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Efficient inference of large prokaryotic pangenomes with PanTA



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Abstract

Pangenome inference is an indispensable step in bacterial genomics, yet its scalability poses a challenge due to the rapid growth of genomic collections. This paper presents PanTA, a software package designed for constructing pangenomes of large bacterial datasets, showing unprecedented efficiency levels multiple times higher than existing tools. PanTA introduces a novel mechanism to construct the pangenome progressively without rebuilding the accumulated collection from scratch. The progressive mode is shown to consume orders of magnitude less computational resources than existing solutions in managing growing datasets. The software is open source and is publicly available at https://github.com/amromics/panta and at 10.6084/m9.figshare.23724705.

Background

Prokaryotic genomes are known for enormous intraspecific variability owing to great variation events such as horizontal gene transfers, differential gene losses, and gene duplication [1]. This led to the introduction of the pangenome concept as a methodology to investigate the diversity of bacterial genomes [2]. Since its inception in 2005, pangenome analysis has been an indispensable tool in microbial genomics studies [3] and has generated novel biological insights in bacterial population structures [4, 5], genetic diversity [6], niche adaptation [7], and genome assembly [8]. Pangenome studies have also been successfully applied into inferring the evolution of lineages of pandemic causing pathogens and identifying lineage-specific genetic features [9, 10], investigating genetic signatures associated with antimicrobial resistance [11], pan-reactome analyses [12], and therapeutic development including vaccine design [13] and novel drug discovery [14, 15].

To address the need for pangenome analysis, a plethora of computational tools have been developed to construct the pangenome of a collection of prokaryotic genomes. Notable examples include PGAP [16], PanOCT [17], Roary [18], BPGA [19], panX [20], MetaPGN [21], PIRATE [22], PPanGGOLiN [23], PEPPAN [24], and Panaroo [25].

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Advances in high-throughput sequencing technologies have recently enabled the exponential growth of microbial genomics data in public databases and in research laboratories around the world. The Genbank database stores hundreds of thousands of genomes for common bacterial species, and the numbers are fast-growing. While these resources contain rich sources of population genomics information, pangenome analysis has not been able to scale with the volume of the data. Most existing pangenome inference methods take days and require large amounts of memory that are typically beyond the capacity of a standard computer to construct the pangenome of just a few thousand isolates. In addition, the genomic databases are growing by nature, accumulating genomes of isolates collected and sequenced at different time points. There currently exists no efficient utility to update an existing pangenome when new genomes become available. In such cases, the pangenomes of the accumulated collection have to be constructed from scratch over and over, leading to the excessive burden of computational resources.

In order to address these challenges, we have developed PanTA, an efficient and scalable pangenome construction tool to keep up with the growth of bacterial genomics data sources. With vigorous computational experiments, we show that PanTA exhibits an unprecedented multiple-fold reduction in both running time and memory usage compared with the current state-of-the-art tools for building the pangenomes of large collections. Crucially, PanTA allows performing pangenome analysis progressively where batches of new samples can be added to an existing pangenome without the need to recompute the accumulated pangenome from scratch. The progressive mode can further reduce PanTA memory usage by half without affecting running time and pangenome accuracy. We also show that, PanTA in progressive mode consumed orders of magnitude less computational resource than existing solutions to manage the pangenomes of growing microbial datasets. Finally, we demonstrate the utility and practicality of PanTA by constructing the pangenome of the entire set of high quality *Escherichia coli* genomes that have been deposited into RefSeq database to date on a laptop computer.

Results

Overview of the pipeline

PanTA is developed with the aim to build the pangenome of a large collection of genomes and to add a set of new genomes to an existing pangenome without rebuilding the accumulated pangenome from scratch. The workflow of PanTA pipeline is summarized in Fig. 1. PanTA takes as input a list of genome assemblies and their annotations. PanTA then extracts the protein coding regions as specified by the



Fig. 1 The schematic depict of PanTA workflow. The flowchart of PanTA pipeline in both single and progressive modes. In single model, the gene clustering process involves the reduction of protein sequences to representative gene sequences using CD-HIT, the all-against-all alignment of the representative sequences by DIAMOND, and the MCL clustering. In progressive mode, new protein sequences are first matched with the existing representative sequences and only unmatched sequences are reduced to form new groups. Pairwise alignments are performed only between new representative sequences against existing representative sequences and among new representative sequences

annotation, and translates them to protein sequences. In the process, it verifies and filters out coding regions that are incorrectly annotated, e.g., those with ambiguous bases or with frame length not divisible by 3, that can potentially introduce noise into the clustering step and the downstream analyses.

