervals in K to obtain the likelihood $gc_prob_{c,b}$ that contig c originates from a molecule with GC content in interval b .

Penalty term. For contig c and interval b , the MILP uses penalty $gc_pen_{c,b}$, which is incurred if c is part of a plasmid whose GC content belongs to b . This penalty is computed as

$$gc_pen_{c,b} = \max_{b' \in K} \{gc_prob_{c,b'}\} - gc_prob_{c,b}.$$

The penalty is thus zero for the interval b where the likelihood of c achieves maximum. The penalty increases with the gap between the proposed GC content interval b and the most likely interval for contig c . For very short contigs, probabilities of different GC content intervals are closer to each

other, and thus the resulting penalties are smaller compared to longer contigs.

Pre-defining GC content intervals. The GC content intervals considered were pre-determined prior to running experiments based on an analysis of the reference dataset. The GC content of most plasmids and chromosomes in the reference dataset was observed to be between 0.4 and 0.6. The interval $[0.4, 0.6]$ was subdivided into 4 equal intervals resulting in $k = 6$ intervals $K = \{[0, 0.4], [0.4, 0.45], [0.45, 0.5], [0.5, 0.55], [0.55, 0.6], [0.6, 1]\}$.

3 Experimental results

3.1 Datasets and tools

We evaluated the following methods: our new tool PlasBin-flow, HyAsP (14), MOB-recon (18), plasmidSPAdes (2) and gplas (4). PlasmidSPAdes and gplas are de-novo binning methods, while MOB-recon, HyASP, and PlasBin-flow rely on a reference database of plasmid genes.

Our dataset consists of 133 bacterial genomes and 377 plasmids from a collection of real bacterial isolates with closed genome assemblies compiled by Robertson and Nash (18). To simulate the use of a realistic reference database, we split our data into a reference set and a test set; the samples released before 19 December 2015 were used to build the reference database and those released after that date formed the test set. The reference set consists of 67 bacterial isolates with 230 plasmids containing 10685 plasmid genes.

The remaining test set consists of 66 samples with 147 plasmids. For the test set, Illumina sequencing data was re-assembled using Unicycler to provide contigs and assembly graphs, except in case of plasmidSPAdes. PlasmidSPAdes takes raw reads as input and constructs its own assembly graph using the SPAdes assembler. Gplas requires the use of mlplasmids (3) to compute the probability of contigs to originate from a plasmid. Since mlplasmids is a species-specific tool, it currently supports only four species, three of which feature in our test dataset (*Escherichia coli*, *Enterococcus faecium*, and *Klebsiella pneumoniae*). From the test dataset, 39 samples with 88 plasmids belong from these three species.

3.2 Results

To evaluate predictions of individual tools on a particular testing sample, we need to compare them to the ground truth determined from annotated finished assembly of the sample as given by Robertson and Nash (18). We consider two sets of measures: *weighted* or *base-pair level* statistics, which consider accuracy of bin predictions weighted by the lengths of individual contigs (effectively counting how many base pairs were involved in correct predictions), and *unweighted* or *contig-level* statistics, which only consider accuracy of predictions for individual contigs, disregarding contig lengths.

Each short-read contig in the isolate assembly is mapped against the plasmids from the corresponding finished assembly (ground truth) using BLAST+(6). Matches that span less than 95% of the contig length are discarded. Thus for each bin u predicted by a particular tool, we have a set of contigs P_u included in the bin u , and for each ground-truth plasmid v , we have a set of contigs T_v that match (via BLAST+) to the ground-truth plasmid v . It should be noted that plasmidSPAdes is run on a different set of contigs than the other 5 methods. Thus, the sets of contigs representing the ground truth plasmids are different in case of plasmidSPAdes.

For the two sets of contigs P_u and T_v , we define $\text{overlap}(P_u, T_v)$ as either $|P_u \cap T_v|$ (in case of unweighted statistics) or the cumulative size in base pairs of all contigs in $P_u \cap T_v$ (in case of weighted statistics). Similarly, we define $\text{size}(X)$ for a set of contigs X as either $|X|$ (in case

Table 1. **Comparison of tool accuracy on 39 bacterial isolates.** The table shows mean values of precision, recall and F1 statistics over 39 testing samples from the three species supported by gplas

Tool	Basepair level (weighted)			Contig level (unweighted)		
	Precision	Recall	F1	Precision	Recall	F1
PlasBin-flow	0.74	0.69	0.66	0.77	0.50	0.54
PlasBin	0.58	0.58	0.56	0.62	0.61	0.59
HyAsP	0.73	0.46	0.53	0.74	0.40	0.49
plasmidSPAdes	0.52	0.72	0.54	0.53	0.84	0.58
MOB-recon	0.82	0.87	0.83	0.80	0.33	0.44
gplas	0.84	0.70	0.73	0.80	0.22	0.33

of unweighted statistics) or the cumulative size in base pairs of all contigs in X (in case of weighted statistics).

