Method

Genealogical inference and more flexible sequence clustering using iterative-PopPUNK

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Bacterial genome data are accumulating at an unprecedented speed due to the routine use of sequencing in clinical diagnoses, public health surveillance, and population genetics studies. Genealogical reconstruction is fundamental to many of these uses; however, inferring genealogy from large-scale genome data sets quickly, accurately, and flexibly is still a challenge. Here, we extend an alignment- and annotation-free method, PopPUNK, to increase its flexibility and interpretability across data sets. Our method, iterative-PopPUNK, rapidly produces multiple consistent cluster assignments across a range of sequence identities. By constructing a partially resolved genealogical tree with respect to these clusters, users can select a resolution most appropriate for their needs. We showed the accuracy of clusters at all levels of similarity and genealogical inference of iterative-PopPUNK based on simulated data and obtained phylogenetically concordant results in real data sets from seven bacterial species. Using two example sets of Escherichia/Shigella and Vibrio parahaemolyticus genomes, we show that iterative-PopPUNK can achieve cluster resolutions ranging from phylogroup down to sequence typing (ST). The iterative-PopPUNK algorithm is implemented in the “PopPUNK_iterate” program, available as part of the PopPUNK package.

[Supplemental material is available for this article.]

A key step in many analyses of collections of bacterial genome sequences is to understand how the sequences are related to each other. For example, to make a clinical diagnosis, a key question is which previously characterized organisms are similar to strains present in the patient sample (Gardy and Loman 2018; Gu et al. 2019). To identify transmission chains, it is necessary to establish which of the sequenced strains share very recent common ancestors, indicating a shared source (Croucher and Didelot 2015; Sintchenko and Holmes 2015). To map genotype-phenotype relationships, it is often helpful to delineate lineages. Organisms from the same lineage have similar sequence content and thus are likely to have similar phenotypes (Falush 2016; Power et al. 2017; Binda et al. 2020). To accurately reconstruct the evolutionary history of a species, it is necessary to obtain a sample of organisms that is broadly representative of the species’ diversity as a whole and is not biased towards specific lineages, for example by containing only multiple closely related sequences from the same environmental source.

A “gold standard” for establishing genetic relationships between strains is through genealogical reconstruction. Any given sample of bacteria are related to each other by a tree, with each node of the tree representing the most recent common ancestor (MRCA) of the strains below it in the genealogy. The genealogical history reflects the clonal relationships between strains; it is unique and unambiguous. Genome sequence data provides partial information on the topology of this tree, its MRCA, and where and when that MRCA existed. In practice, the information that genome sequences provide on genealogical relationships is both incomplete and time consuming to extract. Moreover, a full genealogy is rarely necessary for most practical uses of genetic relationships. This has led to the development of methods that attempt to identify clusters of closely related organisms. A simple interpretation of each cluster is that it corresponds to the set of strains descended from a single node of the genealogical tree. Among these possible methods, PopPUNK is notable because it works from raw sequence data and does not require a genome alignment (Lees et al. 2019). This means that the clustering can be obtained with significantly less computational resources and is much faster than methods based on sequence alignments. Moreover, PopPUNK is easily extendable, as new batches of genomes can be integrated into an existing database without needing to recalculate all pairwise distances and reperform the model fitting step.

PopPUNK avoids the need for an alignment by creating a k-mer profile for each strain, using both short and long k-mer lengths, which is computationally much faster than any form of alignment thanks to highly efficient hashing algorithms. It compares the k-mer profiles of different strains based on their Jaccard similarity. Gene presence-absence polymorphisms create a profile of k-mer similarities which is distinct from that because of single-nucleotide polymorphisms (SNPs), making it possible to estimate distances for core and accessory genome separately by fitting the pattern of distance profiles across multiple k-mer lengths.

