

Annotation Transfer for Genomics: Measuring Functional Divergence in Multi-domain Proteins

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For *Genome Research* (Version gr-final)

ABSTRACT

Annotation transfer is a principal process in genome annotation. It involves "transferring" structural and functional annotation to uncharacterized open reading frames (ORFs) in a newly completed genome from experimentally characterized proteins similar in sequence. To prevent errors in genome annotation, it is important that this process be robust and statistically well characterized, especially with regard to how it depends on the degree of sequence similarity. We and others have previously analyzed annotation transfer in single-domain proteins. Multi-domain proteins, which make up the bulk of the ORFs in eukaryotic genomes, present more complex issues in functional conservation. Here we present a large-scale survey of annotation transfer in these proteins, using scop superfamilies to define domain folds and a thesaurus based on Swissprot keywords to define functional categories. Our survey reveals that multi-domain proteins have significantly less functional conservation than single-domain ones, except when they share the exact same combination of domain folds. In particular, we find that for multi-domain proteins approximate function can be accurately transferred with only 35% certainty for pairs of proteins sharing one structural superfamily. In contrast, this value is 67% for pairs of single-domain proteins sharing the same structural superfamily. On the other hand, if two multi-domain proteins contain the same combination of two structural superfamilies the probability of their sharing the same function increases to 80%. (And in the case of complete coverage along the full length of both proteins, this value further increases to more than 90%.) Moreover, we found that only 70 of the current total of 455 structural superfamilies are found in both single and multi-domain proteins, and only 14 of these were associated with the same function in both categories of proteins. We also investigated the degree to which function could be transferred between pairs of multi-domain proteins with respect to the degree of sequence similarity between them, finding that that functional divergence at a given amount of sequence similarity is always about two-fold greater for pairs of multi-domain proteins (sharing similarity over a single domain) in comparison to pairs of single-domain ones -- though the overall shape of the relationship is quite similar. Further information is available at <http://partslist.org/func> or <http://bioinfo.mbb.yale.edu/partslist/func> .

Introduction

Annotation Transfer

The ultimate goal of the genome projects is to determine the structure and function of all the newly identified gene products. Fundamentally, this will be carried out via annotation transfer, transferring the structural and functional annotation from an experimentally characterized protein (say in a model organism such as *E. coli*) to a predicted protein in a newly sequenced genome that shares similarity in sequence. The degree of annotation transferred will depend on the degree of sequence similarity. This process is shown schematically in Figure 1. In this paper we aim to address this major question in

bioinformatics, specifically focusing on multi-domain proteins, as they make up the bulk of the proteome in eukaryotic organisms (Gerstein, 1998).

Our work is a direct outgrowth of two previous analyses of ours that concentrated on single-domain proteins. In an earlier paper we found that the different structural classes of the scop classification system have different propensities to carry out certain types of function (Hegyí & Gerstein, 1999). In particular, while the alpha/beta folds were disproportionately associated with enzymes and all-alpha and small folds with non-enzymes, the alpha+beta structures had an equal tendency for both enzymatic and non-enzymatic functions. Wilson et al. (2000) compared a large number of protein domains to one another in a pair-wise fashion with respect to similarities in sequence, structures and functions. Using a hybrid functional classification scheme merging the ENZYME and FLYBASE systems (Bairoch, 2000; Gelbart *et al.*, 1997), they found that precise function is not conserved below 30-40% identity, although the broad functional class is usually preserved for sequence identities as low as 20-25%, given that the sequences have the same fold. Their survey also reinforced the previously established general exponential relationship between structural and sequence similarity (Chothia & Lesk, 1986).