In order to compute the *precision*, we assign to each predicted plasmid bin v one of the true plasmids $f(v)$ such that $\text{overlap}(P_v, T_{f(v)})$ is maximized. The precision is then computed as $\text{overlap}(P_v, T_{f(v)})/\text{size}(P_v)$. Similarly, to compute the *recall*, we assign to each true plasmid u one of the predicted plasmid bins $g(u)$ such that $\text{overlap}(P_{g(u)}, T_u)$ is maximized. The recall is then computed as $\text{overlap}(P_{g(u)}, T_u)/\text{size}(T_u)$.

For a sample with a set \mathcal{P} of predicted plasmid bins and \mathcal{T} of true plasmids, the *mean precision* for the sample is then computed as:

$$\frac{\sum_{v \in \mathcal{P}} \text{overlap}(P_v, T_{f(v)})}{\sum_{v \in \mathcal{P}} \text{size}(P_v)}$$

The *mean recall* for the sample is computed as:

$$\frac{\sum_{u \in \mathcal{T}} \text{overlap}(P_{g(u)}, T_u)}{\sum_{u \in \mathcal{T}} \text{size}(T_u)}$$

The F1 measure for a sample is computed as a harmonic mean of mean precision and mean recall for that sample. For the set of 39 samples supported by gplas, we only consider contigs of length more than 1000 bp for our evaluation to be similar to gplas methodology (4). All contigs are considered in the evaluation for the remaining 27 samples.

Table 1 compares both unweighted and weighted accuracy measures to other tools averaged over the 39 testing samples, whereas figures 1 and 2 show the full distribution of these measures. In case of weighted base-pair level measures, the accuracy of PlasBin-flow is lags behind gplas as well as MOB-recon, which ranks as the best tool. PlasBin-flow show significant improvement over its predecessor PlasBin and also has comparable recall to gplas.

Unweighted contig level accuracy is a challenging measure, since short contigs have a higher contribution to these measures. Depending on the exact down-stream application, correct binning of shorter contigs may or may not be important. Shorter contigs are more difficult to classify and bin because corresponding sequence-based statistics have a higher variance and homology-based statistics are often non-informative. When considering unweighted statistics, PlasBin-flow shows comparably high precision to MOB-recon and gplas - both of which are especially selective in their choice of contigs. PlasmidSPAdes is the tool with the highest recall, but it suffers from many false predictions (low precision).

Table 2 compares both unweighted and weighted accuracy measures for the other 27 testing samples. Figures 3 and 4 show the full distribution of these measures. In case of weighted base-pair level measures, PlasBin-flow shows the best accuracy. It ranks second-best in terms of recall behind MOB-recon but shows a better combined F1 accuracy. A similar trend is observed in unweighted contig level accuracy measures. The performance of PlasBin-flow, HyAsP and MOB-recon is comparable, with PlasBin-flow having a slight edge in terms of precision.

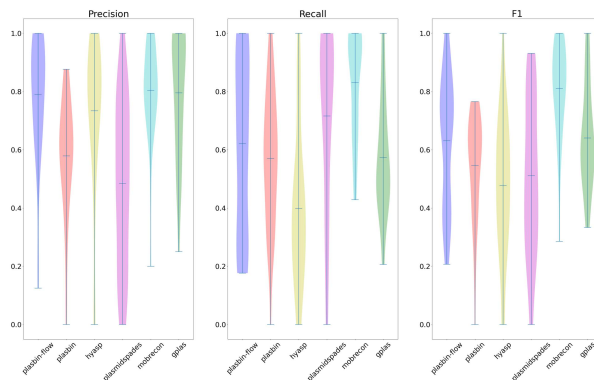


Fig. 1: Distribution of the unweighted precision, recall and F1-score statistics for samples from three species supported by gplas

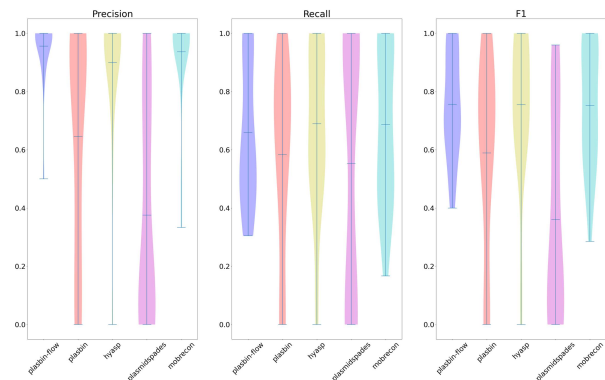


Fig. 3: Distribution of the unweighted precision, recall and F1-score statistics for remaining 27 samples.