Two machine learning algorithms, Bayesian Gaussian mixture model (BGMM) and hierarchical density-based spatial clustering of applications with noise (HDBSCAN), are implemented in PopPUNK to identify components. The component closest to
the origin of the scatterplot of core and accessory distances is defined as “within strain,” corresponding to closely related strains. The other one is defined as “between strain,” corresponding to distant strains. Then these within-strain distances are used as edges to link samples and create PopPUNK clusters. The PopPUNK algorithm identifies a single level of clusters, attempting to find an “optimal” choice. However, the biological interpretation of this optimal clustering is not clear because the genealogical tree underlying the relationships between strains is hierarchical rather than having a single level. Furthermore, in practice, multiple levels of clusters with different genetic similarities are often required to achieve different purposes. For example, in the disease outbreak scenario, epidemiologists require clusters of very closely related or indistinguishable strains to help identify and track outbreaks. For routine surveillance and subtyping, clusters with higher diversity are required to represent strains from a sequence type or clonal group. Population geneticists usually require clusters of population or subpopulation level to conduct species-level-level studies.

Two state-of-the-art pipelines for large-scale genome data sets that allow multilevel clustering are Fastbaps (Tonkin-Hill et al. 2019) and HierCC, as implemented in Enterobase (Zhou et al. 2021). Fastbaps is based on a core genome alignment, which is slow to generate for data sets of even moderate sizes. HierCC requires an appropriate curated database of genes and alleles or a stable strain nomenclature and cannot be the general solution for most bacterial species.

Here we present iterative-PopPUNK, an extension of the PopPUNK software that allows estimation of a partially resolved genealogical tree and provides clustering consistent with that tree at different levels of sequence similarity. This enhancement provides greater flexibility than the original PopPUNK algorithm, allowing the fineness of the clustering to be adjusted according to the purpose at hand. Moreover, the clusters can be interpreted more readily, by reference to the full (partially resolved) genealogy.

Results

Overview of Iterative-PopPUNK

Iterative-PopPUNK allows estimation of a partially resolved genealogical tree and provides clustering consistent with that tree at different levels of sequence similarity. This enhancement provides greater flexibility than the original PopPUNK algorithm, allowing the fineness of the clustering to be adjusted according to the purpose at hand. Moreover, the clusters can be interpreted more readily, by reference to the full (partially resolved) genealogy. The speed of iterative-PopPUNK is similar to PopPUNK, which can complete the analysis of thousands of bacterial genomes in a few minutes, and can scale even to very large-scale genome data sets (e.g., >100,000). Using simulated data, we show the accuracy of the clusters from iterative-PopPUNK at all output levels of similarity. Furthermore, we show that for real data, the produced clusters are normally highly concordant with those obtained by phylogenetic methods, but without requiring an alignment.

An overview of iterative-PopPUNK pipeline is shown in Figure 1A. Six steps are required to implement the algorithm of iterative-PopPUNK. The first two steps proceed as usual using the original PopPUNK algorithm: (1) database construction and distance calculation and (2) model fitting using original PopPUNK algorithm. Iterative-PopPUNK then performs four additional steps to achieve multilevel clustering and generate clusters under the given similarity cutoffs.

Step 3. Multilevel clustering by moving decision boundary iteratively

After an initial model has been fitted, PopPUNK identifies two fitted components representing “within-strain” comparisons between closely related isolates and “between-strain.” However, unlike the original PopPUNK algorithm, which identifies a single boundary for defining a “best” cluster assignment by optimizing a heuristic (the network score), iterative-PopPUNK instead ignores the network score, and places the decision boundary at multiple equally spaced positions perpendicular to the line connecting within- and between-strain components to create multilevel cluster assignments. We have found using simulated data that the clustering accuracy remains high even when the network score is low and we are interested in obtaining clustering at multiple hierarchical levels.

The first viable boundary point is where at least one nontrivial cluster (containing at least two samples) is formed, corresponding to the two most closely related sequences in the data set. The number of boundaries, and therefore the number of starting sets of clusters, can be set by the user. Based on simulation data, we found that the number of estimated clusters increases as the number of decision boundary line increases, and basically reached a plateau when the number of decision boundary line is 30 (Supplemental Fig. S1). To balance the speed and resolution of iterative-PopPUNK, we set 30 as the default cutoff. Users can choose to increase this value if they want a higher resolution.

Step 4. Quality control (QC) of clusters obtained at different levels

Quality control (QC) is performed to remove any duplicated or inconsistent clusters. The clusters obtained under the first viable boundary point are directly defined as QC-passed clusters because of their high accuracy based on the evaluation using simulation data. A new cluster assignment $C_i$ (which is a set of samples) at boundary point $j$ was compared with QC-passed clusters $C_j$ at the previous boundary $i$, and is considered QC-passed if and only if it meets one of the following conditions, as shown in the conceptual outline in Figure 1B:

1. $C_j$ contains all of the isolates from a QC-passed cluster $C_i$ ($C_j \subseteq C_i$, condition 1);
2. $C_j$ is unique and contains none of the isolates from any previous $C_i$ ($C_j \cap C_i = \emptyset$, condition 2);
3. $C_j$ is a strict subset of one of the previous QC-passed clusters $C_i$ ($C_j \subseteq C_i$, condition 3).

