G-SAIP: Graphical Sequence Alignment Through Parallel Programming in the Post-Genomic Era

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ABSTRACT: A common task in bioinformatics is to compare DNA sequences to identify similarities between organisms at the sequence level. An approach to such comparison is the dot-plots, a 2-dimensional graphical representation to analyze DNA or protein alignments. Dot-plots alignment software existed before the sequencing revolution, and now there is an ongoing limitation when dealing with large-size sequences, resulting in very long execution times. High-Performance Computing (HPC) techniques have been successfully used in many applications to reduce computing times, but so far, very few applications for graphical sequence alignment using HPC have been reported. Here, we present G-SAIP (Graphical Sequence Alignment in Parallel), a software capable of spawning multiple distributed processes on CPUs, over a supercomputing infrastructure to speed up the execution time for dot-plot generation up to 1.68× compared with other current fastest tools, improve the efficiency for comparative structural genomic analysis, phylogenetics because the benefits of pairwise alignments for comparison between genomes, repetitive structure identification, and assembly quality checking.

KEYWORDS: G-SAIP, HPC, bioinformatics, dot-plots, graphical alignments, post-genomic era

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Introduction

Bioinformatics is a multi-disciplinary area supporting the discovery of biological information utilizing computational approaches.¹ This research field is at the intersection of several sciences like biology, computer sciences, and mathematics, intending to analyze and classify biological data.² One of the most common bioinformatics tasks, relevant in metagenomics and phylogenetic analysis,³ is the sequence alignment,^{4,5} which consists of comparing 2 (pairwise)⁶ or more (multiple)⁷ nucleotides or proteins sequences against a reference.

One of the most utilized alignment types is the graphical sequence alignment,^{8,9} which provides visualization of rearrangements, insertions, deletions, and other structures found in DNA or protein sequences. Graphical alignments are commonly represented in a dot matrix called dot-plots.¹⁰ Dot-plots are rectangular matrixes where columns and rows represent the residues to be aligned; at each cell, a dot is painted with a grayscale intensity proportional to the degree of similarity of the sequences at that point.¹¹

There are several published software packages able to generate dot-plots. Dotter9 is one of the most popular graphical DECLARATION OF CONFLICTING INTERESTS: The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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pairwise sequence aligners, using dynamic programming¹² and well suited for small DNA or protein sequences (few 1000 of nucleotides).¹³ Other available software programs are Dotlet¹⁴, which runs on a web server; JDotter¹³ is a version of Dotter running on a remote java platform, Tuple_plot¹⁵ which proposed a different way to calculate the dot-plot and can reduce the noise for large sequences (>10kB) and Gepard¹⁶ using a heuristic suffix array method¹⁷ to generate dot-plots of small and large sequences but with a high noise level. The SFILE¹⁸ library was designed to deal with big-data sequences, enhancing out-of-core management and using a k-mer value identification to reduce computational space. More recently, novel software has been published as Flexidot,¹⁹ which generates high-quality dot-plots for small repetitive sequences, and finally D-genies,²⁰ a standalone and web application that uses the minimap²¹ output to calculate the alignment for chromosomes and genomes, D-genies also use MashMap,²² an approximate algorithm for computing local alignment boundaries between long DNA sequences using k-mers and taking advantage of HPC strategies and mapping genome assembly or long reads to other reference sequences. However, despite significant progress in using large sequences datasets, the quality and the execution times of dot-plots still represent a challenge.



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Advances in next-generation sequencing technologies and associated low costs have allowed an exponential increase in available genetic information,²³ known as the sequencing revolution.²⁴⁻²⁷ As a result, the challenges have shifted from sequencing organisms to analyzing their genomes in a post-genomic era.^{28,29} This shift required a new generation of algorithms considering the use of parallel, distributed, and other high-performance computing (HPC) techniques to accelerate the genome data analyses.³⁰

Developments in HPC, supercomputing and parallel programming have improved the execution time in several areas,³¹⁻³⁴ due to parallel programming can launch processes over heterogeneous architectures such as CPU, GPU, or CPU + GPU using libraries for programming in a fast and flexible way, even with shared or unshared memory.³⁵ In bioinformatics, these techniques allow accelerating the analysis of genetic information.³⁶ Different bioinformatics applications use parallel programming approaches.³⁷⁻⁴⁰ These applications focus on multiple sequences alignments⁴¹⁻⁴⁵ and non-graphic paired alignment of protein nucleotides,⁴⁶⁻⁵⁰ including repetitive structures.⁵¹ However, graphic aligners that use parallel strategies are not available.

In this work, we reported G-SAIP (Graphical Sequence Alignment in Parallel), a tool that can be easily integrated into a pipeline and HPC-based strategy that follows the Flynn⁵² taxonomy SIMD (simple instruction multiple data). G-SAIP, taking advantage of MashMap, performs graphical pairwise sequence alignment (1 channel and 8 bits image) at the genomic level on CPU architectures over multiple nodes speeding up execution times through parallel programming in order to provide tools for analyzing a massive amount of data produced by large scales genomic projects such as the 10K plant genomes⁵³ and the Earth BioGenome.⁵⁴ In contrast with other programs, G-SAIP prioritizes process parallelization over programs like gepard to generate dot-plots, as well as uses the principles of sequence mapping to score similarities with programs like minimap2 and MashMap which perform file alignments as results and use multi-threaded processes in their execution. This tool can also be used for quality verification of genome reference-based assemblies.