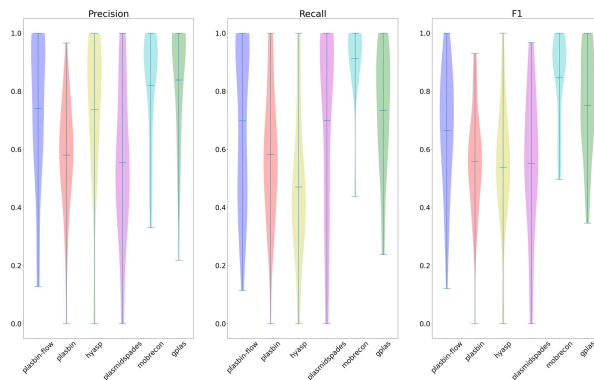


Fig. 2: Distribution of the weighted precision, recall and F1-score statistics for samples from three species supported by gplas

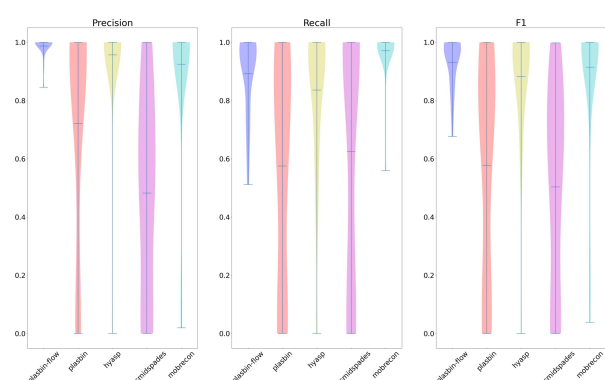


Fig. 4: Distribution of the weighted precision, recall and F1-score statistics for remaining 27 samples.

Table 2. **Comparison of tool accuracy on 27 bacterial isolates.** The table shows mean values of precision, recall and F1 statistics for the remaining 27 testing samples from species not supported by gplas

Tool	Basepair level (weighted)			Contig level (unweighted)		
	Precision	Recall	F1	Precision	Recall	F1
PlasBin-flow	0.98	0.89	0.93	0.96	0.66	0.76
PlasBin	0.72	0.58	0.58	0.65	0.58	0.59
HyAsP	0.96	0.84	0.88	0.90	0.69	0.75
plasmidSPAdes	0.48	0.63	0.50	0.38	0.55	0.36
MOB-recon	0.92	0.97	0.91	0.94	0.69	0.75

Table 3. **Running time statistics comparison.** Median, mean, minimum, and maximum running time (in minutes) over the testing data set

Tool	Minimum	Maximum	Median	Mean
PlasBin-flow	5	120	6	6
PlasBin	5	473	22	117
HyAsP	1	19	3	4
MOB-recon	3	15	5	8
gplas	2	12	3	4
plasmidSPAdes	4	363	16	43

3.3 Computational footprint

We ran our experiments on a standard laptop computer, with a quad-core processor and 16GB of memory. PlasBin-flow relies on an MILP, that typically requires large computing resources (time and memory). However, PlasBin-flow showed significant improvement over PlasBin. In fact, in terms of median and mean running times, the method was comparable to other methods such as HyAsP, MOB-recon and gplas. Excluding outliers, PlasBin-flow usually completed within a few minutes. Over all considered samples, PlasBin-flow had a maximum running time close to 2 hours, with a memory footprint close to 6GB (maximum 10GB). The details have provided in Table 3.

4 Conclusion

In this work, we presented PlasBin-flow, an MILP algorithm that uses network flows for grouping contigs into putative plasmid bins. We compared the results of PlasBin-flow against other state-of-the-art plasmid binning methods: HyAsP, plasmidSPAdes, MOB-recon and gplas using two sets of metrics: weighted (basepair level) and unweighted (contig level). In both cases, PlasBin-flow consistently performed reasonably well in precision statistics. For samples belonging to the *Escherichia coli*, *Enterococcus faecium*, and *Klebsiella pneumoniae*, PlasBin-flow ranked behind MOB-recon and gplas in the average F1 scores. Gplas in particular had a distinct advantage over PlasBin-flow due to its use of species specific training sets. The performance of PlasBin-flow was markedly better on

samples belonging to other species. PlasBin-flow was especially useful in binning contigs of short length, which are typically difficult to bin correctly.

The objective function in PlasBin-flow is significantly changed from that used in PlasBin. The uniformity of %GC content along the contigs in a putative plasmid bin is achieved using a probabilistic model. The use of %GC content penalties discourages the choice of contigs disparate %GC content to be selected in the same bin. Another salient features of this method is the use of flow values as a proxy for read coverage used in a solution. The conditions necessary to maintain a consistent flow ensure the uniform read coverage in a plasmid bin without the need to explicitly compute the mean read coverage of the bin.

In some instances, PlasBin-flow shows low recall. One possible explanation for low recall is that a true plasmid may consist of multiple bins output by PlasBin-flow. This is typically expected when the contigs connecting distinct plasmid bins have low gene density. Gplas which performs well in this regard, is able to avoid this potential pitfall due to its reliance on coverage rather than just gene density.

The performance of PlasBin-flow shows that there is definite room for improvement. Multiple avenues can be explored in this regard. Currently, all terms in the objective function have equal importance. However, PlasBin-flow can be further investigated by assigning different weights to each term to factor in their relative importance. Our MILP formulation can also accommodate other contig features in its objective function such as including a probabilistic value from plasmid contig classification methods such as mlplasmids. Guided by the results of classification methods, the assembly graph can be pruned before the binning stage.

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