Genome analysis

GEnView: a gene-centric, phylogeny-based comparative genomics pipeline for bacterial genomes and plasmids

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Abstract

Summary: Comparing genomic loci of a given bacterial gene across strains and species can provide insights into their evolution, including information on e.g. acquired mobility, the degree of conservation between different taxa or indications of horizontal gene transfer events. While thousands of bacterial genomes are available to date, there is no software that facilitates comparisons of individual gene loci for a large number of genomes. GEnView (Genetic Environment View) is a Python-based pipeline for the comparative analysis of gene–loci in a large number of bacterial genomes, providing users with automated, taxon-selective access to the >800.000 genomes and plasmids currently available in the NCBI Assembly and RefSeq databases, and is able to process local genomes that are not deposited at NCBI, enabling searches for genomic sequences and to analyze their genetic environments through the interactive visualization and extensive metadata files created by GEnView.

Availability and implementation: GEnView is implemented in Python 3. Instructions for download and usage can be found at https://github.com/EbmeyerSt/GEnView under GLP3.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

With the advancement of sequencing technologies and their increasing efficiency, the number of sequenced bacterial genomes in publicly available databases has increased rapidly during the last two decades (Land et al., 2015). The NCBI Assembly database alone currently contains (November 2021) more than 800,000 bacterial assemblies (Kitts et al., 2016), which are used by researchers around the world to investigate questions across a variety of fields, from bacterial evolution to public health, including pathogen virulence and antibiotic resistance (Ebmeyer et al., 2021).

Though many comparative genomics tools are available to date for the analysis of bacterial replicons, such as GeneCO (Jung et al., 2019), BRIG (Alikhan et al., 2011) or MAUVE (Darling et al., 2004), most are developed for comparing full genomes and detecting evolutionary events such as insertions, deletions or rearrangements across those genomes, and are therefore limited in the number of genomes that are feasible to compare. Furthermore, these tools are not developed for comparative analysis of particular gene loci, which may be more suitable when researching e.g. the evolutionary history or taxonomic distribution of single bacterial genes. Such comparative analysis may require retrieval and extensive pre-processing of data, which is often a non-trivial undertaking.

Here, we present GEnView (Genetic Environment View), a fully automated, gene-centric pipeline for comparing genomic regions of several kilobasepairs that combines multiple bioinformatics tools and resources into a workflow that enables visual comparison of gene loci across hundreds of genomes stored locally or in the NCBI Assembly/RefSeq databases. GEnView identifies and processes user specified target genes from either user-provided nucleotide sequences or the genome and plasmid sequences stored in the NCBI Assembly/RefSeq database, creating an interactive visualization of all identified genes, including their phylogeny and their genetic environment. The main computational steps are parallelized which greatly reduces the run-time, making it possible to annotate several gene–loci at once. GEnView is therefore highly suitable for addressing questions about the evolutionary history of prokaryotic genes, such as the horizontal transfer and origins of mobile genes, or their potential (future) spread to other taxa.
within 5.9, 3.1, 1.8, 1.2, 1 and 0.6 h respectively, using an identity cutoff toward the FOX-1 reference protein of 80% (Supplementary Figure S1). In addition to the visualization, the exact sequences, annotations and more meta-information are provided for further in-depth analysis (e.g. co-occurrence with different antibiotic resistance genes or transposases).

3 Conclusion

GEnView provides automated, selective access to thousands of genomes and plasmids from the NCBI Assembly and RefSeq databases. Parallelized, automated annotation and phylogeny-based visualization of genome sequences allow for visual comparison of gene loci from several hundreds of genomes at the same time and greatly reduces the amount of manual work that previously was necessary to perform such tasks. The interactive visualization together with the generated metadata, alignments and phylogenetic trees allow for in-depth analysis of the gene locus of interest.

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Conflict of Interest: none declared.

Data availability

The FOX-1 reference is available at https://card.mcmaster.ca/ontology/38535; the 881 *Aeromonas* genomes are available at https://www.ncbi.nlm.nih.gov/assembly/?term=Aeromonas and easily downloadable using GEnView by following the GEnView tutorial at https://github.com/EbmeyerSt/GEnView/wiki.

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