Other work on establishing relationships between sequence, structure and function

Several other groups have studied the relationship between sequence, structure and function in detail, attempting to determine the extent to which functional transference between matching proteins is feasible (Martin *et al.*, 1998; Shah & Hunger, 1997; Shapiro & Harris, 2000; Thornton *et al.*, 1999; Thornton *et al.*, 2000; Zhang *et al.*, 1999; Todd et al. 2001). Orengo et al. (1999) analyzed protein families in the CATH database and concluded that more than 96% of the folds in the PDB are associated with a single homologous family. By investigating enzymatic folds they also found that more than 95% of homologous families show either single or closely related functions. Pawlowski et al. (2000) studied the relationship between sequence and functional similarity in the twilight zone of 10-15% sequence similarity and found a clear correlation between the two, with functional similarity based on the E.C. classification of enzymes.

Russell et al. (1997) analyzed binding sites in proteins with similar 3D structures and estimated that 90% of new remote homologues have common binding sites and similar functions. Eisenstein et al. (2000) evaluated the first results from the structural genomics projects and found that in many instances the protein structure itself offers an important clue to its biological function. Stawiski *et al.* (2000) found that for just the proteases function could be rather successfully predicted. Devos & Valencia (2000) presented a critical view of function transference between similar sequences, highlighting the limitations of this process due to errors in databases and the inherent complexity of the relationship between protein sequence-structure and function that does not allow "simplistic interpretations." They also found that binding sites are the least conserved features between related proteins while the catalytic activity of enzymes is the most conserved one.

Multi-domain proteins with divergent functions: how common?

Most of these previous investigations focused on single-domain proteins or did not distinguish between single- and multi-domain ones. It is not clear how the multi-domain proteins with various functions behave with respect to functional conservation: whether they are more or less conserved than their single-domain counterparts. In particular, as shown in figure 1, if one multi-domain protein shares a single domain fold with another one it is not clear the degree to which the functional conservation of these proteins is constrained by the shared part and to what degree it is influenced by other domains that are not shared.

Specific groups of proteins that have the same combination of structural domains but dramatically different functions illustrate this situation. One example is the combination of the SH3-domain (scop superfamily identifier 2.24.2) and the P-loop containing NTP hydrolase (3.29.1). While in higher organisms this combination is associated with presynaptic and tumor suppressor functions (Swiss-prot names SP02_HUMAN and DLGI_DROME, respectively), in the lower Dictyostelium it was found in myosin (MYSP_DICDI). Another example is the combination of the FAD/NAD(P)-binding superfamily and FAD-linked reductases C-terminal superfamily (3.4.1 and 4.12.1 superfamilies, respectively). In one group of proteins they appear in enzymes of the oxidoreductase group (e.g. OXDA_CAEEL or PHHY_PSEAE), while in another they are found in a dissociation inhibitor (e.g. GDIA_HUMAN). It should be noted that the proteins are not covered completely by the structural matches, so it is quite possible that the rest of them contain totally different domains that are responsible for the dramatically different functions. However, do these two examples show a rather unique or a more frequent phenomenon? How often do multi-domain proteins, sharing the same structural domain composition, differ in their functions?

In this paper, we attempt to provide a comprehensive answer to this question. This is particularly timely given that most of the unknown proteins in eukaryotic genomes are multi-domain. We use the same approach as in our previous analyses, comparing the sequences of the structural domains in scop to those of Swiss-prot using BLASTP. We focus on the functional divergence of single and multi-domain proteins, extending previous investigations of single-domain proteins. Also, in comparison to previous work, we focus more on non-enzymatic functions and scop structural superfamilies, instead of folds.

Results

Our approach to functional and structural assignment

We used the BLASTP program (version 2.0) (Altschul *et al.*, 1997) to identify the scop 1.39 (Murzin *et al.*, 1995) structural domains in Swiss-prot (version 37) (Bairoch &

Apweiler, 2000) with an e-value cutoff of 10^{-4} . We removed the hypothetical and fragment proteins. This resulted in two sets of proteins.

1818 single-domain. Of the single-domain matches only those were selected that were almost completely covered with a match to a single structural domain. (The maximum number of uncovered residues was set at 70 with an additional condition that a maximum of 40 residues on the N-terminal end and 30 residues on the C-terminus were allowed to be uncovered.) These criteria resulted in 1818 single-domain proteins being selected from Swiss-prot.