Materials and Methods

G-SAIP implementation

G-SAIP was developed using Python 3.8⁵⁵ and with parallel computing support using mpi4py.⁵⁶ It is a DNA graphical aligner that takes advantage of MashMap⁵⁷ for sequence alignment due to its speed, and requires input parameters as the reference and query sequences in FASTA format that must be declared in the command line execution to calculate the dot-plot. Also, users can define optional parameters like MashMap Segment similarity, identity percentage for filtering, *k*-mer size, and G-SAIP output image attributes like image width, height, and word size to enhance the resulting quality.

$$Window_{size} = \frac{Total_sequence_length}{1024}$$
(1)

Equation 1. Calculated window size by G-SAIP, where Total_ sequence_length is the complete size of residues from larger sequence and 1024 is the maximum size of the result image.

To generate the dot-plot, G-SAIP receives the nucleotide FASTA files for reference and query sequences; these files can contain 1 or more sequences. Hence the algorithm joins each file into a unique sequence to facilitate the execution. Thus, the largest sequence file is split into subsequences of length calculated by equation (1), ensuring that each pixel of the default output image size represents the minimum number of nucleotides and keeps the significant information. Then, G-SAIP uses MashMap to calculate the score of each region; the score value is extracted from the result file and mapped from 0% to 100% (identity) to pixel intensity 0 to 255. Next, G-SAIP generates the dot-plot matrix, which is scaled to height and width defined by the user (by default is 1024 pixels and 1024 pixels). The software preserves the intensities extracted from MashMap but, if the user wants to reduce the image noise, G-SAIP has a filter tool that reduces to zero the intensities under a given threshold and assigns the maximum intensity to pixels above the threshold. Another option available is to generate a dotplot with 3 colors. The user is asked for 2 values between 0 and 100; the dots are red-colored for scores under the minimum value. For scores above the maximum number, dots are colored green, and scores between those ranges are orange painted. Finally, the image is saved in SVG, PNG, or PDF format as a specified format by the user (by default, PNG is used), and the algorithm removes temporary files used during the execution.

G-SAIP parallel strategy

G-SAIP uses mpi4py, a Python implementation of Message Passing Interface (MPI),⁵⁸ specifically for Open MPI.⁵⁹ This library allows graphical alignment to be performed parallel over multiple CPU cores belonging to a single node or distributed nodes in an HPC cluster. The software takes all subsequences to calculate the number of sequences that each processor will process. Then, the master process creates 1 file per worker node (with the subsequences that it will use) and sends this file as the same as the shorter-joined sequence to worker processes. Finally, each process runs MashMap with its individual sequences file, shorter-joined sequence, and user parameters. By default, G-SAIP defines MashMap parameters like segment length, *k*mer size, identity percentage, and filter mode as 5000, 16, 95, and None. In this way, the algorithm takes advantage of this HPC strategy to do several alignments simultaneously.

The output file generated by MashMap is processed to extract alignment scores, which are used to generate and fill the $N \times M$ matrix, where N is the subsequence length, and M is the shortest sequence length. Finally, when each process is done,



Figure 1. G-SAIP parallel strategy diagram. Sub-sequence file corresponds to a file containing sub-sequences for the largest FASTA file (can be the reference or the query file). One-sequence file corresponds to the other file (the shorter).

the master process takes all matrixes generated and creates the final dot-plot matrix. Figure 1 resumes the parallel strategy applied by G-SAIP for dot-plot calculation.

In addition, G-SAIP has a specific module to make dotplots to compare the quality of an assembly with a reference genome. Ragtag⁶⁰ is used for contigs and scaffold orders compared with a reference file. This process is executed before the execution of MashMap for alignment to determine the correct quality of an assembly file.

Availability of G-SAIP

The G-SAIP source code is open source and can be found in https://github.com/simonorozcoarias/G-SAIP. Installation instructions, how to run, sample data, and results are also available there.

Computational resources

All experiments were executed using a server with a 64-core Intel (R) Xeon(R) CPU E5-2683, with 2.1 GHz, 256 GB of RAM and the CentOS7 operating system, managed by Slurm.⁶¹

Performance tests

G-SAIP was tested for 2 perspectives to generate self-plots (a dot-plot with the same query and subject sequence) of genomes with different sizes (Table 1). First, the software was executed with different CPU numbers (2, 4, 8, 16, 32, 56, and 62) in a single node, and each execution was performed 10 times to examine the acceleration and speed provided by G-SAIP. Amdahl's law considers the elapsed execution time Table 1. Performance test sequence dataset.

SEQUENCE NAME	SIZE	GENBANK ASSEMBLY ACCESSION
Homo sapiens genome62	3.1 Gb	GCA_000001405.28
<i>Triticum turgidum</i> genome ⁶³	9.9 Gb	GCA_900231445.1
Pinus taeda genome64	22.1 Gb	GCA_000404065.3

sequentially and the execution time for the code parallel section to speed up the calculation. In addition, the execution time of the overall execution was recorded to explore the time added by sequential code.

On the other hand, *Homo sapiens* X chromosomes self-plot was generated with G-SAIP to compare the execution time differences between G-SAIP and other graphical sequence aligners: Gepard, Dotter, D-GENIES, and the MashMap Perl script to make dot-plots.

