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S-plot2: Rapid Visual and Statistical Analysis of Genomic Sequences

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ABSTRACT: With the daily release of data from whole genome sequencing projects, tools to facilitate comparative studies are hard-pressed to keep pace. Graphical software solutions can readily recognize synteny by measuring similarities between sequences. Nevertheless, regions of dissimilarity can prove to be equally informative; these regions may harbor genes acquired via lateral gene transfer (LGT), signify gene loss or gain, or include coding regions under strong selection. Previously, we developed the software S-plot. This tool employed an alignment-free approach for comparing bacterial genomes and generated a heatmap representing the genomes’ similarities and dissimilarities in nucleotide usage. In prior studies, this tool proved valuable in identifying genome rearrangements as well as exogenous sequences acquired via LGT in several bacterial species. Herein, we present the next generation of this tool, S-plot2. Similar to its predecessor, S-plot2 creates an interactive, 2-dimensional heatmap capturing the similarities and dissimilarities in nucleotide usage between genomic sequences (partial or complete). This new version, however, includes additional metrics for analysis, new reporting options, and integrated BLAST query functionality for the user to interrogate regions of interest. Furthermore, S-plot2 can evaluate larger sequences, including whole eukaryotic chromosomes. To illustrate some of the applications of the tool, 2 case studies are presented. The first examines strain-specific variation across the Pseudomonas aeruginosa genome and strain-specific LGT events. In the second case study, corresponding human, chimpanzee, and rhesus macaque autosomes were studied and lineage specific contributions to divergence were estimated. S-plot2 provides a means to both visually and quantitatively compare nucleotide sequences, from microbial genomes to eukaryotic chromosomes. The case studies presented illustrate just 2 potential applications of the tool, highlighting its capability to identify and investigate the variation in molecular divergence rates across sequences. S-plot2 is freely available through https://bitbucket.org/lkalesinskas/splot and is supported on the Linux and MS Windows operating systems.

KEYWORDS: comparative genomics, alignment-free, gene transfer, gene loss

Background

Modern sequencing technologies can quickly and affordably produce genomic sequences for species across the tree of life. Consequently, many new lineages and poorly resolved areas of the tree have been identified.¹⁻³ With tens of thousands of bacterial genomes now publicly available, comparative genomics has produced numerous insights into microbial life.⁴ Several tools are currently used to detect genome similarity through sequence alignment.⁵⁻⁸ In addition, tools employing a graphical “dot plot” approach, such as Gepard,⁹ Serolis,¹⁰ and SeqTools’ Dotter,¹¹ can highlight genomic similarities and rearrangements as well as gene duplications. These tools, however, have their limitations: Serolis¹⁰ is limited in the size of sequence it can analyze (4kb), and Dotter¹¹ is significantly slower than Gepard for larger sequences. Nevertheless, alignment-free approaches, including the aforementioned “dot plot” tools, have a significant advantage over alignment-based methods: they are less computationally expensive (regarding both time and resources) and impervious to synteny-related problems (see reviews of Vinga and Almeida¹² and Bonham-Carter et al¹³).

In addition to sequence similarities, the dissimilarities between genomic sequences can be equally informative.¹⁴ These dissimilarities can indicate strain-specific genes horizontally/laterally acquired rather than vertically inherited. Lateral gene transfer (LGT) is an important force in the evolution of prokaryotes,¹⁵ including the exchange of defense mechanisms and virulence factors.¹³,¹⁵,¹⁶ Although certainly less prevalent (and fiercely debated), LGT between eukaryotes and prokaryotes can also occur.¹⁷⁻²⁰ Disparities between genomic sequences can also be the result of gene loss, another pervasive and often significant driver of evolution in prokaryotic¹²¹,²² and eukaryotic species²³ (see review Albalat and Cañestro²⁴). Moreover, recognition of substantial sequence divergence between orthologous gene sequences can signify genes under strong selection (see review Long et al²⁵). Such genes can provide insight into phenotypic differences between species.²⁶,²⁷

Previously we developed S-plot, a tool for the rapid analysis and visualization of bacterial genomic sequences.²⁸ This tool was applied to the examination of Escherichia,²⁸ Bacillus,²⁹ and Neisseria³⁰ genomes, identifying regions of unusual nucleotide
composition corresponding to LGT events. Herein, we present
the next generation of this tool: S-plot2. Similar to its prede-
cessor, S-plot2 creates an interactive, 2-dimensional heatmap.
Similar to the aforementioned dot-plot tools, S-plot2 captures
the similarities in nucleotide usage between genomic sequences,
but unique to this tool is the fact that it also captures the dis-
similarities in nucleotide usage between genomic sequences.
Through the examination of nucleotide usage, phylogenetic
signals can be uncovered.31 In S-plot2, whole eukaryotic chro-
somes and smaller prokaryotic genomes can be efficiently
compared. Furthermore, the new version includes functionality
to extract, analyze, and automate BLAST queries of regions of
interest within the heatmap. This facilitates the investigation of
quickly evolving coding regions, novel coding regions, and lat-
erally transferred elements.