The core of the pipeline is the clustering of all genes in the collection into gene clusters, which represent the gene families in the collection. PanTA first runs CD-HIT [26] to group similar protein sequences together, and essentially reduces the set of all protein sequences to a smaller set of representative sequences from the groups. The default thresholds for sequence identity and gene length difference are set to 98% and 70% respectively, and the values can be adjusted by users. The representative sequence of each group is the longest sequence of the group. PanTA then performs an all-against-all alignment of the representative sequences with DIAMOND [28] or optionally BLASTP [27]. The resulting pairwise alignments are filtered to retain those that pass certain thresholds of sequence identity (default at 70%), alignment length ratios, and length difference ratios. These alignments are inputted into Markov clustering (MCL) [29] that clusters the representative sequences into homologous groups of genes. Each protein sequence is then assigned to the gene cluster its representative sequence belongs to.

While the clustering strategy employed by PanTA is similar to that of recent pangenome tools such as Roary [18], PIRATE [22], and Panaroo [25], we optimize the pipeline configurations that speed up the process without compromising the clustering accuracy. Notably, during the sequence grouping stage, PanTA runs CD-HIT only once at sequence identity 98% which is similar to Panaroo, instead of conducting multiple rounds of grouping at differing sequence identity levels as Roary and PIRATE. It also uses the word size of 5, which is suitable for such a high level of sequence identity. This word size, also used by PIRATE, enables CD-HIT to operate much faster than Panaroo's use of word size of 2 and, at the same time, produces similar sequence grouping. We also found DIAMOND significantly faster than BLASTP for the all-against-all alignment at the same level of sensitivity, confirming the previous report [28].

PanTA can run in progressive mode where it adds new genomes into an existing pangenome without rebuilding the pangenome from scratch. In this mode, PanTA uses CD-HIT-2D, a tool in the CD-HIT suite [26] to match new protein sequences extracted from the new samples to the representative sequences from the existing groups. The protein sequences that are matched to an existing group are assigned to the groups and by proxy, to the existing gene cluster. Only unmatched sequences are subject to CD-HIT to create new groups (Fig. 1). By running CD-HIT clustering solely on the new genes in the added batch, PanTA significantly reduces the running time and memory usage over grouping all genes in the accumulated collection. Similarly, during the all-againstall alignment step, PanTA first performs alignment of the representative sequences of the new groups against the representative sequences of the existing groups. It then runs the all-against-all alignment of only the new representative sequences, that is, those not aligned to the existing groups to the defined sequence identity threshold. The two sets of alignments after filtering are combined and then subject to MCL clustering. With this strategy, PanTA reduces the number of sequences in the grouping and alignment steps which are the most resource-intensive steps of the whole pipeline. As a result, the process is significantly accelerated.

Finally, PanTA provides options to perform post-processing steps, including splitting paralogous clusters and multiple alignment of genes in each cluster. For split paralogs, PanTA employs the conserved gene neighborhood (CGN) approach as described in [18]. Sequences of each gene cluster are aligned using MAFFT [30] at both DNA and protein levels. PanTA then generates output reports according to the standards set out by Roary, which include a spreadsheet detailing the presence and absence of each gene in each isolate as well as a summary of pangenome statistics.

PanTA is significantly more efficient than existing pangenome inference tools

We evaluated the performance of PanTA and compared it with that of existing pangenome construction methods on collections of bacterial genomes. We sourced the genomes of isolates from three bacterial species *Streptococcus pneumoniae*, *Pseudomonas aeruginosa*, and *Klebsiella pneumoniae* that are known for carrying resistance to multiple antibiotics. These three species were chosen to cover a range of genome sizes and CG content as well as both gram-positive and gram-negative. We selected 600 *S. pneumoniae*, 800 *P. aeruginosa*, and 1500 *K. pneumoniae* isolates to create three datasets, named Sp600, Pa800, and Kp1500, respectively (Table 1). We downloaded their

Dataset	Species	Number of genomes	Genome size	Ave. gene number	CG content	Gram
 Sp600	S. pneumoniae	600	2.0 Mb	2.0k	40%	Positive
Pa800	P. aeruginosa	800	6.1 Mb	6.0k	67%	Negative
Kp1500	K. pneumoniae	1500	5.4 Mb	5.1k	57%	Negative

 Table 1
 Characteristics of the three datasets to evaluate pangenome construction tools

genome assemblies from the RefSeq database [31] and ran Prokka [32] to generate the gene annotations of these genomes in gff3 format. The gffs files were then used as input for the pangenome construction process.