By iterating through all pairs of clusters between the current step and next step forward (as clusters increase in size; equivalently the boundary moves upwards), a new QC-passed $C_j$ was added into the list $C_j$ at the next round, until the QC of all the clusters is completed.

Step 5. Partially resolved genealogical tree construction and average core distance calculation

A partially resolved genealogical tree is then constructed based on QC-passed clusters by nesting clusters within larger ones (Fig. 1D). The algorithm we used to do this is as follows:

1. Place all strains at the root as “cluster zero” and sort all clusters in a descending order according to their size;
2. Run through the QC-passed clusters, from largest to smallest. For each one:
a. Look through all clusters which have been added to the tree and find the smallest cluster which is a superset of the cluster being tested;
b. Nest the new cluster as a child under this parent cluster;
c. Remove the strains in the child cluster from the parent cluster;
d. For the remaining strains (which may be at parent nodes, or the tips) add in a child node;
e. For each of the nodes in the tree, we calculate the average core distances between the strains under this node and set its branch length relative to root accordingly.

Figure 1. Workflow for developing iterative-PopPUNK. (A) Steps for designing iterative-PopPUNK. After inputting sequence data, PopPUNK creates a local sketching database, which is further used to estimate clusters. QC steps for removing inconsistent clusters are shown in the orange box. The green box shows how to nest these estimated clusters into an iterative-PopPUNK tree. Methods for choosing the final set of clusters are presented in the blue box. (B) QC algorithm. Three conditions for determining QC-passed clusters are described in detail in Methods, which in short are as follows: (1) The new clusters contain all of the isolates from previous clusters; (2) the new cluster is unique and contains none of the items from previous clusters; (3) the new cluster is a strict subset of one of the previous clusters. (C) Algorithm for cutting iterative-PopPUNK tree. The goal is to choose the closest node to the cutoff line but with a smaller value. In this example, the cluster (node) annotated using blue color has the maximum value of average core distances (0.6). The red dashed line shows the cutoff is 50% of the MACD (0.3). Therefore, the node with an ACD lower than and closest to 0.3 will be selected. (D) Hierarchical tree assembly. The dashed lines indicated these potential branches during the tree assembly process.
Step 6. Choosing clusters under given similarity cutoffs

Considering the large difference in the genetic diversity of different bacterial species, we designed two types of relative cutoffs to use when quantifying genetic similarity within a species: (1) \( P_{\text{max-cluster-dist}} \), the percentage of maximum value of the average core distance (MACD) of all the iterative-PopPUNK clusters (extracted from local PopPUNK distance database). The cluster with maximum value of the MACD has the highest genomic diversity. We use the percentage of this value as a cutoff to decide which other clusters are selected. (2) \( P_{\text{bet-mean-dist}} \), the percentage of mean pairwise core distance between all isolates (extracted from initial model fitting results).

When one of these cutoffs is provided, iterative-PopPUNK searches all the nodes except root node in the partially resolved tree and extracts the relative similarity values (\( P_{\text{max-cluster-dist}} \) or \( P_{\text{bet-mean-dist}} \)) of all the samples under each node (NP) and its parent node (N + 1P). If a node meets the condition NP \( \leq \) cutoff < N + 1P, all the child strains under this node are assigned into a cluster (Fig. 1C). The remaining unassigned strains were defined as non-clustered strains.

Evaluation of iterative-PopPUNK using simulation data

To test the performance of iterative-PopPUNK, we generated seven data sets of simulated genomes in which the true genealogy is known. We generated 100 genomes with a length of 1000 kb per data set and their corresponding genealogical trees using FastSimBac (De Maio and Wilson 2017), representing bacteria data set and their corresponding genealogical trees using known. We generated 100 genomes with a length of 1000 kb per data set and their corresponding genealogical trees using FastSimBac (De Maio and Wilson 2017), representing bacteria data set and their corresponding genealogical trees using known. We generated 100 genomes with a length of 1000 kb per data set and their corresponding genealogical trees using FastSimBac (De Maio and Wilson 2017), representing bacteria data set and their corresponding genealogical trees using known.