Implementation
Developed in Java, S-plot2 performs pairwise comparisons of
genomic sequences (partial or complete) via a sliding window
approach. Windows can be of a user-defined length (the
“genome approach”) or confined to annotated coding regions
(the “gene-by-gene approach”). The “genome approach” per-
mits windows to be either adjacent or overlapping. Regardless
of the approach selected, each window’s $k$-mer (subsequence of
length $k$) frequencies are enumerated. The similarity/dissimi-
larity between 2 windows is calculated based on these $k$-mer
frequencies, using either the Pearson ($r$) or Spearman rank ($\rho$
) correlation coefficient. The resulting values for each pairwise
window comparison are then graphed as a 2-dimensional heat-
map using Glimpse32 (eg, Figure 1A). Windows with a similar
$k$-mer usage are represented in the heatmap using colors at one
end of the color spectrum, whereas windows with dissimilar
$k$-mer usage are represented by colors at the other end of the
spectrum. Draft genome sequences that include several scaf-
dfold sequences can be examined using the “genome approach”
in S-plot2. The scaffolds can be concatenated, separated by, eg,
N5s, into a single FASTA sequence. S-plot2 does not calculate
frequencies for windows in which greater than half of the
sequence is not A, T, C, or G; thus, windows containing more
than one scaffold will be ignored. The “gene-by-gene approach”
is a new feature released in S-plot2, as is the Spearman rank
correlation coefficient metric for sequence comparison.

Functionality has been developed in S-plot2 to aid in the
interpretation of the heatmap. Users can specify regions of
interest based on window coordinates or select windows meet-
ing specific criteria (eg, regions exhibiting aberrant $k$-mer
usage) and then output or BLAST33 these regions. For instance,
a cluster of genes which appears in one genome and not the

Figure 1. Comparison of *Pseudomonas aeruginosa* PAO1 (x-axis) and PA7 (y-axis) genomes. (A) “Genome approach” comparison with a window and
offset of 5000bp. (B) Genomic island present with the PA7 strain. (C) “Gene-by-gene approach” comparison of protein-coding gene sequences annotated
for the 2 genomes in panel A (*.faa files). Here, the window size is equivalent to a single coding region and $k=3$ is evaluated (the same color bar as shown
in panel A). The comparisons conducted here for both approaches were done using the Pearson correlation coefficient. Sequence similarity is measured
by the frequency of shared $k$-mers, with green signifying low similarity and red signifying high similarity.
other (indicative of a gene loss/gain), such as that shown in Figure 1B, can be queried; in the case in which a gene was acquired via LGT, the putative source can be identified. Queries to National Center for Biotechnology Information's (NCBI) eUtils API were automated using JEutils. All BLAST queries in S-plot2 use the blastn algorithm and remotely query the NCBI nucleotide collection (nt/nt) database. Users can also output statistics computed for the heatmap as well as generate multi-FASTA format files for windows with an $r$ or $p$ value within a user-defined range. The heatmap image itself can be saved to file as a TIF file, implemented using the iCafe package.

An executable jar file, sample sequence data, and a tutorial are freely available through https://bitbucket.org/lkalesinskas/splot. S-plot2 was tested thoroughly on the Windows and Ubuntu operating systems. Due to the lack of support for compatibility profiles on MacOS, rendering and maneuvering within the S-plot2 heatmap are suboptimal (due to incompatibilities with the Glimpse visualization version used) on MacOS. Exploration of the S-plot2 heatmap (scrolling through a sequence, zooming in/out, etc) was optimized for use with the mouse on Windows and Ubuntu.

As the similarity between windows is calculated based on the correlation (either the Pearson or Spearman rank correlation coefficient) of the frequency of shared $k$-mers, the condition $4^k \leq M/n$ where $w$ is the window size must be followed. If $w > 4^k$, then most $k$-mer frequencies will be 0 or 1 and thus unsuitable for the correlation analysis. Run time and memory usage are dependent on the number of windows. For a genome of size $M$, the number of windows, $n$, is $M/w$. Thus, for smaller window sizes, a larger heatmap will be generated. For each window, $k$-mer frequencies are enumerated for the original and reverse-complement sequences. Calculation of $k$-mer usage is linear and values are stored in a sorted array. The run time and memory usage estimate is $O(n^2)$. For instance, the heatmap generated in Figure 1A was generated in 42 seconds. It is important to note that for large sequences, the required RAM may exceed the RAM allocated or available for the Java Virtual Machine (particularly if the user has a 32-bit version installed) in which case the application will not execute. Nevertheless, a complete human chromosome can be compared using less than 8 GB of RAM in a matter of minutes; S-plot2's performance is significantly faster than other graphical alignment-free available graphical tools.