We compared PanTA to the pangenome inference methods that are currently considered state-of-the-art in terms of scalability. Specifically, we included in the comparison Roary [18], PIRATE [22], PPanGGOLiN [23], and Panaroo [25]. Other pangenome construction methods such as panX [20], COGSoft [33], and PEPPAN [24] were reported to be prohibitively expensive for application to thousands of genomes [24, 25] and hence were excluded from the comparison. We ran all the competing tools using their default and recommended parameters. To evaluate the performance of the tools with varying input sizes, we ran them on subsets of these collections, gradually increasing in size. All computational experiments were conducted on a laptop computer with a 20 hyper-thread CPU (Intel Core i7-1280P) and 32 Gb of memory, running Ubuntu Linux 22.0. All methods are parallelized with multi-threading, and we ran them on 20 threads, the number of CPU cores of the computer. We recorded the wall time and peak memory usage of all the runs for comparison.

Most pangenome inference methods have an option to split paralogs where clusters containing paralogous genes are identified and subsequently split into true ortholog clusters. However, they have different definitions of paralogous clusters, and employ different paralog splitting strategies, leading to varying levels of splitting rigorousness. The most rigorous strategy is employed by Roary which considers a cluster paralogous if it contains more than one gene from the same genome. It then uses conserved gene neighborhood information to split homologous groups. This split paralog strategy is also performed by PanTA. PIRATE and Panaroo consider a pair of genes paralogs if they exhibit over 98% sequence identity from the CD-HIT pre-clustering step, resulting in significantly fewer paralogs compared to Roary and PanTA. PPanGGOLiN does not provide the option to split paralogs. Because of the differences in the rigorousness of the tools, we ran the competing tools with the same base configuration, that is without split paralog option. We also excluded the post-processing step that performs multiple alignment of gene clusters, as these tools eventually call a multiple alignment method such as MAFFT [30] for this task.

Figure 2a shows the computational resources in wall time and peak memory against the size of the genome collection for the competing pangenome inference methods on the three datasets. Additional file 1: Fig. S1 shows the differences in the number of folds in resources required by each tool against PanTA. We noted that PIRATE crashed when inferring the pangenomes for the sets of 1200 and 1500 *K. pneumoniae* genomes and PPanGGOLiN ran out of memory (32 Gb) in constructing the pangenomes for 800 *P.*



(c) The concordance in ARI between the gene clustering and the grouping by gene annotation

Fig. 2 The performance of PanTA and existing tools on the three data collections. **a** Wall time and memory usage among the competing tools at various dataset sizes. Note that PPanGGOLiN and PIRATE were unable to complete the pangenome construction for some large datasets. **b** The number of core genes and accessory genes of the pangenomes constructed by all the tools. **c** The concordance in Adjusted Rand Index between the gene clustering and the grouping by gene annotation

aeruginosa genomes and for 900 or more *K. pneumoniae* genomes. Hence, the results for these runs are not included in the comparison. We observed that all methods exhibited an approximately linear increase in time and memory usage against the input size. Strikingly, we found PanTA was significantly faster than the competing methods across three datasets by a large margin (Fig. 2a, top panel). Specifically, it took under 2 min to build the pangenome for 600 *S. pneumoniae* genomes and 0.168 h and 0.207 h to build the pangenomes for 800 *P. aeruginosa* and 1500 *K. pneumoniae* genomes, respectively. The next fastest method is PPanGGOLiN, which took between 1.8 and 2.2 times longer than PanTA on the small dataset Sp600, and the fold difference increased to 2.3–2.7 times in the Pa800 dataset and 3.0–4.5 times in the Kp1500 dataset (Additional file 1: Fig. S1). Panaroo took much longer, over 10 times longer than PanTA for the larger datasets Pa800 and Kp1500. Roary was the slowest, about 15 times slower than PanTA in most cases.

