SitEx: a computer system for analysis of projections of protein functional sites on eukaryotic genes

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ABSTRACT

Search of interrelationships between the structural–functional protein organization and exon structure of encoding gene provides insights into issues concerned with the function, origin and evolution of genes and proteins. The functions of proteins and their domains are defined mostly by functional sites. The relation of the exon–intron structure of the gene to the protein functional sites has been little studied. Development of resources containing data on projections of protein functional sites on eukaryotic genes is needed. We have developed SitEx, a database that contains information on functional site amino acid positions in the exon structure of encoding gene. SitEx is integrated with the BLAST and 3DExonScan programs. BLAST is used for searching sequence similarity between the query protein and polypeptides encoded by single exons stored in SitEx. The 3DExonScan program is used for searching for structural similarity of the given protein with these polypeptides using superimpositions. The developed computer system allows users to analyze the coding features of functional sites by taking into account the exon structure of the gene, to detect the exons involved in shuffling in protein evolution, also to design protein-engineering experiments. SitEx is accessible at http://www-bionet.sscc.ru/sitex/. Currently, it contains information about 9994 functional sites presented in 2021 proteins described in proteomes of 17 organisms.

INTRODUCTION

The functional sites of proteins are the major determinants of the biological function of proteins, being the most conserved portions of protein sequence. The features of structural organization of the protein functional sites at the level of gene exon structure may be consequential for evolutionary events such as formation of protein domain structure and those due to exon shuffling (1), the duplication of the domain and insertion of domain from one gene to another. A protein domain is often encoded by a single or several exons (2). The formation of mosaic proteins composed of several identical and non-identical domains has been demonstrated. Shuffling events of this kind might have significantly affected protein function and they have been considered as major mechanisms underlying forces operating during evolution and shaping the diversity of domain complexity (1).

Databases and program resources for projection of the domains and exon boundaries on amino acids sequences and spatial protein structures are currently available. Resources for projection of domain boundaries are both numerous and robust. These include the well-known databases such as Pfam (3), InterPro (4), TIGRFAMs (5), PROSITE (6), ProDom (7). Certain resources that describe the projections of exon and domain boundaries on primary and tertiary structure of proteins have been developed. Thus, web-resource XdomView (8) contains information on projection of exon and domain boundaries on 3D structure; SEDB (9) maps the exon boundaries and intron phases on multiple structural superimposition of 3D structures; ExDom (10) maps the location of exon boundaries and protein domains on sequence and spatial structure of proteins. However, information about functional sites is not contained in these resources.

A number of resources store data on the functional sites: Catalytic Site Atlas (11), scPDB (12), SitesBase (13), InterPro, PROSITE. They are constructed automatically or under expert control. Based on multiple alignment Gene3D could predict the active sites (14). Earlier we have developed the PDBSite database (15) that contains data on sites automatically extracted from PDB. It was integrated with the PDBSiteScan...
program (16) for recognition of functional sites in protein spatial structure. However, these resources do not contain information on the boundaries of exons and domains.

The aim of this work was to develop a resource that describes projection of protein domains, functional sites and exon boundaries on protein and coding gene sequence.

In analysis of the structural–functional organization of proteins, search of sequence and spatial structure similarity among polypeptides encoded by individual exons appears promising. SitEx was integrated with the BLAST and 3DExonScan programs. BLAST is used to search similarity of a query protein to polypeptides encoded by individual exons stored in SitEx. 3DExonScan searches the similarity between 3D structures of query protein and polypeptides encoded by single exons. It could be helpful for search for the structural similarity between the polypeptides that do not necessarily possess the sequence similarity (17,18).