Results and Discussion

To illustrate the functionality and utility of S-plot2, we conducted 2 case studies. In addition to providing a visualization of the genomic sequences under investigation, the new functionality developed in S-plot2 can lead to a deeper understanding of the variation in molecular divergence rates across sequences.

Case study 1: exploring the evolution of bacterial genomes

The genomes of the opportunistic bacterial pathogen Pseudomonas aeruginosa are highly mosaic and include regions of genomic plasticity. The P. aeruginosa accessory genome exceeds that of its core genome. Figure 1 shows the pairwise comparison of the P. aeruginosa strains PAO1 (NC_002516) and the known “taxonomic outlier” for the species, PA7 (NC_009656). Two comparisons were conducted: the “genome approach” using a fixed window size (Figure 1A) and the “gene-by-gene approach” in which each window is an individual gene (Figure 1C). As even closely related P. aeruginosa strains can be distinguished by single-nucleotide polymorphisms, indels, and inversions, it is thus not surprising to observe genomic variation between the PAO1 and PA7 genomes (Figure 1A and C). The nucleotide sequence of the PA7 region shown in Figure 1B was investigated using S-plot2’s automated BLAST functionality. This region includes numerous transposases and integrases as well as plasmid- and phage-associated genes. It corresponds to the previously identified genomic island RGP42 within the P. aeruginosa PA7 genome. The region shown in Figure 1B is but one of the many genomic islands within these 2 strains. Users can recognize windows of unusual composition visually via the “genome approach” or individual genes of interest via the “gene-by-gene approach” and BLAST the sequences. Furthermore, S-plot2 can automatically identify such regions and BLAST their sequences.

Recombination within P. aeruginosa species is frequent and previous research has found variation in the evolutionary histories of regions of the P. aeruginosa genome. To exemplify how S-plot2 can be used to investigate recombination, 7 genomes included in the comparative genomic study of Dettman et al were selected (Table 1) and pairwise comparisons were performed. Sequence similarity was assessed for each window size of 5000 bp (base pairs) for $k = 6$ using the Pearson correlation coefficient. Figure 2 (panels B, C, and D) shows the pairwise comparisons for PAO1 and C3719, LESB58, and PACS2, respectively. These heatmaps illustrate the presence/absence of unique regions within the genomes and, most notably, rearrangements. The matrices generated by S-plot2 were saved and contiguous 0.2 Mbp regions along the PAO1 genome were evaluated. Thus, an alignment-free approach was used to identify and quantify similarity/dissimilarity between homologous regions of the PAO1 genome and other P. aeruginosa strains. As shown in Figure 2A, different regions of the PAO1 genome are represented by different topologies. Consistent with prior alignment-based analyses, we find that the evolution of the P. aeruginosa genome is not uniform across the entire genome sequence. In this fashion, S-plot2 can provide evidence of evolution across a genome sequence both visually and quantitatively.
Table 1. Seven *Pseudomonas aeruginosa* genomes examined.

<table>
<thead>
<tr>
<th>STRAIN</th>
<th>GENOME SIZE, MBP</th>
<th>NO. OF SCAFFOLDS</th>
<th>NO. OF CODING REGIONS</th>
<th>ASSEMBLY</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAO1</td>
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<td>1</td>
<td>5572</td>
<td>GCA_000006765</td>
</tr>
<tr>
<td>LESB58</td>
<td>6.60</td>
<td>1</td>
<td>6041</td>
<td>GCA_000026645</td>
</tr>
<tr>
<td>C3719</td>
<td>6.22</td>
<td>1</td>
<td>5648</td>
<td>GCA_000152525</td>
</tr>
<tr>
<td>PACS2</td>
<td>6.49</td>
<td>1</td>
<td>5913</td>
<td>GCA_000168335</td>
</tr>
<tr>
<td>JD316</td>
<td>6.19</td>
<td>1882</td>
<td>6590</td>
<td>GCA_000506125</td>
</tr>
<tr>
<td>JD317</td>
<td>6.49</td>
<td>2043</td>
<td>6979</td>
<td>GCA_000506145</td>
</tr>
<tr>
<td>JD320</td>
<td>6.41</td>
<td>2038</td>
<td>6876</td>
<td>GCA_000506165</td>
</tr>
</tbody>
</table>

Sequences were retrieved for genomes (*.genomic.fna.gz) and coding sequences (*.cds_from_genomic.fna.gz).