4763 multi-domain. We selected 4763 multi-domain proteins from Swiss-prot. All of these matched (in different locations) at least two domains of known structure belonging to different scop superfamilies (see schematic in figure 1). We also selected a subset of these proteins that have almost their entire length covered by matches with structural domains (allowing again a maximum of 70 uncovered residues). This selection resulted in 2829 proteins being selected from Swiss-prot. (In all cases, duplicate matches were removed, i.e. a protein at a certain location matches only one structural domain.)

We set out to compare these two sets of proteins for functional divergence. As previously we divided functions into enzyme and non-enzyme (Hegyí & Gerstein, 1999). Enzymatic functions were classified by the EC system (Bairoch, 2000). Comparisons of enzymatic functions were treated the same way as in our earlier analyses, i.e. if they differ in the first 3 components of their respective EC numbers, they were considered different. This implied that our analysis dealt with a total of 112 enzymatic functions. Non-enzymatic functions were classified into 508 different categories based on a simple thesaurus we assembled of synonymous keywords drawn from Swiss-prot description lines. In addition, we created 49 categories for functions that have an enzymatic component but which are not part of the EC system. This gave us a total of 669 functions (112+508+49). (The list of all the functional categories is described further in the caption to Table 2 and also can be found on the web at <http://bioinfo.mbb.yale.edu/partslis/func> or <http://partslis.org/func>.)

Overall distribution of the matches

Figure 2 shows the most commonly observed multi-domain combinations in a set of recently sequenced genomes. The occurrences of further combinations is available from the website. Clearly the distribution is very skewed, with certain combinations, such as 3.29-2.32, and 2.29-4.61 tending to greatly predominate.

Figure 3 shows the overall distribution of the single-domain and multi-domain matches in the different structural classes. The distribution of matches between enzymes and non-enzymes in multi-domain proteins largely agrees with that in the single-domain proteins. The multi-domain matches follow the overall tendency of the alpha/beta folds to be associated with enzymes to a larger extent and the all-alpha and small folds with non-enzymes. However, the values for the multi-domain matches are generally more tempered, less extreme than for single-domains - e.g. the 10-fold difference between

single-domain alpha/beta enzymes and non-enzymes decreases to about two-fold in multi-domain proteins. Another significant difference is the reduction in the number of multi-domain non-enzymes in the all-beta and alpha+beta structural classes compared to the single-domain matches. Altogether, there are more enzymes than non-enzymes among the multi-domain proteins (2805 enzymes vs. 1958 non-enzymes) whereas for single-domain proteins the opposite is true (850 enzymes vs. 968 non-enzymes).

Table 1 summarizes the distribution of superfamilies and superfamily combinations among the major functional classes, i.e. whether they have only enzymatic, only non-enzymatic or both enzymatic and non-enzymatic functionality. Altogether 215 superfamilies were found in single-domain proteins and 310 in multi-domain ones. As 70 superfamilies were found in both, altogether 455 distinct structural superfamilies matched a Swiss-prot protein with our required coverage criteria (described above). Similarly, we apportioned the 281 superfamily combinations observed in multi-domain proteins amongst different broad functional categories.

In single-domain proteins there are about as many superfamilies with exclusively enzymatic functionality as there are those with exclusively non-enzymatic functions (82 vs. 78). In contrast, in multi-domain proteins this ratio increases to almost 3-fold (135 vs. 56). This is in accord with the notion that most enzymes are multi-domain. Another difference between single and multi-domain proteins appears in the ratio of superfamilies with a single function compared to multifunctional ones. As it is apparent from Table 1, about a quarter of the superfamilies matched single-domain proteins with different functions (55 out of 215), whereas in the multi-domain proteins this ratio increased to more than a third (119 out of 310 superfamilies).