Assembly quality test

G-SAIP was used to determine the assembly quality of 2 organisms from raw sequence reads datasets. First, we used a WGS (whole genome sequencing) Illumina paired-end sequence reads of *Arabidopsis thaliana* available in NCBI SRA⁶⁵ repository under SRR10178322 accession number, with 16 Gb of size per file. The second dataset used was a WGS Illumina paired-end sequence reads of *Drosophila melanogaster* with 4Gb of size per file⁶⁵ and accessible in SRA repository with SRR10735526 accession number.

Table 2. Comparative genomic datasets test.

SEQUENCE NAME	SIZE (MB)	NCBI NUMBER
Homo sapiens X chromosome77	151	NC_000023.11
Canis lupus X chromosome78	120	NC_006621.3
Sus scrofa X chromosome79	122	NC_010461.5
Equus caballus X chromosome ⁸⁰	125	NC_009175.3
Mus musculus X chromosome ⁸¹	166	NC_000086.7
Pan paniscus X chromosome82	151	NC_027891.1

Thus, both raw sequence reads were analyzed using FASTQC⁶⁶ and Trimmomatic⁶⁷ to improve the quality of the reads. Then, each dataset was assembled with MEGAHIT,⁶⁸ Velvet,⁶⁹ ABySS,⁷⁰ and MaSuRCA⁷¹ assemblers keeping a minimum contig length of 500 bp and *k*-mer values of 31, 51, 71, and 91. So, assemblies were checked with BUSCO⁷² and QUAST⁷³ to calculate the N50 value and other significant metrics to define the best assembly. Finally, G-SAIP was executed with a quality module activated in order to execute ragtag for assembly ordering and compare each assembly with references genomes of *Arabidopsis thaliana*⁷⁴ (116 Mb) and *Drosophila melanogaster*⁷⁵ (139 Mb) to find out a relation between G-SAIP dot-plot and variables extracted with BUSCO and QUAST.

Comparative genomic test

Finally, to define the G-SAIP usefulness to perform a comparative analysis of sequences, the X chromosome of the *Homo sapiens* genome was compared against other X chromosomes of mammals listed in Table 2 because this chromosome is the most conserved during the species evolution.⁷⁶ All chromosomes were joined in a unique file for better visualization, and then a G-SAIP self-plot was generated.

Scalability test

For this test, G-SAIP was executed with 62 CPUs distributed over 1, 2, 3, 4, 5, and 6 nodes with distributed memory, using *Triticum turgidum* genome and running the algorithm 10 times for each number of nodes in order to determine if G-SAIP can run in several nodes without a significant reduction of performance compared to its execution on a single node. Finally, a strong and weak scaling test was performed to verify the software scalability.

For the strong scaling, we follow the Amdahl's law⁸³ that can be formulated as follows:

$$SpeedUp = \frac{1}{s + \frac{p}{N}}$$
(2)

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Equation 2. Amdahl's law formula for strong scaling

Where *s* is the serial time execution of G-SAIP, *p* is the proportion of execution times and N is the number of processors. In weak scaling, Gustafson's law⁸⁴ provides the formula for scaled speedup:

$$ScaledSpeedUp = s + p^*N \tag{3}$$

Equation 3. Gustafson's law formula for weak scaling

Where s, p, and N have the same meaning as in Amdahl's law.

Results

Performance test

G-SAIP was executed with a specific window size for each genome size according to equation (1), and MashMap segment length of 50000 for Homo sapiens and Triticum turgidum genomes, and 500 000 for Pinus taeda, because of its exceptionally large genome size. G-SAIP averaged execution times of genomes in Table 1 with 2, 4, 8, 16, 32, 56, and 62 cores are plotted in Figure 2a, obtaining a reduction in times from twelve (12) to seven (7) minutes for Homo sapiens, from ~1.2 hours to 23 minutes for Triticum turgidum and up to 50 minutes for Pinus taeda. Also, speed-up was calculated by taking the time with 2 cores as the nominal time due to G-SAIP using 1 working process as a master process, achieving a speed up even of $3.0 \times$ (*Triticum turgidum*) compared with nominal time. The speed-up of G-SAIP with each genome was calculated by dividing averaged times for 2 CPUs between the averaged time obtained with each other CPUs; these values are drawn in Figure 2b. We timed only the code section, which is executed in parallel (shown in Figure 1.)

For G-SAIP execution time comparison against Dotter, Gepard, D-GENIES, and MashMap dot-plot tool, a joined file with X chromosomes of Table 2 was used to generate selfplots with these graphical aligners. Table 3 demonstrates the overall execution time registered of each software and speed-up of G-SAIP against each other software; these times show performance up to 1.68× from G-SAIP concerning to current software tested. All software was executed in the same computational architecture using 62 CPUs. G-SAIP was executed with 62 CPUs, a window calculated with (1) and MashMap segment length of 50000; Dotter execution time, with default parameters, is the estimated given by the software because of the considerable time. D-Genies was installed in standalone mode, changing in the configurations files the number of CPUs to execute this software from 8 to 62, using a maximum of RAM memory up to 80 GB, changing the maximum file of the input files, and using also MashMap to calculate the alignment, other parameters were set by default for D-GENIES. Gepard dot-plot matrix was calculated with EDNA substitution



matrix, and MashMap dot-plot tool was executed with output for *Homo sapiens* with segment similarity of 50 000. All experiments can be consulted G-SAIP repo under the folder Test.