In terms of memory usage, PanTA was also the most memory-efficient, requiring only 5.1 Gb of memory for all 1500 *K. pneumoniae* genomes. Panaroo used more than twice as much memory (11.8 Gb) for the same dataset, and generally the fold difference tended to increase with larger datasets. PIRATE exhibited similar memory usage profiles, but it was unable to construct the pangenomes for 1200 and 1500 *K. pneumoniae* genomes. Roary consumed 22.4 Gb of memory for the Kp1500 dataset, which is 4.4 times more than PanTA. While PPanGGOLiN was the second fastest, about twice as slow as PanTA, it required the most memory, about 7 times more than PanTA for the large datasets. Specifically, it required 26.3 Gb and 23.1 Gb of memory for analyzing 600 genomes of *P. aeruginosa* and *K. pneumoniae* respectively; it also encountered memory issues when analyzing configurations with more than 600 genomes of these species.

Figure 2b compares the numbers of core genes and of accessory genes in the pangenomes inferred by the completing methods. The numbers of genes are also shown in Additional file 1: Table S1. Note that for the Pa800 and Kp1500 datasets, PPanGGOLiN did not complete constructing the pangenomes beyond 600 genomes.

We show the statistics from the pangenome constructed from the 600 genomes of each species for a fair comparison of all five methods. We observed that the pangenomes produced by Panaroo and PanTA contained a consistent number of gene families as the result of the same sequence identity threshold (70%). Roary, which used a higher threshold (95%), resulted in many more gene clusters in its inferred pangenomes. On the other hand, PIRATE applied a series of thresholds ranging from 50 to 95% giving rise to the smallest number of gene clusters. All the methods however inferred similar numbers of core genes, in that PanTA pangenomes reported within 5% of core genes with the corresponding pangenomes produced by the other methods (Fig. 2b).

We further assessed the accuracy of the pangenomes constructed by competing methods. While there is no established benchmark to assess the accuracy of pangenome inference methods, we used the degree of concordance of the gene family clustering and the gene annotations. We collected all genes in the collection of genomes annotated by Prokka to a known gene family, that is, excluding genes that are marked as hypothetical protein. In other words, the benchmark considered genes that were aligned by Prokka to the same gene family in the common protein databases (such as UniProt, HMM, and species-specific databases) to be in the same cluster. We note that the average proportions of hypothetical genes in a sample were 0.391 (std = 0.014), 0.409 (std = 0.018), and 0.262 (std = 0.020) for S. pneumoniae, P. aeruginosa, and K. pneumoniae respectively. Concordance was assessed by calculating the Adjusted Rand Index (ARI) [34], which is a measure of similarity between clustering results. An ARI value of 1.0 indicates a perfect match between two clusterings, while a value of 0.0 indicates random grouping. The ARI of the competing methods on the three datasets is presented in Fig. 2c. We found that the gene clustering by PIRATE was markedly different from that inferred from gene annotation, indicated by the low ARI values (Fig. 2c). We attribute this discordance to the use of a series of sequence identity thresholds ranging from 50 to 95% by PIRATE during gene clustering, as opposed to a fixed threshold for gene annotation by Prokka. Among all the methods that use a fixed sequence identity threshold, PanTA had comparable ARI scores with others while being significantly faster, which urderscores the efficiency of the approach.

PanTA progressively builds pangenome

We next evaluated the performance of PanTA in progressive mode where it updates an existing pangenome when new samples are added without the need of rebuilding the pangenome from scratch. For each of the aforementioned datasets, we ran PanTA to construct the pangenome of the smallest partition and progressively added the genomes of the subsequent partitions into the pangenome. We noticed Panaroo also offered a similar functionality, namely Panaroo-merge, that merges the pangenomes of multiple collections together. For comparison, we ran Panaroo on each partition of the dataset, and then applied Panaroo-merge to merge the partition collections together. In these experiments, we collected the wall times for each pangenome as the sum of the wall time of each step and the peak memory usage as the maximum amount of memory at each step. Figure 3 presents the computational resources consumed by both methods on the three datasets. We also included the resources needed by both methods when rom-puting the pangenomes from scratch as part of the comparison. As presented in Fig. 3,



(a) Computational resources of PanTA and Panaroo in single and progressive modes

(b) Concordance of pangenomes constructed in single and progressive modes by PanTA and Panaroo