CONSTRUCTION AND CONTENT

The SitEx database provides data on mapped positions of exon and domain boundaries, also on functional sites in full-length protein and coding gene sequences. Here we examine only proteins with known 3D structure and corresponding genes. Figure 1 presents the SitEx structure and the main information tables in relational database constructed with MySQL 5.1. They are as follows. Chain gives information on the coding sequences and the proteins they encode, also on the identifiers from the Ensembl (19) database; Site contains information on protein functional sites, names, references to PDB (20), functional site description and some other information; ExonPos and SitePos provide information on positions of exon boundaries and functional site amino acids/codons in coding gene sequence and the corresponding protein. The other tables are designed for storing references to external databases, also for links between the tables.

Those protein structure, whose descriptions contained functional sites were selected from the PDB database. Their sequences and their encoding gene sequences, exon and domain borders (according to the Pfam classification) were extracted from the Ensembl database. To identify the amino acid residues in a functional site, the same approach was used as for PDBSite. Information on amino acid positions of functional site in the PDB sequence was retrieved from the SITE record. Information on the ligands, sequences, organisms, protein names and site identifiers was retrieved from PDB, too. The numberings of amino acid residues in PDB and Ensembl may differ. With this mind, we analyzed the sequence equivalence. Using the PDB identifier and ClustalW (21) alignment, the information on coding gene was retrieved from Ensembl (version of July 2009). Also, using ClustalW alignments, amino acid positions of functional site, exon, Pfam domains and SCOP structural region boundaries were identified in polypeptide chain. Information on folds was retrieved from the SCOP database (22). The programs retrieving the information from the external data and constructing SitEx database were developed in Perl. As a result of data integration, we retrieved 2021 non-redundant protein sequences that belong to 17 genomes of different organisms.

Search in SitEx using BLAST

Based on the data on exon boundaries in gene sequence stored in SitEx, two BLAST (23) databases were developed for sequences, exons and the polypeptides they encode. As a result, search of sequences involved in coding or in the formation of a protein functional site became available.

The 3DExonScan program

The 3DExonScan program aims for search of structural similarity with polypeptides that are encoded partly or entirely by a single exon. 3DExonScan uses the files in PDB format that are prepared for every such polypeptide, if the tertiary structure even for the part of it is known.

3DExonScan is based on the SSM and CE algorithms (24,25). The 3D protein structure is represented as elements of the secondary structure, an α-helix or a β-strand. This makes feasible quick comparisons of protein 3D structures with those in the SitEx database. This representation sets limitations on the length of polypeptide sequences.

3DExonScan ignores exons encoding polypeptides whose secondary structure consists of less than two elements. Many exons code for short protein regions that may possibly be only single α-helix, β-strand or loop. 3D structures of single α-helaces or single β-strands from different proteins are very similar and structural superimposition will result, in most cases, in small RMSD. To avoid this, short exons that encode single helices or β-strands were excluded from analysis. Being aware of the limitations of the approach, we intend to use a strategy we developed earlier to speed up structural superimposition of functional sites in the PDBSiteScan.

Using the 3DExonScan program, the user obtains the structural homologues from database of 3D structures of polypeptides that are encoded by single exon. The results are presented as a table that contains reference to exon description in the SitEx, PDB identifier, protein description and the Z-score of structural superimposition. The threshold for the Z-score is set empirically at 3.2 (this corresponds to $P = 0.9993$). Superimpositions that have lower Z-score are eliminated by program. Having chosen the most suitable superimposition, the user gets a visualization of superimposed 3D structures of query protein and polypeptide with Jmol 3D structure viewer (26)(Figure 2). It is possible to download the structural superimposition in a PDB format.

SitEx content

A statistics page is provided. The database includes:

- 17 organisms (list of organisms is presented in Table 1): 75% of protein functional sites are...
represented by human proteins, 10% mouse, 5% rat, 5% bovine, 2% nematode, the remaining proteins occur singly;  
- 715 ligands;  
- 2021 non-redundant Ensembl protein sequences;  
- 9994 (of 10 887) unique functional sites (sites that are different in amino acid composition and positions in similar PDB chains within one PDB entry);  
- proteins were assigned to 13 groups according to their type; the enzyme group is widely represented (Table 2); and  
- the functional sites are assigned to 14 groups according to the ligand type they bind (Table 3).