Assembly quality test

Read sequencing data of *Arabidopsis thaliana* and *Drosophila melanogaster* was analyzed with FASTQC to examine the sequences quality and adapter's presence. Then, Trimmomatic was executed to cut sequences with substandard quality. Next, FASTQC was re-executed for new sequences to visualize the new quality. Secondly, VELVET, MEGAHIT, ABySS, and MaSuRCA were executed using the trimmed data and *k*-mer values of 13, 51, 71, and 91. All completed assemblies had a minimum contig length of 800 bp. These assemblies' results were the input of QUAST to evaluate and compare the best assembly. Moreover, BUSCO was executed for *Arabidopsis* and *Drosophila* genomes to complement the QUAST results. QUAST results for each assembly with its N50 value are in Figure S1. Figure S2 shows BUSCO results for all assemblies.

Finally, each assembly was compared with its respective reference genome (*Arabidopsis thaliana or Drosophila melanogaster*) using G-SAIP with a window size of 113 000 for *Arabidopsis* and 135 000 for *Drosophila*, the rest of the parameters were assigned by default, and quality assembly parameter was set to true for ragtag execution before dot-plot calculation. Figure 3a and c showed the highest quality assembly dot-plot for Arabidopsis and Drosophila, respectively, against the reference genome, and Figure 3b and d displayed lower quality assembly dot-plot for each organism against the reference. In addition, the N50 value and the complete and single-copy BUSCO's percentage were added to each image to analyze them in the discussion section. All details about this section are in

SOFTWARE	ELAPSED TIME	G-SAIP SPEED-UP	MAXIMUM MEMORY CONSUMPTION IN GB FOR X CHROMOSOMES
G-SAIP	5.9 min	1×	1.6
D-GENIES	9.95 min	1.68×	0.95
MashMap tool	50 min*	8.4×	1.8
Gepard	19h	196×	0.98
Dotter	1699 y**	151 447 165×	0.95

This table shows how many times it is faster G-SAIP against the other tools.

*This time is from MashMap execution added the tool published in their repository to make dot-plots available in: https://github.com/marbl/MashMap/tree/master/scripts with name marbl MashMap/scripts.

**Time estimated by Dotter.

Table 3. Execution times comparison.

supplementary data, and all dot-plots are in the same document from Supplemental Figures S3 to S30.

In Table 4 are lists values of N50 and complete and single copy BUSCO's percentage for dot-plots in Figure 3.

Comparative genomic tests

In this experiment, G-SAIP generated a self-plot with X chromosomes in Table 2. Thus, all X chromosomes were pre-joined in a single FASTA file to generate a unique dot-plot with all comparisons. In this case, G-SAIP was executed with a window size of 160 000 because this is the window for the shortest chromosome to analyze. Also, MashMap segment length was set at 5000, 90% identity percentage was chosen, and k-mer size of 16 to enhance the dot-plot quality. In addition, for better visualization, the draw sequences limits were set to true to draw lines at the end of sequences. Figure 4 demonstrates the dot-plot for the X chromosomes comparison. G-SAIP generated this image in 5.9 minutes (358 seconds) for 860 Mbp compared with GEPARD which performed this alignment in 19.3 hours.

Scalability test

G-SAIP was executed with the *Triticum turgidum* genome using 62 CPUs distributed in a different number of nodes. Each run had a memory consumption of <32 Gb. We ran each test 10 times to do a box-plot graph. Figure 5 shows that G-SAIP execution times are similar between each computing node with few differences according to the number of nodes.

This experiment shows that are no significant time differences changing the number of computing nodes parameter, the performance of the time is associated with the process saturation in the cluster where the experiment was executed. In addition, weak and strong scaling experiment Formulas (2) and (3) were applied to execution times for different cores over the same node in the cluster. Results are painted in Figure 6. In GitHub repository are all scripts and result files executed for this article in "test" folder.

Discussion

Over the last years, software that performs graphical sequence aligners have proposed different approaches to generate dotplots, such as DOTTER, which has high performance for short sequences using dynamic programming, software with heuristics methods to calculate dot-plot as GEPARD or r2cat,80 and recently published as Flexidot or D-GENIES. Nevertheless, this software presents some issues with big size sequences (greater than 3Gb); some of them are no longer available, or the languages they were developed are depreciated. Software like D-GENIES report execution times shorter than G-SAIP, but we present a software that is easily integrated into pipelines, also, G-SAIP is easier configurable than D-GENIES which is a web application useful for making dot-plots within an interactive interface, and the default parameters like maximum RAM memory, maximum file size are not intuitive modifiable. In addition, the application of HPC has demonstrated high performances for several bioinformatics tasks such as multiple sequence alignment,⁴¹⁻⁴⁵ sequence mapping,^{45-48,81} analyzing transposable elements,⁸²⁻⁸⁴ and identification of transposon insertion polymorphisms,83 among others. Nevertheless, HPC software has not been deployed for graphical alignment. In this way, this software will be helpful to process big data supported by the availability of clusters around the world18 and the number of massive sequencing projects.

However, G-SAIP reduces the execution time for dot-plot calculation, generating dot-plots in times under 30 minutes for sequence size of <3 GB and speeding up the algorithm up to $3 \times$ times faster, increasing the number of CPUs. In the same way, G-SAIP can generate dot-plots of sequence greater than 3 GB in size, even *Pinus taeda* genome (21 GB) in a few hours in contrast with other software that does not support large files. Also, G-SAIP executes graphical aligners with chromosome sequences even $196 \times$ times faster than Gepard with detail in the output image because this tool does not apply any parallel strategy to calculate the graphical alignment.