Single-domain proteins

In Table 2 we listed the two functionally most diverse structural superfamilies in single-domain proteins with some representative functions. The most diverse superfamily, the 3.38.1 Thioredoxin-like, has 11 different functions associated with it, most of them with an oxidoreductase mechanism. For instance, THIO_BPT4 is a small disulphide-containing thioredoxin that serves as a general disulphide oxidoreductase, while TDX2_BRUMA is almost twice as long (199 aa) and serves as a thiol-specific antioxidant that acts against sulfur-containing radicals. Another interesting example of functional diversity is provided by the Scorpion toxin-like superfamily (7.3.6). While BRAZ_PENBA is a small protein that is known to be 2000 times sweeter than sucrose, the other members of the superfamily are associated with different host-defense mechanisms. In insects the superfamily possesses antifungal activity (DMYC_DROME) or acts as a toxin (SCX5_BUTEU). Interestingly, in plants it can also act as an antifungal (AF2B_SINAL) or as an inhibitor of insect alpha-amylases (SIA1_SORBI). It appears that many single-domain proteins are toxins or allergens or are related in other ways to a host – defense response.

Based on the data we can also determine the probability that if two single-domain proteins match domains in the same superfamily category they also carry out the same function. Using Bayes' theorem:

$$P(F|S) = P(F)P(S|F) / (P(F)P(S|F) + P(\sim F)P(S|\sim F)) \quad (1)$$

where

S: 2 proteins share the same superfamily

F: 2 proteins have the same function

$\sim F$: 2 proteins do not have the same function

Rearranging and simplifying the equation we get:

$$P(F|S) = 1 / (1 + N(S, \sim F) / N(S, F)) \quad (2)$$

where N is the number of times that the two events in the parentheses occur together in our database of 1818 single-domain proteins. This results in

$$P(F|S) = 1 / (1 + 8501 / 12516) = 68\%.$$

That is, the probability that two single-domain proteins that have the same superfamily structure have the same function (whether enzymatic or not) is about 2/3.

Multi-domain proteins

Table 3 lists the combinations of superfamilies that have been associated with the greatest number of different functions in multi-domain proteins, with representative entries in Swiss-prot. The combination with the greatest number of different functions is that of 1.95.1 and 7.33.1. Although it has twice as many different functions as the most diverse superfamily in the single-domain proteins (22 vs. 11, respectively) careful examination reveals that all the proteins in this category are DNA-binding and most of them act as hormone receptors.

The second entry listed in the table is the combination of 3.4.1 and 4.48.1 superfamilies associated with the FAD/NAD(P)-linked reductases. It is an all-enzymatic combination and always carries out an oxido-reductase function. All the proteins in this category are completely covered by matches with these two superfamilies. The 1.78.1-2.1.1 hemocyanin-immunoglobulin combination seems also to be fairly conserved: although the proteins in this category are called by 8 different names, most of them turn out to be extracellular larval storage proteins, except for the copper-containing oxygen carrier hemocyanin itself (HCY_PALVU).

Following the same logic, we can also determine the probability that two proteins that have the same superfamily combination share the same function, viz:

$$P(F|S) = 1/(1+32242/134230) = 81\%$$

This means that we have significantly greater certainty in determining the function of a multi-domain protein with a particular superfamily combination than that of a single-domain protein containing a particular superfamily. We also determined a similar probability for those proteins that have an almost complete coverage with exactly the same type and number of superfamilies, following each other in the same order. The probability that the functions are the same in this case was 91%, a considerably higher value than that arrived at above. However, if two multi-domain proteins share only a single superfamily, the probability that they share the same function drops to only 35%! This greater functional certainty from sharing a combination of superfamilies rather than just one is also reflected in Table 1. While one-fourth of the single-domain proteins and one-third of singularly matching superfamilies in multi-domain proteins have multiple functions, only about one-fifth of the multi-domain *combinations* possess multiple functions (60 out of 281). It is also clear from the data that domains in larger proteins often lose their original function and no longer have an autonomous function.

