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A Phylogenomic Gene Cluster Resource: The Phylogenetically Inferred Groups (PhIGs) Database

Paramvir Dehal, Wayne Huang and Jeffrey Boore

Abstract

We present here the PhiGs database, a phylogenomic resource for sequenced genomes, although many methods exist for clustering gene alimitis, very free watempt to create truly orthologous clusters sharing descent from a single ancestral gene across a range of evolutionary depths. Although these non-phylogenetic gene family clusters have been used broadly for gene annotation, errors are known to be introduced by the artificitual association of slowly eoving paralogs and lack of annotation for those more napidy evolving. A full phylogenetic framework is necessary for accurate inference of function and for many studies that address pattern and exchanism of the evolution of the genome. The automated generation of evolutionary gene clusters, creation of gene tess, determination of orthology and paralogy relationships, and the is an important resource to the socient for communic context

The PhIGs database currently contains 23 completely sequenced genomes of lungi and metazon, containing 406,653 genes that have been grouped into 42,645 gene clusters. Each gene cluster is built such that the gene sequence distances are consistent with the known organismal relationships and in so doing, maximizing the likelihood for the clusters to represent truly orthologous genes. The PhIGs website contains tools that allow the study of genes within their phylogenetic framework through keyword searches on annotations, such as GO and InterPro assignments and sequence similarity searches by BLAST and HMM. In addition to visualizing the gene clusters, the website also allow users to troves multi-species syntery maps.

Accurate analyses of genes and genomes can only be done within their full phylogenetic context. The PhiGs database and corresponding website URL: http://phigs.org addresses this problem for the scientific community. Our goal is to expand the content as more genomes are sequenced and use this framework to incorporate more analyses.

http://phigs.org

Common problems using pairwise BLAST for orthology determination

Unequal rates of change



Human Gene

Rat Gene 1

..... Human Gene 2

Rat Gene 2

Mouse Gene 2

Mouse Gene

Hidden paralogy

Gene deletions following duplication events may result in situations where paralogous genes may appear

Species Tree

Illustration of the clustering method

The tree shown on the left side of the figure indicates the evolutionary relationships among several hypothetical organisms, four from Clade A, how from Clade B, and one that is an outgroup. The right side of the figure illustrates a protein distance graph with circles representing proteins colored to conform to each organism, with the spatial distance of the circles proportional to their sequence distance. The duster is created by identifying a pair of sequences (a seed) that is the shortest distance from any Clade A protein to any Clade B protein. The cluster is then grown by adding all proteins that have a shorter distance than the seed until no additions can be made. The blue cloud represents no such cluster.

Protein Distance Graph



PhIGs

Phylogenetic Clustering of Eukaryotic Genomes



the settice evolutionary tes of the organisms beginning at the base, we ensure that the most cardy diverging gene families create the most comprehensive clusters, with later established families properly assigned to the lineages in which they arose. Genes with a highly accelerated amilion acid substitution rate, such that they are more distantly related to their sister genes than those sister genes are to a gene from the outgroup, are always excluded, since this cannot be differentiated from ancestral parallogy. The results of clustering 23 Opishtoken

By using an iterative approach, working through

genomes (Fungi - Metazoa) is shown in this figure. With 324.193 protein cooling genes defined by these genomes, 232,750 (-72%) are placed within one of the 42,645 resulting PhiOs gene cluster. The percentage of protein coding genes within a PhiOs cluster for each organism ranges from 92% to 44% and averages 70%.

Multiple Sequence Alignment construction

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Clustatify program. A summary graphic of the M&I represented on the web page for each cluster. This figure shows the portions of the sequence which align in blue and gaps in the alignment as dashes. This allows the user to quickly determine large scale alterations in alignments caused by intercina and deletions. Additionally, the exon showing exon boundaries as ventical lines and the size of each exon, in nucleotides, contend understand the exon. This is useful for examining gene structure and protein domain alterations.

Multiple sequence alignments are created using the

A more detailed examination of the MSA can be performed by either downloading the MSA file or by using the "jalview" java applet shown below.

Hidden Markov Models

The Multiple Sequence Alignments are used to create Hidden Markov Models to facilitate searching the clusters and to provide a resource for placing genes from genomes too sparsely sampled to be included in this comprehensive analysis.