Application of iterative-PopPUNK to real data shows consistency with phylogenetic results

We applied iterative-PopPUNK with default settings to analyze the genomic data sets of eight bacterial species with different mutation and recombination rates (Supplemental Table S1) and obtained the cluster assignments under six relative similarity cutoffs (\( P_{\text{max-cluster-dist}} \): 1%–99%). In addition, we performed core-genome alignments for these data sets, and constructed maximum-likelihood (ML) phylogenetic trees based on core-genome SNPs. To measure the consistency between iterative-PopPUNK and alignment-based method, we mapped the clusters obtained under different cutoffs to the corresponding ML trees, and calculated matching accuracy (i.e., the proportion of matched clusters to all clusters) in the same manner as described above.

We collated the data set for Escherichia coli by subsampling 500 genomes from 50 largest lineages (10 per lineage), which together represent the entire genomic diversity of human-isolated E. coli (Horesh et al. 2021). The entire, fully sampled, data set has previously been analyzed by PopPUNK, and these 50 lineages were defined based on PopPUNK clusters. An ML phylogenetic tree based on core-genome SNPs was constructed for this same set of E. coli genomes to evaluate the matching rate of sequence typing (ST) groups and this ML tree (Fig. 4A). The result showed that these 55 ST groups from E. coli genomes were poorly coordinated with the ML tree, as only 26 of them can exactly be matched to the tree nodes whereas the other 29 failed to find any corresponding nodes from the ML tree. The high mismatch rate between ST and the ML tree is because of our very strict calculating method: these strains from unmatched clusters were closely related in the simulated tree, although not fully clustered together. However, there was nevertheless a high matching rate between the ML tree and iterative-PopPUNK clusters, with 75% being completely congruent (Fig. 4B). If we look at the more permissive statistic of strain concordance (the total proportion of isolates that are assigned to one cluster by both methods), the matching rate is very high (91%) (454/500) (Supplemental Fig. S4).

Using a medium cutoff (\( P_{\text{max-cluster-dist}} \geq 20\% \)), iterative-PopPUNK estimated 42 clusters which exactly corresponded to 42 clusters using phylogenetic method with 20% SNP distance as cutoff (Fig. 4A). Under a large cutoff (\( P_{\text{max-cluster-dist}} = 50\% \)), iterative-PopPUNK cluster closely corresponded to strains from a phylogroup. Of these nine obtained clusters, iterative-PopPUNK failed to find phylogroup C, which appears to be a subclade of B1 according to the phylogenetic tree and so does not deserve a phylogroup label according to current whole-genome analysis method. The definition of phylogroup C relied instead on virulence genes (Moisenson et al. 2010; Clermont et al. 2011, 2013). Six perfectly corresponded to phylogroups (A, B2, E, F, Shigella flexneri and Shigella sonnei) and the other two corresponded to the
subclades of phylogroup D. This example highlighted the advantage of iterative-PopPUNK, which allows multilevel clustering in a unified way and users can select their “level” (e.g., sequence type or phylogroup level) according to their analytic purpose.

We also applied iterative-PopPUNK to analyze a “benchmark” data set of *Vibrio parahaemolyticus*, consisting of 2640 genomes, that had been previously analyzed using phylogenomic methods (Yang et al. 2022). This data set contains groups from clonal level (strains with ≤2500 SNPs) to outbreak level (within 1-mo clusters of strains with ≤6 SNPs) representing putative outbreaks. For this data set, iterative-PopPUNK succeeded in estimating 23 clusters that exactly corresponded to 23 pathogenic clonal groups (PCGs) described in Yang et al. (2022) (Fig. 5A; Supplemental Table S3). Two major PCGs, PCG3 and PCG189 (including 1768 and 273 pathogenic genomes, respectively), were extracted from this data set to explore iterative-PopPUNK’s ability in resolving more fine-scale structure of closely related isolates. The results showed that iterative-PopPUNK clustering assignments had high concordance with outbreak-level phylogenomic clusters, with strains concordance of 90% (1371/1535) and 95% (259/273). However, as