70 common superfamilies and their functions compared in single-domain and multi-domain proteins

As we mentioned above, of the total of 455 superfamilies in our analysis only 70 occur in both single- and multi-domain proteins. Even more surprising is the small number of structural superfamilies (14) that have the same function in both single- and multi-domain proteins. These listed in Table 4; 12 of them have enzymatic function, supporting the notion that enzymes are more conserved during evolution than non-enzymes. The two non-enzymatic superfamilies are the 4.29.1 ribosomal superfamily and the 5.4.1 superfamily in penicillin-binding proteins.

Table 5 presents several examples of the converse situation, shared superfamilies that have different functions in single and multi-domain proteins. Comparing parts A and B of the table highlights the fact that although both superfamilies in a multi-domain protein are often present in single-domain form as well, the functions in the different settings are only vaguely related. One example is the combination of the lipocalin superfamily (2.45.1) with that of the BPTI-like or Kunitz inhibitor (7.7.1), which in higher organisms forms a complex protein called alpha-1-microglobulin (AMBP_RAT). Another interesting example is the combination of the 2.5.1 Cupredoxin (occurring in the single-domain blue-copper protein, SOXE_SULAC) and the 6.5.1 Membrane all-alpha (single-domain representative: BACT_HALVA, a sensory rhodopsin) superfamilies into a component of the respiratory chain, cytochrome C oxidase II (COOX_ZOOAN). All these examples demonstrate the evolutionary advantage of a domain fusion event, which creates a function that is more complex than either of the components.

Multifunctionality vs. sequence similarity

Previously we presented a variety of graphs that show how the probability that two domains would share the same function varied with respect to sequence similarity (Hegyí & Gerstein, 1999; Wilson *et al.*, 2000). Figure 4 shows a similar graph, now with the calculations extended to multi-domain proteins. It shows that the functional divergence of a single domain in multi-domain proteins dramatically increases, more than two-fold, compared to the single-domain ones. This reinforces our findings above, based only on superfamily content, that the certainty with which we can predict the function of a protein based on its sequence similarity with a domain in another multi-domain protein is considerably less than for a comparable single-domain situation.

Discussion and Conclusions

Here we build on our previous studies on the relationship between protein structure and function to develop new results related to multi-domain proteins. Throughout the paper we focused on superfamilies instead of folds, as the members of a superfamily are presumably of common evolutionary origin (Murzin *et al.*, 1995).

We found that the 4763 multi-domain and 1818 single-domain proteins that met our selection criteria have about the same distribution of structural classes, with more enzymatic functions associated with the alpha/beta structural classes and more non-enzymatic ones with the all-alpha and small classes. We identified more than 3 times as many multi-domain proteins were enzymes than single-domain ones (2805 and 850, respectively) and, conversely, about twice as many multi-domain proteins as single-domain ones that were non-enzymes (1958 vs. 968).

We focused on the functional divergence of the two groups and found that about a quarter of the superfamilies in single-domain proteins are associated with multiple functions, whereas only about a fifth of the multi-domain superfamily combinations are. Therefore, we can conclude that a combination of specific superfamilies results in a more specific functional assignment for a particular protein. However, about one-third of the superfamilies in the multi-domain proteins were associated with multiple functions, underlining the lesser autonomy of a domain function in multi-domain protein.

This latter finding was also supported by the difference in functional divergences between the two groups of proteins based on particular sequence similarities between the domains and Swissprot proteins: as is shown in Figure 4, the average functional divergence of a single domain is much larger (more than two-fold) in multi-domain proteins than in single-domain ones.

We also found that only 70 of a total of 455 superfamilies are shared between the multi-domain and single-domain proteins and only a small fraction, 14, of these share their functions. This was rather surprising to us, and should be taken into consideration in functional characterization and annotation of new gene products. When the functions were related in single- and multi-domain proteins, we could observe an increasing functional complexity with the appearance of large multi-domain proteins.