A demonstrated application of dot-plots is to check assembly quality based on a genome reference.⁸⁵ For this reason, G-SAIP is also a helpful tool showing through dot-plots the



Figure 3. G-SAIP assemblies versus reference genome dot-plots: (a) MaSuRCA assembly for Arabidopsis, (b) Velvet k-mer 31 assembly for Arabidopsis, (c) MaSuRCA assembly for *Drosophila*, and (d) Velvet k-mer 91 assembly for Drosophila.

FIGURES	ORGANISM	QUALITY	N50	BUSCOS [%]
Figure 3a	Arabidopsis	High	110900	97.8823529
Figure 3b	Arabidopsis	Low	3110	8.7
Figure 3c	Drosophila	High	9973	95.6862745
Figure 3d	Drosophila	Low	2428	0

Table 4. N50 values and BUSCO's percentages for best and worst assemblies.

similarity of respective assemblies against the reference genome using ragtag⁵⁹ tool to make a previous ordering with the reference and complementing results given by other metrics such as BUSCO and N50 score. For example, in Figure 3a a diagonal line across the image is drawn, which shows that *the Arabidopsis* reference genome and the MaSuRCA⁷¹ assembly are close similar, squared regions marked in some points of the images are telomeres and centromeres of chromosomes, these regions are difficult to assemble and are masked by N. This similarity between no recognized regions in assembly and reference genomes are translated in black marked zones in dot-plot and are related to higher values of N50 and percentage of complete and single-copy BUSCOs. In contrast, Figure 3d is no diagonal lines in the image; this denotes that the assembly had very short contigs, and it was not possible to rebuild any part of the genome in relation to N50 value and percentage of BUSCOs that has low values for this assembly.

Moreover, comparative genomic analysis tools are essential to characterize genomes and sequences, focusing on variations between genomes of 2 or more individuals.⁸⁶ However, visualizing this data is not easy, and it would be even more complex with large sequences because the standard tools to make comparative genomic generate text files as results.⁸⁷ A first approximation for dot-plot usage in comparative genomics with software like DOTTER restricted the size of sequences and execution times required for these tasks. In this way, G-SAIP offers the possibility of handling large sequences (up to 21 GB) and executing fast graphical alignment with high quality (in order of minutes or few hours) even in multiples nodes with distributed memory and few latencies compared to 1 node execution. Furthermore, this tool provides a new way to visualize the similarity between sequences, allowing tuning option as similarity percentage to show repetitive sections, deletions, insertion, and rearrangement given by speciation events and delivering a result more comprehensively for researchers.

Conclusions

Currently, there is a necessity for tools to process large-scale genomic data sets in short periods. HPC clusters are growing

worldwide in computational capacity offering the opportunity to researchers to process bigger and more complex information. G-SAIP is a novel graphical sequence aligner able to produce dot-plots at a genomic scale, using the computational resources available in HPC clusters and the data produced by massive sequencing projects.^{53,54} Due to the parallel strategy used by the software and the scalability provided by HPC techniques, G-SAIP can accelerate the graphical alignments up to $1.68 \times$ times than other current software tested. This tool provides the opportunity to analyze, at a genomic scale, complete genomes with sizes up to 21 GB (*Pinus taeda*), facilitating processes such as assembly quality checking, comparative genomics, and identification of structures in DNA as transposable elements and other repetitive sequences.



Figure 4. Dot-plots for X mammals' chromosomes.



Figure 5. Boxplot of G-SAIP execution times over several nodes with 62 CPUs. This experiment was tested in computing cluster that has its queue with SCLURM scheduler.





Author Contributions

The authors confirm contribution to the paper as follows:

Johan S. Piña conceived and designed the software, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.

Simon Orozco-Arias conceived and designed the software, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the article, and approved the final draft.

Nicolas Tobón-Orozco analyzed the data, prepared figures and/or tables, and approved the final draft.

Leonardo Camargo-Forero contributed to the design of the parallel architecture for the software.

Reinel Tabares-Soto analyzed the data, authored or reviewed drafts of the article, and approved the final draft.

Romain Guyot conceived and designed the experiments, authored or reviewed drafts of the article, and approved the final draft.

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Supplemental material

Supplemental material for this article is available online.