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**Figure 2.** The performance of iterative-PopPUNK on simulated data. The chosen simulated data set for A and B has a recombination rate \( r = 0.001 \) and mutation rate \( \mu = 0.01 \). (A) A genealogical tree with nine nodes (clusters) estimated by default PopPUNK. (B) The same genealogical tree with 69 clusters estimated by iterative-PopPUNK, among which 68 can be totally matched with genealogical tree nodes although one of them cannot be matched, with strains in the unmatched cluster shown in blue. (C) A line graph shows the average number of estimated clusters by PopPUNK and by iterative-PopPUNK. The solid lines present the results for iterative-PopPUNK, and the dash lines are for default PopPUNK. (D) The accuracy results for both default PopPUNK and iterative-PopPUNK. The accuracy is calculated from the total number of matched nodes divided by the total number of estimated clusters. (E) Robinson-Foulds distances between true trees and iterative-PopPUNK trees. \( r/\mu \) stands for recombination rate divided by mutation rate.
expected, the number of estimated “outbreak-level” iterative-PopPUNK clusters is less than that of phylogenomic clusters, because multiple phylogenomic clusters were assigned into one iterative-PopPUNK cluster. The number of clusters can be used as a rough quantification of the resolution. Using iterative-PopPUNK, 80 clusters were identified for PCG3 and 25 clusters for PCG189.

The concordance of iterative-PopPUNK clustering with the identified outbreak groups was 60% (48/80) for PCG3 and 80% (20/25) for PCG189. In both cases, the resolution is much finer than clonal level (Fig. 5B,C; Supplemental Table S3).

The data sets of the other six species were assembled from publicly available genomes in NCBI RefSeq database. Iterative-
Figure 4. Population structure of *E. coli* at different iterative-PopPUNK levels. (A) Comparison of PopPUNK clusters and a phylogenetic tree. The outer solid ring represents 9 PopPUNK clusters at 50% of the MACD, corresponding to nine *E. coli* phylogroups. The middle ring contains 42 clusters at 20% of the MACD, corresponding to 42 clusters from phylogenetic tree at 20% SNP distance. The inner solid ring shows 94 PopPUNK clusters at 7.9% of the MACD, corresponding to 92 *E. coli* ST groups. Colors in the tree leaves and branches are used to present these 92 STs and nine phylogroups, respectively. The parameters used for estimating PopPUNK clusters were all set to default with BGMM model (two components). (B) The distribution of matching rate and the number of clusters being estimated at multiple levels for this set of *E. coli* genomes. The tree was plotted using iTOL (https://itol.embl.de/). The comparison of PopPUNK clusters and an iterative-PopPUNK tree is available as Supplemental Figure S2. The original figure with high resolution is available as Supplemental Figure S3.
PopPUNK did not produce useful results for two species with low genetic diversity, namely *Mycobacterium tuberculosis* and *Bacillus anthracis*. For the other four species, namely *Helicobacter pylori*, *Vibrio parahaemolyticus*, *Klebsiella pneumoniae*, and *Bacillus cereus*, the average matching rates between phylogenetic methods and iterative-PopPUNK were 85%, with higher rates in species with high recombination rates such as *Vibrio parahaemolyticus*. The matching rates did not show clear differences between different similarity cutoffs (Fig. 6; Supplemental Figs. S5–S8).

**Discussion**

Over the last two decades, genome data have been accumulating at an unprecedented speed. For some common pathogens such as *Salmonella enterica* and *Escherichia coli*, there are now more than 200,000 genomes in public databases such as EnteroBase (Alikhan et al. 2018) and Genometracker (Brown et al. 2021). Analyzing large-scale genome data sets is still challenging, and alignment-free methods such as PopPUNK are one of the best solutions to address this. PopPUNK was initially designed to identify a single level of clusters by optimizing a heuristic, but in practice different cluster resolutions are required, depending on the purpose. To solve this problem, we developed iterative-PopPUNK, which allows estimation of a partially resolved genealogical tree and enables clustering at different levels of sequence similarity. Iterative-PopPUNK inherits the advantages of PopPUNK, which is fast and consumes few computational resources, and it is easily extendable when new data needs to be added (Lees et al. 2019).