Altogether, with the recent sequencing of the human genome and genomes of other model organisms, we hope that this work can contribute to the successful annotation of the individual gene products and will help to avoid some pitfalls associated with the functional characterization of large, complex proteins.

References

- Altschul, S. F., Madden, T. L., Schaffer, A. A., Zhang, J., Zhang, Z., Miller, W. & Lipman, D. J. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic Acids Res* **25**, 3389-402.
- Bairoch, A. (2000). The ENZYME database in 2000. *Nucleic Acids Res* **28**, 304-5.
- Bairoch, A. & Apweiler, R. (2000). The SWISS-PROT protein sequence database and its supplement TrEMBL in 2000. *Nucleic Acids Res* **28**, 45-8.
- Chothia, C. & Lesk, A. M. (1986). The relation between the divergence of sequence and structure in proteins. *EMBO J.* **5**, 823-826.
- Devos, D. & Valencia, A. (2000). Practical limits of function prediction. *Proteins* **41**, 98-107.
- Drawid, A & Gerstein, M (2000). A Bayesian system integrating expression data with sequence patterns for localizing proteins: comprehensive application to the yeast genome. *J Mol Biol* 301 : 1059-75.
- Eisenstein, E., Gilliland, G. L., Herzberg, O., Moulton, J., Orban, J., Poljak, R. J., Banerjee, L., Richardson, D. & Howard, A. J. (2000). Biological function made crystal clear - annotation of hypothetical proteins via structural genomics. *Curr Opin Biotechnol* **11**, 25-30.
- Gelbart, W. M., Crosby, M., Matthews, B., Rindone, W. P., Chillemi, J., Russo Twombly, S., Emmert, D., Ashburner, M., Drysdale, R. A., Whitfield, E., Millburn, G. H., de Grey, A., Kaufman, T., Matthews, K., Gilbert, D., Strelets, V. & Tolstoshev, C. (1997). FlyBase: a Drosophila database. The FlyBase consortium. *Nucleic Acids Res* **25**, 63-6.
- Gerstein, M (1997). A structural census of genomes: comparing bacterial, eukaryotic, and archaeal genomes in terms of protein structure. *J Mol Biol* 274 : 562-76.
- Gerstein, M. (1998). How representative are the known structures of the proteins in a complete genome? A comprehensive structural census. *Fold Des* **3**, 497-512.
- Harrison, P, Echols, N & Gerstein, M (2001). Digging for Dead Genes: An Analysis of the Characteristics of the Pseudogene Population in the C. elegans Genome. *Nuc. Acids. Res.* 29 : 818-30.
- Hegyí, H. & Gerstein, M. (1999). The relationship between protein structure and function: a comprehensive survey with application to the yeast genome. *J Mol Biol* **288**, 147-64.
- Lin, J & Gerstein, M (2000). Whole-genome trees based on the occurrence of folds and orthologs: implications for comparing genomes on different levels. *Genome Res* 10 : 808-18.

