PHOREST: a web-based tool for comparative analyses of expressed sequence tag data

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**PROGRAM NOTE**

**PHOREST: a web-based tool for comparative analyses of expressed sequence tag data**

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**Abstract**

Comparative analysis of expressed sequence tags is becoming an important tool in molecular ecology for comparing gene expression in organisms grown in certain environments. Additionally, expressed sequence tag database information can be used for the construction of DNA microarrays and for the detection of single nucleotide polymorphisms. For such applications, we present **PHOREST**, a web-based tool for managing, analysing and comparing various collections of expressed sequence tags. It is written in PHP (PHP: Hypertext Preprocessor) and runs on UNIX, Microsoft Windows and Macintosh (Mac OS X) platforms.

**Keywords:** comparative analysis, expressed sequence tags, software

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Expressed sequence tag (EST) analyses are based on single-pass, large-scale DNA sequencing of reverse-transcribed messenger RNA (cDNAs) and have become a widely used approach for transcriptome analyses within functional genomics. Genes expressed under given growth conditions or at various developmental stages are characterized by EST sequencing and, due to the redundancies of transcripts, the expression levels can be inferred (i.e. transcript profile) (Qutob et al. 2000; Davey et al. 2001). There are currently more than 600 different species with ESTs publicly available through dbEST (Boguski et al. 1993). Within molecular ecology, EST database information can be used for the construction of DNA microarrays and for the detection of single nucleotide polymorphisms (Picoult-Newberg et al. 1999; Gibson 2002; Oleksiak et al. 2002).

In this study we present **PHOREST**, which is a novel web-based tool for comparative studies across multiple EST projects (EST libraries). The tool is designed to facilitate the storage, handling and mining of EST sequence data. Projects can easily be shared between multiple users via an intranet or the Internet and can be made public with **PHOREST** features by adding a guest user. **PHOREST** differs from other software packages specializing in EST data analysis, such as **RED** (Everitt et al. 2002), **XGI** (Waugh et al. 2000), **STACKPACK** (Miller et al. 1999) and **PIPEONLINE** (Ayoubi et al. 2002), in that the comparative approach is well developed and that management is possible through a web interface, requiring a minimum of computational skills. **PHOREST** can handle many EST projects simultaneously and compare the redundancy levels (i.e. the frequencies of a given transcript in the different EST projects). **PHOREST** also greatly facilitates the exchange and collaboration possibilities between researchers by using the web-based approach. By having restrictions on various viewing and editing options the security of the data is ensured. The system also allows for several different databases running on the same **PHOREST** installation. The users will only see the databases to which they have access. This makes it possible to have **PHOREST** as a service available to many research groups from a central server. The data analysed and maintained through **PHOREST** can be used for various research areas, for example to generate a uniset for construction of DNA microarrays. In addition, the results can be exported in a number of formats to facilitate external analysis of the **PHOREST** data. For example, it is possible to export the annotations from **PHOREST** into a pathological format (Karp et al. 2002) which can then be directly used for metabolic pathway reconstruction.

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PHOREST main view

To compare various collections of EST sequences, PHOREST can handle information from more than one EST database and each database can contain information from several EST projects (corresponding to one biological sample, cDNA library). After logging into PHOREST, the databases and EST projects to which the user has access are displayed. Each database has a statistics page calculated by PHOREST which provides information on the number of clones uploaded into each EST project, the number of contigs after assembly, the number and relative fraction of singletons, orphans and low-, medium- and high-score clones after assembly and BLAST similarity search. The page is of use both for EST projects where sequencing is not yet finished (validation of quality) as well as computing summary statistics once the EST sequencing is finished.

The main page of the PHOREST web interface shows a list of EST clones from one or several of the EST projects in the selected database (Fig. 1, no. 1). The top frame contains a search option that can be used for filtering and displaying subsets of the EST clones or contigs (Fig. 1, no. 3). For example, it is possible to search and then display ESTs with specific clone names, sizes (sequenced base pairs), specific score values (from BLAST searches), GenBank descriptions, cluster sizes and redundancy levels. The search profile can be reviewed and modified using the ‘Advanced search’ option. The results can either be listed as individual clones (‘Listed by clones’) or as contigs (‘One clone per cluster’) (Fig. 1, no. 7). The contig list provides a uniset of unique transcripts within the selected project(s). For the analysis of a subset of the ESTs/contigs there is a function for selection that can be activated by using the flag hyperlink (Fig. 1, no. 6). Once a subset of clones/contigs has been flagged it can be further analysed, exported in FASTA format or as a tab-delimited table suitable for Microsoft Excel and other programs. The table includes information on the transcript profiles in the various projects, BLAST scores, the GenBank description lines and Accession nos as well as results from functional annotations. In addition, users of PHOREST can...
perform BLAST (Altschul et al. 1990) searches against any locally stored sequence database (both public such as nr. from GenBank and custom databases) (Fig. 1, no. 2). The PHOREST software will detect whether it is a protein or nucleotide database and run the appropriate BLAST program.

