

Genome analysis

PanViz: Interactive Visualization of the Structure of Functionally Annotated Pangenomes

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Abstract

Summary: PanViz is a novel, interactive, visualization tool for pangenome analysis. PanViz allows visualization of changes in gene group classification as different subsets of pangenomes are selected, as well as comparisons of individual genomes to pangenomes with gene ontology based navigation of gene groups. Furthermore it allows for rich and complex visual querying of gene groups in the pangenome. PanViz visualizations require no external programs and are easily sharable, allowing for rapid pangenome analyses.

Availability and implementation: PanViz is written entirely in JavaScript and is available on <https://github.com/thomasp85/PanViz>. A companion R package that facilitates the creation of PanViz visualizations from a range of data formats is released through Bioconductor and is available at <https://bioconductor.org/packages/PanVizGenerator>.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Visualization plays an integral part in modern biology research, as the size and complexity of biological data has increased (Land *et al.*, 2015). Many new tools for analysis of pangenomes have recently been published (Rokicki *et al.*, 2014; Lechat *et al.*, 2012; Grant *et al.*, 2012; Hallin *et al.*, 2008). However, several of these do not scale well. A general tendency for pangenome visualizations is to use the chromosome of a reference genome as an axis and plot synteny between genomes along that. Visualization based on a reference genome fails to take into account novel pan-genes not in the reference. As more genomes are added, the reference genome becomes less representative of the full dataset. Some attempts have been made to create reference free, scalable pangenome visualization of different types. GenoSets (Cain *et al.*, 2012) is a visualization that uses parallel sets to facilitate gene group selections based on presence-absence in pangenome subsets and GenomeRing (Herbig *et al.*, 2012)

tries to overcome the reference bias by merging all chromosomes into a superchromosome that can be used as a backbone for visualization. Here we present a new interactive visualization, PanViz, aimed at letting users explore the structure of functionally annotated pangenomes and pangenome subsets, while performing visual queries to search for gene groups. PanViz is based purely on GO annotation and the presence/absence pattern of gene groups, and is thus not dependent on a single reference genome

2 Implementation

PanViz is written entirely in JavaScript using D3 (Bostock *et al.*, 2011). It is completely self-contained, embedded in a single HTML file, and does not require any connection to external sources. A companion R package, PanVizGenerator, has been released on Bioconductor (Gentleman *et al.*, 2003; Huber *et al.*, 2015) that allows for the creation of PanViz visualizations from a range of different data types, including the

pangenome classes defined in FindMyFriends (Pedersen, 2015). The gene groups in the pangenome must be annotated using Gene Ontology (GO) (Ashburner *et al.*, 2000), using e.g. Blast2GO (Conesa *et al.*, 2005) on representatives for each gene group. Example data, the resulting PanViz and a tutorial for PanVizGenerator are available in the supplementary material.

3 Overview

PanViz consists of four main areas (see figure S1 in supplementary material), with the main pangenome visualization in the middle. The left part is reserved for (pan)genome navigation, while the right part is for legends and additional lookup information. The bottom part contains a list of all the currently selected gene groups as well as tools to modify the selection.

3.1 Genome overview

The genomes in the pangenome are represented by a dual linked view with a principal component analysis / multidimensional scaling based scatterplot on top, and a hierarchical clustering in the bottom (figure S1A). Both views support selecting single genomes in order to transition into the genome-pangenome comparison state, and the dendrogram allows for selection of subsets of the pangenome by selecting the branch points of the dendrogram. The overview plots are also linked to the gene group table (figure S1C) so that all genomes containing the gene group currently hovered over will be highlighted.

3.2 Pangenome overview

The main view of the visualization is a radial representation of the 3 presence-based gene group groupings in the pangenome: Core, Accessory and Singleton gene groups (figure S1D). Each of these is furthermore divided based on the distribution of top level biological process GO terms. As different sub-pangenomes are selected, the changes in the pangenome are animated by moving sections of each GO arc around. After the animation ends the dynamics can furthermore be shown as chords when hovering over a specific GO arc.

3.3 Genome-pangenome comparison

When one or two genomes are selected the main view transitions into a stacked bar chart showing the pangenome in the middle (figure S1E). The genomes are represented by their GO term composition and weighted bezier curves connects the genes in the genomes to their location in the pangenome (if present). If two genomes are selected the proportion of each GO term they share with each other and the pangenome is visible as a darker shaded bezier curve.

3.4 Gene ontology navigation

To gain insight into the distribution of lower level GO terms a treemap weighted by the number of gene groups in the current pangenome having a specific term is available upon selecting a top level GO term bar from the pangenome (figure S1F). The treemap is zoomable and features descriptions of each included GO term.

3.5 Visual querying

As each visual element represent a set of gene groups it makes sense to build a querying mechanism based on set arithmetic (union, intersection, complement, etc.). An icon on top of the gene group table indicates the different set operations available (figure S1C). The operations will be performed between the current content of the table and the gene

groups contained in the visual element selected. Based on the six different operations it is possible to intuitively create very complex gene group queries guided by the insights gained from the visualization.

4 Conclusion

PanViz offers a novel and unbiased approach to visualizing the structure of pangenome data. Interactions and animations are utilized to invite users to investigate the data, and the reliance of a single self-contained HTML file makes it easy to share with fellow researchers. While the main visualization is fully scalable to thousands of gene groups as it relies on summaries, the genome navigation panel is currently limited to ~50 genomes in order for the leafs in the dendrogram to remain discernible. Future work will focus on improving the genome overview, as well as implementing state saving within the URL to facilitate sharing of different states of a PanViz visualization.

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