**UTILITY**

**Interface**

The web service SitEx consists of a database covering the protein functional sites, Pfam and SCOP domain projections, and provides search of the homologous exons and regions of the polypeptide chain encoded by a single exon (Exon BLAST Search page), and also search of the structurally similar polypeptides encoded by a single exon (3D Exon Search page).

One can access the information stored in SitEx through a user-friendly web-interface. The description page of the exon contains information on the exon identifiers, exon length, its boundaries, details concerning the coding protein according to the PDB and Ensembl data. An exon is in bold uppercase in both a coding sequence and an amino acid sequence. The functional sites are represented with uniquely colour background. Color symbols highlight sequence regions that belong to a single domain or structure (Figure 3A).

The description page of a functional site contains information on: (i) the protein that contains known functional site, the different chains on which a functional site is located according to the PDB data; (ii) the site amino acid positions in protein and in coding sequence; (iii) the discontinuity coefficients of a site on amino acid sequence and through exons; (iv) exon borders. A functional site is in bold uppercase in both a coding and an amino acid
sequences. The exons are represented with uniquely coloured background. Color symbols highlight sequence regions that belong to a domain or a structure (Figure 3B).

There is the page for the current statistics for SitEx. Search queries are performed through the PDB, SCOP and Ensembl identifiers, names of the Pfam domains, organism, ligand, length and ordering of exon or any keywords (Figure 3C). Search of the exon length is based both on the number of nucleotides and that of amino acids encoded by the exon. A list of exons (and the polypeptides they code) together with the identification of encoded sites (Figure 3D) results from the query.
CONCLUSIONS

The SitEx system stores information on projection of exon and domain boundaries, positions of functional sites on protein sequence and the coding sequence of the gene. This system provides opportunities for analysis of the relationships between the exon–intron structure of genes and functional sites of proteins. BLAST allows the search for the coding sequences and polypeptides encoding by them corresponding to single exon. 3DExonScan supplies the search through database of 3D structures of polypeptides encoded by single exon with structural superimposition. Using these programs, it is possible to investigate the sequence and structural conservation of the single exon coded polypeptides involved in the formation of a protein functional site.

SitEx is applicable in the study of the structural-functional organization of the gene and the features of coding and evolution of the functional sites by taking into account the exon structure of the gene; detection of exons involved in shuffling in an evolutionary perspective; it is helpful in rational design of novel proteins composed of fragments encoding individual exons from distinct genes.

In the nearest future we are going to expand the number of genomes represented in SitEx by fungi and plant Ensembl databases. During the next step, we intend to associate the functional site positions in gene encoding proteins with known 3D structure and exon boundaries of orthologous genes.

Table 3. Ligand classification in SitEx

| Ligand class                             | Number of entries |
|-----------------------------------------|-------------------|
| Metal ions                              | 2944              |
| Acid anions                             | 2434              |
| Organic acids                           | 551               |
| Nucleotide phosphates                   | 799               |
| Sugars                                  | 308               |
| Proteins                                | 73                |
| Amino acids and their compounds          | 164               |
| Coenzymes                               | 89                |
| Alcohol and its compounds               | 665               |
| Atoms and inorganic molecules            | 351               |
| Amines and amides                       | 1112              |
| Porphyrins                              | 59                |
| Other ligands (alkaloids, ketones, pigments, etc.) | 1354              |
| Unknown ligands                         | 396               |

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Figure 3. Visualization of internal pages from SitEx. (A) exon description page. The start and end positions of the exon are shown. They correspond to the Ensembl numeration. AA, amino acid; bp, base pair; (B) a fragment of a page for description of functional site; (C) a query page; (D) result of query using the keyword ‘ATP’ in the ligand line. The functional site, information on the sequence, list of exons, which encode the sequence, their length measured in codons is displayed.
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