On this basis, iterative-PopPUNK provides greater flexibility and allows the resolution of the clustering to be adjusted according to the specific purpose. However, default PopPUNK may be a better solution for analyzing large data sets in which a single level of clustering needs to be imposed, for example to define “nonredundant” genomes, that are representative of the species as a whole. In this case, the network score provides a mathematically founded method for choosing a “best” clustering, which is a pragmatic way of choosing a subset of isolates that best represent the overall diversity of the data set in a more compact form.

The traditional method for establishing the relationships between bacterial strains is phylogenetic analysis, in which SNPs within a DNA sequence alignment are used to reconstruct a tree. These methods usually require a reference genome, and inappropriate reference selection may introduce bias. This approach is also not typically scalable to large-scale genome data sets (e.g., >10,000) and runtimes can become prohibitively long, even with generously provisioned computational resources. For example, on a high-end workstation a phylogenomic analysis of ~10,000 *Klebsiella pneumoniae* genomes took 14 d and required a maximum of ~500 GB memory. Furthermore, phylogenetic methods may be inaccurate in the presence of high homologous recombination rates, because individual recombination events can change multiple nucleotide positions within the alignment, making the accumulation of polymorphisms over time less clock-like. More sophisticated approaches that differentiate between recombination and mutation events can be even more time consuming.
Another kind of method for classifying strains is based on gene-based subtyping information, including the classic multilocus sequence typing (MLST) which is based on several housekeeping genes (Maiden et al. 1998), and more recently developed core-genome MLST (cg-MLST) or whole-genome MLST (wg-MLST). Compared with genome-wide variation such as SNPs, subtyping information is typically much more parsimonious at cost of the loss of some phylogenetic resolution, making the analyses scalable to large data sets. HierCC is a state-of-the-art pipeline for multilevel clustering analysis of large-scale data sets based on cg-MLST (Zhou et al. 2021). HierCC hierarchical clusters (HC) can be used for multilevel strain assignment. For example, in Salmonella, HC900 (<900 allele difference) and HC2500 (<2500 allele difference) represent strains from the same sequence type complex and species/subspecies, respectively. However, performing gene-by-gene subtyping-based clustering first requires an appropriate curated database of

Figure 6. The distribution of matching rate and the number of clusters being estimated at multiple levels. A cluster is defined as a group with two or more strains (so nonclustered strains are not considered in counting the number of clusters).
genes and a stable strain nomenclature. These have not yet been constructed for most bacterial species, especially for nonpathogenic species, and require ongoing maintenance, making this method unlikely to be the general solution for microbes. Both PopPUNK and iterative-PopPUNK do not require a curated set of reference genomes and allow to train their own model based on any collection of genomes. The PopPUNK package does provide curated databases of several species, which users can use for quickly clustering new genomes.

The current version of iterative-PopPUNK has two main limitations. First, it does not perform well on species with low genetic diversity because boundaries tend to connect clusters together in a nontransitive manner. This can be solved by making a neighbor joining tree within each strain using the core distances calculated by PopPUNK. Second, the range of the levels of clusters obtained depends on the model fit of within- and between-strains components. Sometimes, only a very limited number of levels can be obtained, especially when using the second type of relative similarity cutoff $P_{\text{bet-mean-dist}}$.

With the wide use of whole genome sequencing and rapid accumulation of genome sequences, large-scale genome data set analysis may become normal in the near future. However, there is currently no ideal multilevel clustering method for large-scale data set analysis. Iterative-PopPUNK is alignment-free, highly efficient even for very large-scale data sets, and does not rely on a curated set of reference genomes database and nomenclature, which can solve the limitations of the above methods, and we believe it can be a solution for fast multilevel clustering of most bacterial species. However, where data size and computation time is not an issue, methods that use alignments and evolutionary models to reconstruct the individual genetic events involved in strain divergence capture more information and have the potential to achieve slightly higher clustering resolution.

Methods

Implementation of iterative-PopPUNK algorithm

Iterative-PopPUNK procedure is implemented in the “poppunk_iterate” program installed as part of PopPUNK package. Source code is available on GitHub (see Software availability) and can be installed through conda, pip, or manually. Users can run a multilevel clustering procedure using the flags “--fit-model refine --multi-boundary” in the PopPUNK software. The subsequent QC procedure is implemented in the “poppunk_iterate” program and uses output from multilevel clustering step as input.