Fig. 3 Performance of PanTA in progressive mode. **a** Comparison of computation times and memory usage of PanTA and Panaroo in single and progressive/merge modes. **b** The number of gene families and core genes inferred by PanTA and Panaroo in single and progressive modes. **c** Concordance in Adjusted Rand Index of pangenomes constructed in single and progressive modes by PanTA and by Panaroo. PanTA in the two modes produced near identical pangenomes while the pangenomes inferred by Panaroo and Panaroo-merge had lower concordance

Panaroo-merge improved memory usage by 20% over Panaroo at the cost of 70% longer running time. On the other hand, PanTA in progressive mode saved memory usage by half while maintaining a similar running time over the single mode. All in all, PanTA in progressive mode consumed only 25% and 15% of the amount of memory required by Panaroo and Panaroo-merge, respectively, while was 10 and 17 times faster.

We analyzed the concordance of the pangenomes constructed by the two modes by calculating the ARIs between the two clusterings. In this calculation, we used all the genes present in the collection instead of only the annotatable genes. As shown in Fig. 3b, PanTA in progressive mode produced almost identical clusterings to that in single mode (ARI > 0.99 for Sp600 and Pa800 and > 0.975 for the Kp1500). The pangenomes produced by the two versions of Panaroo are much less concordant, with ARI values of 0.93, 0.92, and 0.89 for the three datasets, respectively.

We posed a hypothetical scenario that the datasets were generated in specified batches, each at a different time point. This reflects the nature of collecting and sequencing bacterial isolates in most research laboratories, infectious disease surveillance centers, and healthcare facilities. We further posited that the computational costs were measured by the time required to run on a computer with specific CPU and memory configurations, similar to those offered by a cloud computing service. We then measured the cumulative computation resources in CPU hours required to compute the pangenomes each time a batch became available. For all methods, including Panaroo and PanTA in single mode, the computation resources would include that for recomputing the pangenomes from scratch. Panaroo-merge would only need to compute the pangenome of the new batch and then merge the pangenome of the batch to the existing pangenome. PanTA in progressive mode would add the new batch of genomes to the pangenome. Figure 4 shows the fold differences of all the methods against PanTA-progressive. As expected, PanTAprogressive required only a small fraction of computing resources compared to all other methods after a few batches. The two methods that could complete the construction of the pangenome in the Kp1500 dataset, Roary and Panaroo, respectively consumed 45.2



Fig. 4 The fold difference in computational resources for both CPU time (top panel) and memory (bottom panel) between existing methods and PanTA-progressive

and 30.4 times more CPU-hours than PanTA-progressive, in addition to 11.7 and 6.1 times more memory. Although PPanGGOLiN was only 4.5 times slower than PanTA in constructing the pangenome of 600 *K. pneumoniae* genomes, the total time to compute the initial pangenome and recompute the updated pangenome was 6.9 folds higher than that of PanTA-progressive after two batches. The actual computational cost was even much higher considering that PPanGGOLiN required > 14.5 times as much memory. PIRATE required approximately 25–30 times more CPU hours and 3–5 times more memory compared to PanTA-progressive after processing 3–4 batches across the three datasets. PanTA-progressive also saved 60–70% of both CPU hours and memory usage compared to PanTA single mode.

It is expected that the pangenome inference methods in single mode required higher and higher accumulated computational resources than PanTA-progressive did as more batches of data became available. We examined the resources consumed by Panaroomerge which employs a similar approach to PanTA-progressive. Indeed, the increase of fold difference between Panaroo-merge and PanTA-progressive was much slower than other methods in single mode. However, it exhibited a large factor of fold difference, and the factor tended to increase with the genome size: 6–7X for *S. pneumoniae* (genome size 2.1 Mb), 15X for *K. pneumoniae* (5.6 Mb), and > 20X for *P. aeruginosa* (6.1 Mb).

Building the pangenome of a growing genome collection

The primary goal of PanTA is to analyze and manage the extensive and fast-growing collections of microbial genomes. We demonstrate this utility by applying PanTA to a realistic and expanding collection of bacterial genomes. To this end, we collected all *Escherichia coli* genomes that were deposited into the RefSeq database [31] during the three years 2020, 2021, and 2022. *E. coli* is one of the most well-studied model prokaryotic organisms and is known for its genotypic diversity and pathogenic for both humans and animals [35]. After removing outliers, we obtained a dataset of 12,560 genomes (the "Methods" section). To demonstrate the growing nature of the dataset, we grouped the samples based on the quarters in which they were released. Table 2 shows the breakdown of the samples.

