Sequence analysis

BADASP: predicting functional specificity in protein families using ancestral sequences
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ABSTRACT
Summary: Burst After Duplication with Ancestral Sequence Predictions (BADASP) is a software package for identifying sites that may confer subfamily-specific biological functions in protein families following functional divergence of duplicated proteins. A given protein phylogeny is grouped into subfamilies based on orthology/paralogy relationships and/or user definitions. Ancestral sequences are then predicted from the sequence alignment and the functional specificity is calculated using variants of the Burst After Duplication method, which tests for radical amino acid substitutions following gene duplications that are subsequently conserved. Statistics are output along with subfamily groupings and ancestral sequences for an easy analysis with other packages.

Availability: BADASP is freely available from http://www.bioinformatics.rcsi.ie/~redwards/badasp/

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Supplementary information: A manual with further details can be downloaded from http://www.bioinformatics.rcsi.ie/~redwards/badasp/

INTRODUCTION
The divergence of proteins following gene duplication has long been recognized as an important process in the evolution of new or specific protein functions. Functional divergence is proposed to occur through some combination of neofunctionalization—the evolution of novel gene function—and subfunctionalization—the partitioning of two or more existing gene functions between paralogues (genes related by duplication) (Zhang, 2003). Although no consensus has yet been reached as to which process is more important, the distinction is somewhat irrelevant for the bioinformatic prediction of sites important for differences in gene function between paralogues (though it is vitally important for the interpretation of results). Instead, it is more pertinent to consider the types of substitution that occur at these sites and the phylogenetic signal that they leave.

Sites of functional change following duplication can be broadly classified into two categories, which Gu has named Type I and Type II (Gu, 2001). Type I functional divergence shows a change in selective constraint on a site following duplication, either by relaxation of existing purifying selection or by gaining functional importance at a previously unimportant site. In contrast, sites experiencing Type II divergence remain important in both duplicates but a different amino acid is favored in each duplicate. Both Type I and Type II divergence can occur as the result of either neofunctionalization or subfunctionalization. For example, subfunctionalization may occur by partitioning domain functions, with different domains maintained in different paralogues (Type I divergence), or by each paralogue specializing for a given set of existing substrates (Type II divergence). Similarly, new gene function may arise at previously unimportant sites (Type I) or by recruiting existing functional sites to the new function (Type II), while the paralogue fulfills the previous roll of the ancestral protein.

Several methods now exist to detect either Type I or Type II divergence (Lichtarge et al., 1996; Caffrey et al., 2000; Hannenhalli and Russell, 2000; Johnson and Church, 2000; Gu, 2001; del Sol Mesa et al., 2003; Kalinina et al., 2004a; Abhiman and Sonnhammer, 2005a). Many of these, however, are complex methods that lack simple software implementations and/or rely on additional information, such as structural data, which is not always available. Although there are available tools for state of the art predictions for divergence of either Type I (Gu and Vander Velden, 2002) or Type II (Kalinina et al., 2004b) for single protein families, there is still the need for a simple analysis package that can be run from the command line for multiple families and can potentially detect both Type I and Type II divergence. BADASP provides software to implement the previously published Burst After Duplication (BAD) algorithm (Caffrey et al., 2000), plus two variants for identifying Type I and Type II divergence that have been used successfully in identifying functionally interesting sites in platelet signaling proteins (unpublished data).

ALGORITHM
BADASP implements three versions of the BAD algorithm (Caffrey et al., 2000). All three versions are built on the same underlying assumption that sites critical to changes in gene function between paralogues are marked by a burst of radical amino acid substitutions directly after duplication, which are subsequently conserved within orthologous groups. This is calculated by comparing the changes in physiochemical properties along the relevant branches for each site:

\[
\text{BAD} = \text{RC} - \text{AC},
\]

where AC is the ‘Ancestral Conservation’ score, calculated as the change in physiochemical properties between the duplication node and the ancestral node for the subfamily; RC is the ‘Recent Conservation’, calculated as the mean change in properties between the ancestor of the subfamily and each orthologous (leaf) sequence.
(1) BADT explicitly analyses two subfamilies, related by a single duplication event, for Type II divergence and is simply the sum of the BAD scores (1) for the two subfamilies.

(2) BADX looks for Type I neofunctionalization in a specific Query subfamily by comparing it with to its duplicate only:

\[ \text{BADX} = \text{RC} - \text{ACX}, \]  

(2)

where ACX is a modified AC score, calculated as the change in physicochemical properties between the two post-duplication nodes; RC is for the Query subfamily only. This method is more robust to incorrect ancestral sequence assignment at the duplication node.

(3) BADN looks for Type II divergence across multiple subfamilies:

\[ \text{BADN} = \frac{\text{BADS}}{N-1}, \]  

(3)

where BADS is the sum of the BAD scores (1) for each subfamily; N is the number of subfamilies. AC values (1) are calculated using the ancestor of each subfamily and the root of the tree, which should be the most ancient gene duplication.

As with other methods, all three BAD algorithms are obviously sensitive to alignment and tree quality. In addition, incorrect ancestral sequence prediction will give erroneous results.

IMPLEMENTATION

BADASP has been implemented using a set of open source Python modules. By default, the amino acid property matrix of Livingstone and Barton (1993) is used. Ancestral sequences are calculated using GASP (Edwards and Shields, 2004). This algorithm was specifically designed with BAD in mind and will reconstruct ancestral sequences for gapped columns of the alignment, allowing the use of partial sequences and/or BADN calculations for sites that are gaps in one or more subfamilies. In addition to the GASP ancestral sequences and BAD statistics, BADASP will also calculate a number of additional specificity and sequence conservation statistics to assist the interpretation of results.

Main BADASP output falls into three primary categories: (1) statistics for a given residue; (2) statistics for a given window size across (a) the whole alignment, (b) the Query protein of interest (if given) and (c) the predicted ancestral sequence of each subfamily; and (3) predicted ancestral sequences at (a) the root and (b) the ancestor of each subfamily. This output is in a tab- or comma-delimited file for easy manipulation or viewing with other programs. A batch mode with the option to output results from multiple data-sets as a MySQL database is planned. In addition to this flat file, the standard ancestral sequence output of GASP (Edwards and Shields, 2004) and a file containing subfamily grouping data is also produced.

A manual with full details of algorithms, acceptable input formats, sequence statistics, output files and parameters can be found at http://www.bioinformatics.rcsi.ie/~redwards/badasp/

DISCUSSION

Sophisticated methods are now available for predicting sites of functional divergence. Abhiman and Sonnhammer have recently performed a large-scale analysis of FunShift (Abhiman and Sonnhammer, 2005a) to test its ability to discriminate between functionally diverged and functionally conserved enzymatic activities in related subfamilies (Abhiman and Sonnhammer, 2005b). However, there is currently no good dataset of individual residues responsible for functional specificity, with which different methods can be compared. Complexity is not always beneficial, and what one gains in the swings of sensitivity, one can lose in the roundabouts of interpretation. A place still exists, therefore, for an open source implementation of a simple algorithm, the results of which may be easier to understand and analyze. BADASP provides such an implementation that is suitable for use in a high-throughput automated analysis of many families. Alternatively, BADASP could provide useful supplemental data for a more focused analysis on a given family when used in parallel with a more complex method, such as DIVERGE (Gu and Vander Velden, 2002) for Type I or SDPpred (Kalinya et al., 2004b) for Type II divergence, or Rate Shift analysis (Abhiman and Sonnhammer, 2005a). The open source Python implementation allows extra measures of specificity or conservation to be added with relative ease.

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REFERENCES