REFERENCES

- López-Gartner G, Agudelo-Valencia D, Castaño S, et al. Identification of a putative ganoderic acid pathway enzyme in a ganoderma australe transcriptome by means of a hidden Markov model. In: Overbeek R, Rocha MP, Fdez-Riverola F, de Paz JF, eds. 9th International Conference on Practical Applications of Computational Biology and Bioinformatics, Spain, 3-5 June 2015. Springer International Publishing; 2015:107-115.
- Arango-López J, Orozco-Arias S, Salazar JA, Guyot R. Application of data mining algorithms to classify biological data: the Coffea Canephora genome case. In: Solano A, Ordoñez H, eds. *Advances in Computing*. Springer International Publishing; 2017:156-170.
- Borozan I, Watt S, Ferretti V. Integrating alignment-based and alignment-free sequence similarity measures for biological sequence classification. *Bioinformatics*. 2015;31:1396-1404.
- Luscombe NM, Greenbaum D, Gerstein M. A Proposed definition and overview of the field. *Methods Inform Med.* 2001;40:346-358.
- Gamermann D, Montagud A, Alberto Conejero J, de Córdoba PF, Urchueguía JF. Large scale evaluation of differences between network-based and pairwise sequence-alignment-based methods of dendrogram reconstruction. *PLoS One*. 2019;14:1-13.
- Clark C, Kalita J. A comparison of algorithms for the pairwise alignment of biological networks. *Bioinformatics*. 2014;30:2351-2359.
- Edgar RC, Batzoglou S. Multiple sequence alignment. Curr Opinion Struct Biol. 2006;16:368-373.
- Duret L, Gasteiger E, Perriere G. Lalnview: a graphical viewer for pairwise sequence alignments. *Bioinformatics*. 1996;12:507-510.
- Sonnhammer ELL, Durbin R. A dot-matrix program with dynamic threshold control suited for genomic DNA and protein sequence analysis. *Gene.* 1995; 167:GC1.
- Gibbs AJ, Mcintyre GA. The diagram, a method for comparing sequences: its use with amino acid and nucleotide sequences. *EurJ Biochem.* 1970;16:1-11.
- Trelles-Salazar O, Zapata EL, Dopazo J, Coulson AFW, Carazo JM. An imageprocessing approach to dotplots: an x-window-based program for interactive analysis of dotplots derived from sequence and structural data. *Bioinformatics*. 1995;11:301-308.
- zu Siederdissen CH, Prohaska SJ, Stadler PF. Algebraic dynamic programming over general data structures. *BMC Bioinformatics*. 2015;16:1-13.
- 13. Brodie R, Roper RL, Upton C. JDotter: a Java interface to multiple dotplots generated by dotter. *Bioinformatics*. 2004;20:279-281.

- Junier T, Pagni M. Dotlet: diagonal plots in a web browser. *Bioinformatics*. 2000; 16:178-179.
- Szafranski K, Jahn N, Platzer M. tuple_plot: fast pairwise nucleotide sequence comparison with noise suppression. *Bioinformatics*. 2006;22:1917-1918.
- Krumsiek J, Arnold R, Rattei T. Gepard: a rapid and sensitive tool for creating dotplots on genome scale. *Bioinformatics*. 2007;23:1026-1028.
- Manber U, Myers G. Suffix arrays: a new method for on-line string searches. SIAM J Comput. 1993;22:935-948.
- 18. Trelles O. Dotplots: dealing with Big-Data. Subbmited to ECCB; 2012.
- Seibt KM, Schmidt T, Heitkam T. FlexiDot: highly customizable, ambiguityaware dotplots for visual sequence analyses. *Bioinformatics*. 2018;34:3575-3577.
- Cabanettes F, Klopp C. D-GENIES: dot plot large genomes in an interactive, efficient and simple way. *PeerJ*. 2018;2018:e4958.
- Li H. Minimap2: pairwise alignment for nucleotide sequences. *Bioinformatics*. 2018;34:3094-3100.
- Jain C, Dilthey A, Koren S, Aluru S, Phillippy AM. A fast approximate algorithm for mapping long reads to large reference databases. *J Comput Biol.* 2018;25:766-779.
- Quail MA, Smith M, Coupland P, et al. A tale of three next generation sequencing platforms: comparison of Ion Torrent, Pacific Biosciences and Illumina MiSeq sequencers. *BMC Genomics*. 2012;13:341.
- Rishishwar L, Wang L, Clayton EA, Mariño-Ramírez L, McDonald JF, Jordan IK. Population and clinical genetics of human transposable elements in the (post) genomic era. *Mob Genet Elements*. 2017;7:1-20.
- Auerbach D, Thaminy S, Hottiger MO, Stagljar I. The post-genomic era of interactive proteomics: facts and perspectives. *Proteomics*. 2002;2:611-623.
- Ow DW. Recombinase-directed plant transformation for the post-genomic era. *Plant Mol Biol.* 2002;48:183-200.
- Medini D, Serruto D, Parkhill J, et al. Microbiology in the post-genomic era. Nat Rev Microbiol. 2008;6:419-430.
- Greene CS, Tan J, Ung M, Moore JH, Cheng C. Big data bioinformatics. J Cell Physiol. 2014;229:1896-1900.
- Orozco Arias S, Isaza G, Guyot R. Retrotransposons in plant genomes: structure, identification, and classification through bioinformatics and machine learning. *Int J Mol Sci.* 2019;20:1-31.
- Khan AA, Hassan L, Ullah S. Open MP-based parallel and scalable genetic sequence alignment. J Eng Appl Sci. 2015;34:29-34.
- Khaitan SK. A survey of high-performance computing approaches in power systems. *IEEE Power and Energy Society General Meeting, Boston, MA*, 17-21 July, 2016. IEEE.
- Procopiou AT, Quiros-Tortos J, Ochoa LF. HPC-based probabilistic analysis of LV networks with EVs: impacts and control. *IEEE Trans Smart Grid.* 2017;8: 1479-1487.
- 33. Fox GC, Qiu J, Kamburugamuve S, Jha S, Luckow A. HPC-ABDS high performance computing enhanced apache big data stack. *Proceedings 2015 IEEE/ACM 15th International Symposium on Cluster, Cloud, and Grid Computing, CCGrid 2015, Shenzhen, China, 4-7 May, 2015.* IEEE; 2015:1057-1066.
- Jha S, Fox G. Understanding ML driven HPC: applications and infrastructure. Proceedings - IEEE 15th International Conference on eScience, eScience 2019, San Diego, CA, 24-27 September, 2019. IEEE; 2019:421-427.
- Tabares Soto R. Programación paralela sobre arquitecturas heterogéneas. 2016: 80. Accessed September 1, 2020. http://www.bdigital.unal.edu.co/54267/
- Orozco-Arias S, Tabares-Soto R, Ceballos D, Guyot R. Parallel programming in biological sciences, taking advantage of supercomputing in genomics. In: Solano A, Ordoñez H, eds. *Advances in Computing*. Springer International Publishing; 2017:627-643.
- Mikailov M, Luo F jyh, Barkley S, et al. Scaling bioinformatics applications on HPC. BMC Bioinformatics. 2017;18:501.
- Li J kun, Zhang L, Xiao M. The high performance computing applications for bioinformatics research. ICBBS'17: 6th International Conference on Bioinformatics and Biomedical Science, Singapore, Singapore, 22-24 June, 2017. ACM; 2017:1-6.
- Rucci E, Garcia C, Botella G, de Giusti A, Naiouf M, Prieto-matias M. Accelerating Smith-Waterman alignment of long DNA sequences with OpenCL on FPGA. *Bioinform Biomed Eng.* 1900;2:500-511.
- Ren S, Ahmed N, Bertels K, Al-Ars Z. GPU accelerated sequence alignment with traceback for GATK HaplotypeCaller. *BMC Genomics*. 2019;20:184.
- Milne I, Lindner D, Bayer M, et al. TOPALi v2: a rich graphical interface for evolutionary analyses of multiple alignments on HPC clusters and multi-core desktops. *Bioinformatics*. 2009;25:126-127.
- Borovska P, Gancheva V, Georgiev I. Hybrid parallel implementation of multiple sequence alignment software ClustalW on Intel Xeon Phi. Sixth International Conference on Advances in Computing, Electronics and Communication - ACEC 2017, Rome, Italy, 9-10 December, 2017.
- Kim J, Warnow T. PASTA: ultra-large multiple sequence alignment for nucleotide and amino-acid sequences. J Comput Biol. 2015;22:377-386.
- Orobitg M, Guirado F, Cores F, Llados J, Notredame C. High performance computing improvements on bioinformatics consistency-based multiple sequence alignment tools. *Parallel Comput.* 2015;42:18-34.