Quarter	#isolates	#isolates accum.
Q1-2020	534	534
Q2-2020	713	1247
Q3-2020	830	2077
Q4-2020	1109	3186
Q1-2021	1166	4352
Q2-2021	694	5046
Q3-2021	1411	6457
Q4-2021	866	7,323
Q1-2022	1645	8,968
Q2-2022	1233	10,201
Q3-2022	1214	11,415
Q4-2022	1145	12,560

Table 2 Number of *E. coli* samples deposited into RefSeq database between 2020 and 2022 by quarter



Fig. 5 Computational resources for constructing the pangenome for E. coli

We ran PanTA to build the initial pangenome of the genomes collected in the first quarter. We then progressively add genomes from subsequent quarters into the collection. For comparison with PanTA in single mode, we also ran PanTA on the accumulated data at each quarter. As shown in Fig. 5, PanTA in progressive mode needed only 16.6 Gb of memory to construct the pangenome for over 12000 *E. coli* genomes while the single mode consumed 30.1 Gb of memory. Both modes exhibited similar running times, around 6.5 h.

Encouraged by the scalability of PanTA, we proceeded to build the pangenome for the entire set of *E. coli* genomes from the RefSeq database. We downloaded all *E. coli* genomes that were released prior to 2020, and after filtering outliers, we obtained 15,625 genomes in addition to the previously collected set (the "Methods" section). We divided these genomes into batches of maximum 1000 genomes each and iteratively added these batches into the *E. coli* pangenome with PanTA-progressive. In effect, we constructed the pangenome of all 28,275 high-quality *E. coli* genomes from the RefSeq database. For this experiment, we used another laptop computer equipped with a 32-core CPU and 64 Gb of memory. Strikingly, the pangenome of *E. coli* species was inferred on a laptop computer with a total time of 32 h, including the time to build the pangenome from the past 3 years. The peak memory recorded during the pangenome construction was 39.9 Gb.

Discussion

Bacteria are among the most diverse life forms on earth, evidenced by the high level of variability of gene content across strains in a species. It is therefore possible but undesirable to use the genome of a single isolate as the reference to represent a clade. Pangenome analysis offers an alternative approach where all gene families of the clade constitute the pangenome that represents the total diversity of the clade.

Most computational methods for pangenome construction usually apply clustering of gene sequences. These methods in most cases run multiple times of CD-HIT clustering on different levels of sequence similarity in order to achieve stability of clustering. In developing PanTA, we use only one round of CD-HIT clustering and yet we obtain the near identical pangenomes with existing tools on the same sequence identity threshold. PanTA is shown to be multiple times faster than and requires less than half of the memory consumed by the current state of the arts.

The bacterial genome collections are growing by nature as more and more genomes are routinely sequenced in laboratories as well as in research and medical settings around the world. PanTA addresses the complexity of rebuilding pangenomes by providing the progressive mode where new genomes are added to an existing pangenome. By utilizing the group membership information of the existing clustering, PanTA needs to compare the genes in the new genomes with only existing groups and thereby are significantly faster than rebuilding the pangenomes from scratch. Interestingly, we found that building the pangenome progressively from batches of genomes takes a similar amount of time to build from the whole collection, while reducing the memory requirements by half. The scalability of PanTA is demonstrated by the ability to construct the pangenome for the E. coli species from the entire set of 28000 genomes from RefSeq database on a laptop computer in an unprecedented 32-h timeframe. More strikingly, PanTA can construct the pangenome progressively when new samples are added into the collection, without recomputing the accumulated collection from scratch. PanTA in progressive mode is shown to consume orders of magnitude less computational resources than existing solutions in analyzing growing data collections. This makes PanTA practically suitable for analysis of the large collections of bacteria in the sequencing ages.

PanTA is designed to be compatible with existing pangenome tools and as such it can be seamlessly integrated into existing bacteria genomics analysis pipelines [36, 37]. Specifically, it takes as input the annotations in GFF3 format, a standard set by Prokka [32], and followed by the modern annotation tools including DFAST [38], PGAP [31], and Bakta [39]. The outputs of PanTA including gene family memberships and gene alignment are also formatted according to the standards in the field. The gene clustering produced by PanTA can also be the foundation for constructing the population graph-based pangenome representation. The graph-based pangenome presentation has been employed for error-correction by Panaroo and for depiction of overall genomic diversity by PPanGGOLiN. This features will be considered to be implemented in the future releases of PanTA.

