

Phydbac (phylogenomic display of bacterial genes): an interactive resource for the annotation of bacterial genomes

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ABSTRACT

Phydbac is a web interactive resource based on phylogenomic profiling, designed to help microbiologists to annotate bacterial proteins. Phylogenomic annotation is based on the assumption that functionally linked protein-coding genes must evolve in a coordinated manner. The detection of subsets of co-evolving genes within a given genome involves the computation of protein sequence conservation profiles across a spectrum of microbial species, followed by the identification of significant pairwise correlations between them. Many ongoing studies are devoted to the problem of computing the most biologically significant phylogenomic profiles and how best identifying clusters of 'functionally interacting' genes. Here we introduce a web tool, Phydbac, allowing the dynamic construction of phylogenomic profiles of protein sequences of interest and their interactive display. In addition, Phydbac can identify *Escherichia coli* proteins exhibiting the evolution pattern most similar to arbitrary query protein sequences, hence providing functional hints for open reading frames (ORFs) of hypothetical or unknown function. The phylogenomic profiles of all *E.coli* K-12 protein-coding genes are pre-computed, allowing queries about *E.coli* genes to be answered instantaneously. The profiles and phylogenomic neighborhoods are computed using an original method shown to perform better than previous ones. An extension of Phydbac, including precomputed profiles for all available bacterial genomes (including major pathogens) will soon be available. Phydbac can be accessed at: <http://igs-server.cnrs-mrs.fr/phydbac/>.

INTRODUCTION

Determining protein functions from genomic sequences is one of the main challenges of bioinformatics. To this purpose, alignment methods based on sequence similarity, such as PSI-BLAST (1) or Pfam (2), are the most heavily used and are still being refined. Yet, they are only capable of providing reliable functional predictions for ~50% of the open reading frames (ORFs) of most newly sequenced microorganisms (3), corresponding to the proportion of already functionally annotated protein coding genes. Besides the experimental determination of gene functions, escaping this vicious circle requires the development of bioinformatic approaches going beyond the recognition of sequence similarity and functional signatures. Phylogenomic profiling is one of these new methods. It is based on the assumption that proteins involved in a common metabolic pathway or constituting a multi-molecular complex are likely to evolve in a correlated manner. We use the term co-evolution throughout this paper to designate such a behavior. This paradigm, originally named phylogenetic profiling, was first put to use by Pellegrini *et al.* (4) who demonstrated that some information on the function of a protein could be retrieved by analyzing the functions of its phylogenomic neighbors, this neighborhood being defined as the subset of the best co-evolving genes in the same genome (e.g. *Escherichia coli*). This approach has been subsequently used in many studies (5–7) and the definition of a meaningful neighborhood refined in various ways (8–10). Other phylogenomic methods have been proposed, such as the analysis of gene co-localization (11,12), as well as the systematic search for gene fusion events (13,14). At the moment, Phydbac only uses the initial concept of co-evolution, but the co-localization information will be added in the future.

In the current version, Phydbac emulates two main different modes of operation. In the first mode, the software allows researchers to build, display and compare the evolution profile(s) of their protein(s) of interest, for instance to see if they exhibit any evidence of co-evolution. The phylogenomic profile is computed on line, using an ORF database derived from 71 bacterial and archaeal (non-redundant) species. By analyzing each sequence conservation profile individually and/or

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and phylogenomic neighbors. *E.coli* genes can be retrieved by their names, the presence of a keyword in their annotation or by similarity with a user-provided query sequence. Upon the selection of one or several genes, their profiles are displayed (Fig. 1). Clicking on the icon near the name of the genes identifies its 10 closest phylogenomic neighbors and gives access to their conservation profiles. As the number of neighbors is arbitrary, the possibility of getting more or less than 10 neighbors is offered (using a plus or minus button near the query's name).

In addition to the profiles, accessory information is also displayed for any gene list (manually selected genes or a query and its neighbors). When two or more genes of a list belong to the same pathway, to the same operon, have a significant sequence similarity or are found to be co-localized, it is indicated in the four corresponding columns, right before their profiles. For instance, Figure 1 shows that *caiA* and *caiD* are involved in common pathways (carnitine degradation and carnitine metabolism—CoA-linked) and that they belong to the same operon. The 'Paralogs' column shows that *fixB*, *ydiR* and *ycgQ* share some significant sequence similarity. Finally, the 'Colocalization' column indicates that *paaH* and *ydiC* are co-localized with *caiA* (i.e. their respective homologous sequences are separated by <2000 nucleotides in more than three genomes). As co-localization is not a transitive property (*paaH* and *ydiC* are not co-localized), each gene found co-localized with other genes in the list generates its own column of colored markers, the darkest hue being associated with each reference gene. For selected genes (checkboxes are on the left of the gene names), annotations and an unrooted tree, built as described before, can be displayed in pop-up windows. Finally, expression intensity values (e.g. from DNA-chip experiments) can be loaded in parallel and displayed next to the profiles of the selected genes (or any of its neighbors). The visual comparison of phylogenomic versus expression intensity profiles may help in the generation of hypotheses on putative functions and metabolic processes.

FUTURE PROSPECTS

Future versions of Phydbac will extend the mode of operation currently limited to *E.coli* ORFs to all fully sequenced micro-organisms. Priority will be given to major human pathogens. This will require the integration/incorporation of the functional annotations contained in databases on different bacterial genomes and the storage of the profiles of all bacterial ORFs.

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