- Lassmann T. Kalign 3: multiple sequence alignment of large datasets. Bioinformatics. 2019;36:1928-1929.
- Rodrigues FM, von Mering C. Sequence analysis HPC-CLUST: distributed hierarchical clustering for large sets of nucleotide sequences. *Bioinformatics*. 2014;30:287-288.
- Sawyer SE, Drive D, Horton MD, Brook RG. HPC-BLAST: distributed BLAST for Xeon Phi clusters categories and subject descriptors. *BCB'15*, *Atlanta, GA, 9-12 September, 2015.* ACM; 2015:512-513.
- Driscoll AO, Belogrudov V, Carroll J, et al. HBLAST: parallelised sequence similarity: a Hadoop MapReducable basic local alignment search tool. *J Biomed Inform.* 2015;54:58-64.
- Nowicki M, Bzhalava D, BaŁa P. Massively parallel implementation of sequence alignment with basic local alignment search tool using parallel computing in Java library. J Comput Biol. 2018;25:871-881.
- Zhang J, Lan H, Chan Y, Shang Y, Schmidt B, Liu W. BGSA: a bit-parallel global sequence alignment toolkit for multi-core and many-core architectures. *Bioinformatics*. 2018;35:2306-2308.
- Orozco Arias S, Liu J, Tabares Soto R, et al. Inpactor, integrated and parallel analyzer and classifier of LTR retrotransposons and its application for pineapple LTR retrotransposons diversity and dynamics. *Biology*. 2018;7:32.
- Tamayo M, Tabares R, Montes N. Three-dimensional indexing in GPU for numerical approximation of solutions of the Laplace equation. *Rev Antioq.* 2015;5:37-42.
- Cheng S, Melkonian M, Smith SA, et al. 10KP: a phylodiverse genome sequencing plan. *Gigascience*. 2018;7:giy013.
- Lewin HA, Robinson GE, Kress WJ, et al. Earth BioGenome project: sequencing life for the future of life. *Proc Natl Acad Sci USA*. 2018;115:4325-4333.
- 55. van Rossum G, Drake FL. The Python Language Reference Manual. Network Theory Ltd.; 2011.
- Dalcín L, Paz R, Storti M, D'Elía J. MPI for python: performance improvements and MPI-2 extensions. J Parallel Distrib Comput. 2008;68:655-662. doi:10.1016/j.jpdc.2007.09.005
- Jain C, Koren S, Dilthey A, Phillippy AM, Aluru S. A fast adaptive algorithm for computing whole-genome homology maps. *Bioinformatics*. 2018;34:i748-i756.
- Bruck J, Dolev D, Ho CT, Roşu MC, Strong R. Efficient message passing interface (MPI) for parallel computing on clusters of workstations. *J Parallel Distrib Comput.* 1997;40:19-34.
- Graham RL, Shipman GM, Barrett BW, Castain RH, Bosilca G, Lumsdaine A. Open MPI: a high-performance, heterogeneous MPI. 2006 IEEE International Conference on Cluster Computing, Barcelona, Spain, 25-28 September 2006. IEEE; 2006:1-9.
- Alonge M, Soyk S, Ramakrishnan S, et al. RaGOO: fast and accurate referenceguided scaffolding of draft genomes. *Genome Biol.* 2019;20:1-17.
- Jette M, Yoo A, Grondona M. SLURM: simple linux utility for resource management. In: Feitelson D, Rudolph L, Schwiegelshohn U, eds. *Job Scheduling Strategies for Parallel Processing. JSSPP 2003. Lecture Notes in Computer Science.* Vol 2862. Springer; 2003:44-60.
- Law WD, Warren RL, McCallion AS. Establishment of an eHAP1 human haploid cell line hybrid reference genome assembled from short and long reads. *Genomics*. 2020;112:2379-2384.
- Gornicki P, Zhu H, Wang J, et al. The chloroplast view of the evolution of polyploid wheat. *New Phytol.* 2014;204:704-714.
- Neale DB, Wegrzyn JL, Stevens KA, et al. Decoding the massive genome of loblolly pine using haploid DNA and novel assembly strategies. *Genome Biol.* 2014;15:R59.