Methods

Pangenome pipeline

PanTA accepts input genomes in GFF3 files which store gene annotations in gff format followed by the genome assembly in fasta format. This format is the output from Prokka [32] and has been popularized for the pangenome analysis started by Roary [18]. Each genome is associated with a unique ID which can be input by the user or generated by PanTA. The ID of each contig in the genome, as well as each annotated coding sequence, must be unique. Coding sequences are extracted and translated into protein sequences. Coding sequences that are less than 120 nucleotides in length or lack both a start and stop codon are excluded. Protein sequences containing more than 5% of unknown amino acids are also removed. Next, a fast sequence grouping is performed using CD-HIT [26] with an identity threshold of 98%. The representative sequences from CD-HIT are compared all-against-all by DIAMOND [28] or BLASTP [27]. The e-value threshold is set to 10e-6 by default. To reduce the time required for all-againstall alignment, the list of representative sequences is divided into smaller chunks of up to 20,000 sequences to enable parallel matching. The identified matches are filtered to retain those with sequence identity above a threshold (default at 70%). The DIAMOND result is then input into MCL [29], which uses a normalized bit score for clustering with an inflation value of 1.5. Finally, the removed sequences in the CD-HIT step are merged back into the MCL clusters. The detailed parameters of the tools are listed as follows.

- CD-HIT: cd-hit -s 0.98 -c 0.98 -T <number_thread> -M 0 -g 1 -d 256
- DIAMOND: diamond blastp -p <number_thread> -evalue 1e-06 -outfmt
 6 qseqid sseqid pident length mismatch gapopen qstart qend sstart send evalue bitscore qlen slen -max-target-seqs 2000
- BLASP: blastp -query <chunked_file> -db <blast_db> -evalue 10-e6 -num_threads 1 -outfmt "6 qseqid sseqid pident length mismatch gapopen qstart qend sstart send evalue bitscore qlen slen" -max_target_seqs 2000
- MCL: mcxdeblast -m9 -score r -line-mode=abc <input> | mcl - abc -I 1.5 -te <num_thread>

Add samples pipeline

First, the protein sequences of the new samples are compared and matched with CD-HIT's representative sequences from the previous collection. This is performed by CD-HIT-2D with the identity threshold of 98%. The protein sequences that are matched to a representative sequence are assigned to the represented group. The unmatched sequences are clustered by CD-HIT to create new groups with new representative sequences. The new representative sequences are then subject to all-against-all alignment by the alignment method of choice, i.e., DIAMOND or BLASTP. The new representative sequences are also aligned against the existing representative sequences. The two sets of alignments are then filtered according to the criteria and then combined with the existing set of alignments in the pangenome. Finally, MCL is applied to the combined set of alignments as described above.

CD-HIT-2D cd-hit-2d -i <existing_group> -i2 <new_sequences> -s 0.98 -c 0.98
 -T <num_threads> -M 0 -g 1 -d 256

Annotating clusters

For each cluster, PanTA maintains a list of all the gene names and gene products of all genes in the cluster. It also keeps gene length statistics such as the number of genes, minimum, maximum, and average gene length in the cluster. The cluster is assigned a name taken from one of the annotated genes. The gene product for the cluster is the concatenation of all the gene products of the gene members. PanTA also picks the longest gene sequence to be the representative sequence for the cluster.

Post-processing and output

PanTA presents the pangenome following the standard set out by Roary. Specifically, the presence and absence of genes in each sample are presented in CSV and Rtab formats. Upon users' request, PanTA performs multiple sequence alignment of all gene clusters. Either or both genomic and protein sequences can be aligned. In addition, PanTA stores the existing all-against-all alignments and the existing CD-HIT groupings for subsequent analyses.

Performance comparisons of pangenome inference methods

The lists of isolates in the three datasets Sp600, Pa800, and Kp1500, together with their accession IDs and the URLs of their genome sequences, are provided in the supporting data (see Availability of data and materials). Their genome sequences were downloaded and were subject to annotation by Prokka in its recommended parameters (prokka – force – -cpus 8 – -addgenes – -mincontiglen 200 – -prefix <accession_id> – -locus <accession_id> – -genus <genus> – -species <species>). The resulting annotations in GFF3 format are also provided in the supporting data. The three datasets were split into batches of 150, 200, and 300 samples respectively based on the order specified in the lists.