In the lower frame, either all EST clones (‘Listed by clones’) or only one clone per cluster (contig) (‘One clone per cluster’) can be listed (Fig. 1, no. 7) as an html table. The columns of the table contain information on the clone name and the number of sequenced base pairs (in parentheses) (Fig. 1, no. 8), descriptions from homology searches (Fig. 1, no. 9) and the similarity score and the expected frequency value of the top hit from the BLAST search (Fig. 1, no. 10) (Altschul et al. 1990). The EST clones can be sorted based on the information in any of these columns. The EST sequences are automatically reblasted as new updates of the GenBank nonredundant (nr) database (10) are released. In addition, clones can be manually selected for updates (Fig. 1, no. 6). New top hits are indicated by red text (‘New’) and sequences with new top hits can be listed separately by asking for ‘Only new’ sequences in the search function available in the top frame (Fig. 1, no. 3).

Clustering and annotation

Results from the clustering analyses using the contig assembly program (CAP) (Huang 1992) are shown as the number of clones in the contig (Fig. 1, no. 11). The column with the clustering data also provides a link to a page showing the alignment of the sequences in the cluster (Fig. 1, no. 11).

The ESTs are annotated according to the four categories used by MIPS (Mewes et al. 1997): protein classes, functional classes, protein complex classes and EC numbers. A fifth category (origin) is used to describe the type of sample (tissue, developmental stage or species) (Fig. 1, no. 12). Linked to this column is a page that gives a list of BLAST hits for the six possible open reading frames of the EST clone. The description column provides links to the GenBank nr database, the ExPaSy ENZYME database, LocusLink and the MIPS yeast genome database (Burks et al. 1985; Mewes et al. 1997; Bairoch 2000; Wheeler et al. 2000). Links are automatically displayed when the description field in GenBank contains an EC number or when the EST sequence displays similarity to sequences in Saccharomyces cerevisiae or LocusLink organisms. Simultaneous examination of ESTs assembled into a given contig can reduce the time needed for annotation. In addition, the annotation process can be speeded up by allowing multiple users to have full access to the EST projects.

Graphical display

In PHOREST, all or a subset of ESTs from one or more projects can be displayed using various graphical interfaces. The global distribution of contigs among projects can be viewed in a triangle diagram. Based on normalized redundancies, assembled contigs are distributed along the axes of a triangle. Contigs at the corners of the triangle represent transcripts unique to one of the EST projects, whereas contigs found at the centre are equally redundant in all projects. Clicking anywhere within the triangle brings up a table of all contigs with corresponding distributions and these can be used for further analysis. The distribution of ESTs or contigs in different functional categories can be presented in tables, pie charts or bar diagrams. The functional distribution of ESTs and contigs in different projects is presented to provide a comparative overview of multiple EST projects.

Administration of PHOREST

In order to ensure the security of the information in the PHOREST database a number of security measures is in place. One person has the administration rights of the system and will have total control. The PHOREST administrator can add new users as well as change users’ access rights to the different databases so that a user can only see the databases that he/she has access rights to. In addition, the PHOREST administrator can also add new databases and EST projects by simply filling in a form on the administration page. The input file required is a FASTA file for each EST project. Users with full access rights to a specific database can upload new sequences from the administration page.

System requirements and availability

PHOREST only requires freely available programs. These include the MySQL relational database (www.mysql.com) and the web server Apache (www.apache.org). To cluster the EST sequences, CAP (Huang 1992) is used and the homology searches are performed using BLAST standalone (Burks et al. 1985). PHOREST is written in PHP and runs unchanged on UNIX, Microsoft Windows and Macintosh (Mac OS X) platforms. As most administrative tasks are handled through the administration page in PHOREST, no advanced computer knowledge is required (i.e. no UNIX or programming skills).

PHOREST is freely available for academic use from the corresponding author. A manual and PHOREST demonstration database are available on the web (http://www.biol.lu.se/phorest/).

Implementation

We have used PHOREST to analyse EST data generated from parasitic and symbiotic fungi (Tunlid & Ahrén 2001; Johansson et al. 2003). Both studies involve comparative analyses of ESTs from a number of different growth stages.

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More than 22,000 EST sequences in five databases containing 13 separate EST projects are currently being handled by phorest on a Linux computer (2 x 1.8 GHz, 1 GB RAM). Two other research groups are currently testing the phorest software for their EST projects. The fact that several groups have actively used phorest in research for several years has ensured that phorest is now a stable software tool with a minimum of bugs.

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References