Gene Structure and Domain Comparisons

The genomic location and exon structure of each gene is also provided, enabling analysis of such issues as whether the parallating analysis of such issues as whether the gene of the physically clustered within or whether the gene family is widely dispersed. Alterations in gene intron and exon structure (and sizes) relative to other members of the cluster may be the result of biological forces acting on the geneme or may simply be indicative of poor gene modeling. By examining the MSA, the user can determine whether poort) aligning or mission genes of a gene contains a protein domain which may indicate the gain or loss of some function.

Automated Creation of Phylogenetic Trees for each gene cluster



This is one output of the PhiCs analysis that is shown on the Cluster View webpage. Instead of simply listing the members of a cluster, a phylogenetic tree is created showing the evolutionary relationships of this multigene family. In this example, we can see that this family had gene deplection events at the base of verticatuss and in the fash optication events at the base of verticatus and in the family of amino and substitutions, we can see how rates of evolution have varied.

Orthology and Paralogy Determination

By reconciling the gene trees with the known evolutionary tree of the organisms, we can sort all genes into their orthologous and paralogous relationships.

Searching



Searches of the database can be done by sequence similarity or by text matches to annotation fields. Text searches can be done on gene names, defines or interPro annotations. Because these are associated with individual genes, the search function can be used to either return a list of genes from a selected set of taxa that contain the search term or it can return a set of clusters which contain genes matching the search term. Because all clustering is done at the protein level, sequence similarity searches can only be performed against protein database. An individual sequence can be aligned against the proteins contained in the database using the BLAST program. Matches to the sequence can there bus used as an entry in the totaken i which they belong. Alternatively, reMMG) generated from the MSA of the clusters using the IMMRP program. Noce a match has been made, the user can assay download either the rave fasts file of the cluster or the MSA file to create a tree incorporating the new sequence.

Whole Genome Synteny Maps



Genes ranging from number 205 through 301 on chicken chromosome 2 (numberd as they occur from the ptelomere to q-telomere along the chromosome) are shown as rectangles in the center of the diagram. An the left and right are the orthologs of these genes found in the human and mouse genomes as determined by the PhIOs analysis, shown as they are arranged. Black connecting lines join orthologs in the same relative transcriptional orientation whereas red lines indicate intervening genes without identified orthologs in the genomes being compared. Quan rectangles Indicate intervening genes without identified orthologs wist in chicken (the query genome), but not in the portion specified for this page.

Conclusion

The npidyl increasing number of sequenced genomes allows us to study genes and genomes within an evolutionary context. Not only does this assist in the transfer of anotations between genes, but also allows us to uncover how the forces of evolution have shaped each genome. The PhiCs database project seeks to facilitate comparative genomic, phylogenomic and functional genomic studes by providing a comprehensive resource for the determination of the evolutionary history for all genes from the fully sequenced genome projects. The two main properties that differentiate the PhiCs database from other clustering methods are the use of the known evolutionary history species to create gene clusters representing the descondants of a single ancestrating area and the creation of a complete phylogenetic gene tese functional annotation, genes structure, genomic position and other datasets, the PhiCs database will prove to be a valuable resource for all fields of biology currently using genomic data.

The scientific applications of the PhiSs database are broad, extending beynd practical genome annotation and analysis. For instance, ordivous applications are he use of orthogous gene ductuses for: (1) organismal physigenetic reconstruction; (2) the study of genome evolution by gene duplication; (3) gene structure evolution through the gain and loss of exons, introns, and domains; (4) the identification of gene family expansions and losses and (3) genome evolution. The PhiSs analyses have already been used to compare specifically the whole genomes of a lunicate, fish, mouse, and human, demonstrating that the relative positions in the human genome of paralogs generated by duplications at the base of verticates provide care verdence in two or of the cortentions hypothesis of the vorteod of undel genome duplication having occurred at the base of the vertebrates, and maybe providing the raw material for vertebrate complexity. Further applications can be develoced to mere duck of ther advicture community.

Future development includes improvements to the underlying clustering method, incorporation of more anotation data. Treation of more analysis tools and more rapid updates of newly available genomes. The functionality of the PhiCs database is currently accessible though the web interface and data files of orthology nelationships for download. Our goal is to convert this into an open source project to help maintain and expand this as a resource for the scientific community.