The competing methods Roary, PIRATE, Panaroo, and PPanGGOLiN were installed with their stable releases via conda. They were run with their parameters as follows:

- Roary: roary -p 20 -s -f <output_folder> <list_of_samples>
- PIRATE: PIRATE –para–off –t 20 –z 0 –o <output_folder> –i <folder_containing_ samples>
- Panaroo: panaroo -merge_paralogs –t 20 -clean-mode strict –o <output_ folder> –i <samples>
- Panaroo in merge mode: the pangenome for a new batch of genomes was generated with the parameters as above, and was merged into the existing pangenome with the parameters panaroo-merge – -merge_paralogs -t 20 -o <output_folder> -d <existing_pangenome> <new_batch_pangenome>
- PPanGGOLiN: ppanggolin workflow –anno <sample_list> –verbose 2 –c 20 –o
 <output_folder> –identity 0.7
- PanTA: panta main –dont–split –o <output_folder> –g <samples>

PanTA in progressive mode: panta add – –dont–split –c <existing_pangenome> –g <samples>

Running times and memory usage of the computational experiments were collected with the time utility, i.e., having /usr/bin/time –v preceding the command line. The wall time was determined from the "Elapsed" field, whereas memory usage was from the "Maximum resident set size." For Panaroo in merge mode and PanTA in progressive mode, the total time of constructing the pangenome was the sum of the wall times from all preceding steps, while the memory usage was the maximum.

Data collection for the E. coli dataset

The set of genomes available on RefSeq database was downloaded from https://ftp.ncbi. nlm.nih.gov/genomes/refseq/assembly_summary_refseq.txt (accessed February 22, 2023). We selected only genomes of samples belonging to *E. coli* species. The genome sequence (fna file) and genome annotation (GFF file) for each sample were downloaded and combined to generate a GFF3 format file. Coding sequences that were shorter than 120 bp or contained non-canonical nucleotides were removed. To remove outliers, we inspected the histograms of genome sizes, number of genes, and N50 statistics (Additional file 1: Fig. S2) and selected genomes that were between 4.2 Mb and 5.9 Mb long, contained between 4200 and 5500 genes, and having N50 statistics of 50 kb or higher. These genomes were grouped into quarters based on their release dates. The Jupyter notebook and the script that was used to download and process the dataset and to run pangenome construction were provided in the supporting data.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13059-024-03362-z.

Additional file 1: Table S1. The numbers of core-, accessory- and total genes in the pangenomes constructed by the completing methods. The gene counts were from the pangenomes of 600 genomes for all the datasets so that PIRATE and PPanGGOLIN could complete the pangenome construction. Fig. S1. Fold difference between computational timeand memoryrequired by existing methods against PanTA. Fig. S2. Distributions of the gene number, genome size and N50 of the *Escherichia coli* genomes collected from RefSeq database.

Additional file 2: Review history. The peer review history.

Review history

The review history is available as Additional file 2.

Peer review information

Andrew Cosgrove was the primary editor of this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.

Authors' contributions

MDC, SHN, TN, and HAN conceptualized and designed the project. DQL, TAN, SHN, and MDC implemented the software. DQL, TAN, SHN, TTN, TN, and MDC performed the experiments and data analysis. CHN, HTP, THH, and NSV managed the project. DQL, TAN, and MDC drafted the first version of the manuscript. All authors wrote, revised, and approved the final manuscript.

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Availability of data and materials

The PanTA software is open source and is publicly available on GitHub at https://github.com/amromics/panta [40] or on Figshare at DOI 10.6084/m9.figshare.23724705 [41] under an MIT license. The software is also distributed via pip. An official docker container (amromics/panta) is available on Docker Hub at https://hub.docker.com/r/amromics/panta [42]. The genome assemblies for datasets were downloaded from RefSeq database [31]. Supporting data and the source code version used in the manuscript are available on Figshare at DOI 10.6084/m9.figshare.23724705 [41].

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

TN, HAN, and MDC are founders of AMROMICS JSC. DQL, TAN, and SHN are consultants to AMROMICS JSC. All other authors declare no competing interests.

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