Iterative-PopPUNK cluster accuracy evaluation using simulation data

We performed simulation using FastSimBac (De Maio and Wilson 2017) to test the accuracy of iterative-PopPUNK clusters. Fifteen parameter combinations were used in simulation to represent bacteria with different mutation and recombination rates (Supplemental Table S1). Simulation outputs were converted to genome sequences in FASTA format for further iterative-PopPUNK analysis using integrated simulator, msformatter and ms2dbsa (available from http://guanine.evolbio.mpg.de/bioBox), and the corresponding genealogical trees were extracted for comparison with iterative-PopPUNK clusters.

To analyze the simulated genomes, we set the maximum accessory distance parameter of iterative-PopPUNK to 1.0 to retain all isolates. In the first “model fitting” step, BGMM from original PopPUNK was chosen to fit the distribution of core and accessory distances. Two components (K = 2), one for within-strain and the other one for between-strain were applied. The component closest to the origin is defined as within-strain, in which all these shorter pairwise distances are from one cluster; although the between-strain distances refer to these larger ones that may be from different clusters. The other parameters were set to default values ($k = 15–29$ in steps of 4; sketch size $10^5$), which have been shown to work well across most bacterial species.

Based on the node information of simulated genealogical trees, we assign simulated strains into true tree clusters. Each iterative-PopPUNK cluster was compared with the true tree clusters using the ete3 toolkit (Huerta-Cepas et al. 2016). If the strain assignment of an iterative-PopPUNK cluster is identical to one of these simulated tree clusters, it is defined as a match iterative-PopPUNK cluster, and associated nodes were labeled using green/red faces alternatively in the simulated tree; otherwise, it is unmatched, and the leaves would be annotated using a certain color for each unmatched cluster. Accuracy is defined as the proportion of matched clusters to all iterative-PopPUNK clusters. The method of Robinson and Foulds (1981) was applied to compare the simulated genealogical tree and the inferred iterative-PopPUNK tree using the ete3 toolkit in Python.

Application of iterative-PopPUNK in real data

The publicly available, assembled genomes of seven species were downloaded from the National Center for Biotechnology Information (NCBI) Reference sequence (RefSeq; https://www.ncbi.nlm.nih.gov/ refseq/) database based on the taxonomy identifier (Supplemental Table S2). Two steps of genome sequences QC were performed. First, we used CheckM (Parks et al. 2015) to assess the genome quality and only genome sequences with completeness >90% and contamination <5% were kept in further analysis. Second, we set 70% as whole genome coverage threshold to remove low-quality genomes or outlier samples that may not be part of the species. After QC, the same iterative-PopPUNK pipeline used for simulation data were applied to real data analysis. The 500 E. coli genomes were collected from Horesh et al. (2021). Additionally, a separate set of benchmark data comprising 2640 Vibrio parahaemolyticus genomes (distinct from those obtained from the NCBI database) were obtained from Yang et al. (2022). The original data set consisted of 3460 genomes, but singletons and nonpathogenic isolates were removed for this analysis. The Sankey diagrams were plotted using Plotly graphing library for Python (available from https://github.com/plotly/plotly.py).

QC-passed genome sequences were further aligned to the reference genome of each species (Supplemental Table S2) using MUMmer (Kurtz et al. 2004) to generate whole genome alignments. Core-genome (regions present in >99% isolates) SNPs were called using snap-sites (Page et al. 2016) based on the alignments. ML phylogenetic trees were constructed using FastTree (Price et al. 2010) based on core-genome SNPs. We adopted the method used in iterative-PopPUNK cluster accuracy evaluation to calculate the matching rate of iterative-PopPUNK clusters and ML trees, by replacing the simulated trees to ML trees. To show how the number of estimated clusters changes for different species, we used 1% MACD as a gap to move the cutoff from 1% MACD to 99% MACD. For practical usage of iterative-PopPUNK, users can adjust these cutoffs as needed.

Software availability

The source code and tutorial are available in Supplemental Code. The program “poppunk_iterate.py” is available on GitHub (https://github.com/bacpop/PopPUNK), installed as a part of
PopPUNK package. Documentation for tutorial can also be found at https://poppunk.readthedocs.io/en/latest/poppunk_iterate.html. Online examples from tutorial for each data set are listed in Supplemental Table S3.

Competing interest statement
The authors declare no competing interests.

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