- Leinonen R, Sugawara H, Shumway M, Collaboration INSD. The sequence read archive. *Nucleic Acids Res.* 2010;39:D19-D21.
- Andrews S, Krueger F, Segonds-Pichon A, Biggins L, Krueger C, Wingett S. FastQC. Babraham Institute; 2012.
- 67. Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for illumina sequence data. *Bioinformatics*. 2014;30:2114-2120.
- Li D, Liu CM, Luo R, Sadakane K, Lam TW. MEGAHIT: an ultra-fast singlenode solution for large and complex metagenomics assembly via succinct de Bruijn graph. *Bioinformatics*. 2015;31:1674-1676.
- Zerbino DR, Birney E. Velvet: algorithms for de novo short read assembly using de Bruijn graphs. *Genome Res.* 2008;18:821-829.
- Simpson JT, Wong K, Jackman SD, Schein JE, Jones SJM, Birol I. ABySS: a parallel assembler for short read sequence data. *Genome Res.* 2009;19: 1117-1123.
- Zimin AV, Marçais G, Puiu D, Roberts M, Salzberg SL, Yorke JA. The MaSuRCA genome assembler. *Bioinformatics*. 2013;29:2669-2677.
- Seppey M, Manni M, Zdobnov EM. BUSCO: assessing genome assembly and annotation completeness. *Methods Mol Biol.* 2019;1962:227-245.
- Mikheenko A, Prjibelski A, Saveliev V, Antipov D, Gurevich A. Versatile genome assembly evaluation with QUAST-LG. *Bioinformatics*. 2018;34:i142-i150.
- Pucker B, Holtgräwe D, Rosleff Sörensen T, Stracke R, Viehöver P, Weisshaar B. A De Novo genome sequence assembly of the Arabidopsis thaliana Accession Niederzenz-1 displays presence/absence variation and strong synteny. *PLoS One*. 2016;11:e0164321.
- 75. Hoskins RA, Carlson JW, Wan KH, et al. The release 6 reference sequence of the Drosophila melanogaster genome. *Genome Res.* 2015;25:445-458.
- Murphy WJ, Pevzner PA, O'Brien SJ. Mammalian phylogenomics comes of age. Trends Genet. 2004;20:631-639.
- 77. Ross MT, Grafham D v, Coffey AJ, et al. The DNA sequence of the human X chromosome. *Nature*. 2005;434:325-337.
- Lindblad-Toh K, Wade CM, Mikkelsen TS, et al. Genome sequence, comparative analysis and haplotype structure of the domestic dog. *Nature*. 2005;438: 803-819.
- 79. Fang X, Mou Y, Huang Z, et al. The sequence and analysis of a Chinese pig genome. *Gigascience*. 2012;1:16.
- Wade CM, Giulotto E, Sigurdsson S, et al. Genome sequence, comparative analysis, and population genetics of the domestic horse. *Science*. 2009;326:865-867.
- Church DM, Goodstadt L, Hillier LW, et al. Lineage-specific biology revealed by a finished genome assembly of the mouse. *PLoS Biol.* 2009;7:e1000112.
- Prüfer K, Munch K, Hellmann I, et al. The bonobo genome compared with the chimpanzee and human genomes. *Nature*. 2012;486:527-531.
- Gene DR, Amdahl M. Validity of the single processor approach to achieving large scale computing capabilities. *AFIPS Conference Proceedings – 1967 Spring Joint Computer Conference, AFIPS 1967, Atlantic City, NJ, 18 April 1967.* ACM; 1967:483-485.
- Orozco-arias S, Tobon-orozco N, Piña JS, Jiménez-Varón CF, Tabares-Soto R, Guyot R. TIP _ finder: an HPC software to detect transposable element insertion polymorphisms in large genomic datasets. *MDPI Biol.* 2020;9: 1-17.
- 85. Gustafson JL. Reevaluating Amdahl's law. Commun ACM. 1988;31:532-533.
- Husemann P, Stoye J. r2cat: synteny plots and comparative assembly. *Bio-informatics*. 2010;26:570-571.
- Zhang X, Wang J, Li J, Chen W, Liu C. CRlncRC: a machine learning-based method for cancer-related long noncoding RNA identification using integrated features. *BMC Med Genomics*. 2018;11:120.