Figure 2. Evolution of the *Pseudomonas aeruginosa* chromosome. (A) Comparison of cluster topologies based on sequence similarity based on 6-mer usage for window size = offset size = 5000 bp over 0.2 Mbp regions of the PAO1 genome. Heatmaps for (B) PAO1 vs C3719, (C) LESB58, and (D) PACS2. The same color scale as Figure 1 is used here: sequence similarity is measured by the frequency of shared k-mers, with green signifying low similarity and red signifying high similarity.
Case study 2: exploring the evolution of primate chromosomes

S-plot2 is also capable of evaluating whole eukaryotic chromosomes. As such, it can be used to estimate chromosome-specific molecular divergence rates, estimate lineage-specific contributions to divergence, and identify regions that are significant contributors to observed divergence. As a case study of S-plot2, we performed pairwise comparisons for all homologous human, chimpanzee, and rhesus autosomes (window size = offset size = 100 Kbp for \( k = 6 \) using the Pearson correlation coefficient). Each chromosome was also compared with itself using the same window size, offset size, and \( k \). This self-sequence comparison provides a baseline for the variation within a chromosome relative to that observed between species (see Supplemental File 1). Prior whole genome comparison studies between human and chimpanzee found \( \approx 1.4\% \) sequence divergence\(^{43} \) and 23 inversions,\(^{44} \) as well as other differences (for a review, see the work by Kehrer-Sawatzki and Cooper\(^{45} \)). Sequence analysis of human-chimpanzee chromosome pairs suggests that recombination, proximity to telomeres, bias in repair mechanisms, and GC content are all exerting influence on genetic variation.\(^{46–50} \)

Here, we present a comparison between human chromosome 17, chimpanzee chromosome 17, and rhesus chromosome 16. As the heatmaps in Figure 3 show, the pericentric inversion previously found between these sequences\(^{44} \) can be identified through the pairwise comparisons of the human, chimpanzee, and rhesus autosomes. The heatmaps for these 3 pairwise comparisons, however, do not readily present how

Figure 3. Comparison of human (Homo sapiens) chromosome 17 (H17), chimpanzee (Pan troglodytes) chromosome 17 (C17), and rhesus (Macaca mulatta) chromosome 16 (R16). Sequence similarity is measured by the frequency of shared \( k \)-mers, with green signifying low similarity and red signifying high similarity. The inlay shows the divergence between H17 and C17 (red), C17 and R16 (yellow), and H17 and R16 (blue), relative to the window's GC content. The \( x \)-axis is representative of the divergence calculated for a window relative to its GC content.
these chromosomes are evolving. For instance, the differences observed between the homologous human and chimpanzee chromosomes may be the result of changes within the chimpanzee chromosome or changes within the human chromosome. Comparisons of both chromosomes to the rhesus chromosome let us distinguish between these 2 scenarios. If we oversimplify the process of species divergence to a single point in time (thus ignoring subsequent gene flow), one could assume that the chromosomal sequences are essentially identical. Thus, for a window in the human chromosome, its homologous window in the chimpanzee genome would have the same sequence (and thus nucleotide composition). As such, the heatmap for an individual chromosome compared with itself would be indiscernible from the comparison of the chromosome to its homolog. Post-spe­ciation, the 2 genomes would begin to diverge and this divergence can be quantified by the cross­species comparison value (eg, human vs chimpanzee) relative to the intraspecies comparison (eg, human vs human). The matrices of r values were retrieved for each of the plots shown in Figure 3 and used to calculate the divergence between species (see Supplemental File 1 for details regarding this calculation). The inlay in Figure 3 shows the results of this calculation for human vs chimpanzee (red), chimpanzee vs rhesus (yellow), and human vs rhesus (blue). In this figure, the x-axis is representative of the divergence calculated for a window relative to its GC content. As shown in the inlay in Figure 3, regions in the human genome with a GC content ≈45% are the most divergent windows from chimpanzee; these regions are evolving within the human lineage.

**Conclusions**

S-plot2 provides a means to visually and quantitatively compare genomic sequences ranging from microbial genomes to eukaryotic chromosomes. These comparisons can be generated in a matter of seconds to minutes (depending on the size of the sequence under consideration). S-plot2 includes functionality to aid in the analyses of genomic sequences, allowing users to quickly investigate their data and test hypotheses based on either observed patterns or statistics capturing both the similarities and dissimilarities of sequences. The case studies presented highlight just some of the applications of S-plot2. Furthermore, the analyses performed for the *Pseudomonas* genomes and human-chimpanzee-rhesus autosomes illustrate the variation in molecular divergence rates across sequences.

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**Author Contributions**

LK and EC implemented the application. YF and CP designed the project. LK, EC, and CP performed the analyses of the case studies. CP was a major contributor in writing the manuscript. All authors read and approved the final manuscript.

**REFERENCES**