- Martin, A. C., Orengo, C. A., Hutchinson, E. G., Jones, S., Karmirantzou, M., Laskowski, R. A., Mitchell, J. B., Taroni, C. & Thornton, J. M. (1998). Protein folds and functions. *Structure* **6**, 875-84.
- Murzin, A., Brenner, S. E., Hubbard, T. & Chothia, C. (1995). SCOP: A Structural Classification of Proteins for the Investigation of Sequences and Structures. *J. Mol. Biol.* **247**, 536-540.
- Orengo, C. A., Pearl, F. M., Bray, J. E., Todd, A. E., Martin, A. C., Lo Conte, L. & Thornton, J. M. (1999). The CATH Database provides insights into protein structure/function relationships. *Nucleic Acids Res* **27**, 275-9.
- Pawlowski, K., Jaroszewski, L., Rychlewski, L. & Godzik, A. (2000). Sensitive sequence comparison as protein function predictor. *Pac Symp Biocomput* 42-53.
- Pearson, W. R. (1994). Using the FASTA program to search protein and DNA sequence databases. *Methods Mol Biol* **25**, 365-89.
- J Qian, B Stenger, C Wilson, J Lin, R Jansen, W Krebs, V Alexandrov, N Echols, S Teichmann, J Park, M Gerstein.
"PartsList: a web-based system for dynamically ranking protein folds based on disparate attributes, including whole-genome expression and interaction information." *Nucleic Acids Res* 29: 1750-64 (2001).
- Russell, R. B., Saqi, M. A., Sayle, R. A., Bates, P. A. & Sternberg, M. J. (1997). Recognition of analogous and homologous protein folds: analysis of sequence and structure conservation. *J Mol Biol* **269**, 423-39.
- Shah I, Hunter L (1997). Predicting enzyme function from sequence: a systematic appraisal. *Proc Int Conf Intell Syst Mol Biol* 5:276-83
- Shapiro, L. & Harris, T. (2000). Finding function through structural genomics. *Curr Opin Biotechnol* **11**, 31-5.
- Stawiski EW, Baucom AE, Lohr SC, Gregoret LM (2000). Predicting protein function from structure: unique structural features of proteases. *Proc Natl Acad Sci U S A* 97(8):3954-8
- Thornton, J. M., Orengo, C. A., Todd, A. E. & Pearl, F. M. (1999). Protein folds, functions and evolution. *J Mol Biol* **293**, 333-42.
- Thornton, J. M., Todd, A. E., Milburn, D., Borkakoti, N. & Orengo, C. A. (2000). From structure to function: approaches and limitations [In Process Citation]. *Nat Struct Biol* **7 Suppl**, 991-4.
- Todd AE, Orengo CA, Thornton JM (2001). Evolution of function in protein superfamilies, from a structural perspective. *J Mol Biol* **307**,1113-43
- Wilson, C. A., Kreychman, J. & Gerstein, M. (2000). Assessing annotation transfer for genomics: quantifying the relations between protein sequence, structure and function through traditional and probabilistic scores. *J Mol Biol* **297**, 233-49.

Zhang, B., Rychlewski, L., Pawlowski, K., Fetrow, J. S., Skolnick, J. & Godzik, A. (1999). From fold predictions to function predictions: automation of functional site conservation analysis for functional genome predictions. *Protein Sci* **8**, 1104-15.

Figure Captions and Table Legends

Table 1, Overall distribution of structures and functions

The basic functional distribution of the superfamilies in single and multi-domain proteins and the functional distribution of multi-domain combinations. The first row lists the number of scop superfamilies that were associated only with enzymatic function in each category. The second row lists the number associated only non-enzymatic functions, the third row indicates the number of superfamilies that were associated with both types of function. Altogether we characterized $160+55=215$ single-domain and $191+119=310$ multi-domain superfamilies, 70 of which overlapped in the two categories.

Table 1. Functional distribution of singledomain, multidomain superfamilies and multidomain combinations

	Singledomain superfamilies		Multidomain superfamilies		Multidomain sfam combinations	
	Single function	Multiple function	Single function	Multiple function	Single function	Multiple function
enzymatic	82	11	135	42	151	16
nonenzymatic	78	23	56	30	70	27
both functions	-	15	-	47	-	17
total	160	55	191	119	221	60

Table 2, Most versatile single-domain superfamilies

The most versatile superfamilies in single-domain proteins as determined from their functional description in Swiss-prot, with some representatives. The keyword combinations in the 4th column were either based on the first 3 components of their EC numbers (for enzymes) or derived automatically by comparing the DE description line of Swiss-prot entries to a list of synonymous keywords at <http://bioinfo.mbb.yale.edu/partslist/func>. (A keyword number starting with a D indicates an enzyme that does not have an assigned EC number in its description in Swiss-prot.)

No. func	No. prot	Sfam comb	Function	Swissprot ID	Swissprot function
11	69	3.38.1	E1.11.1 263# D260# 268# 266# 269# 272# D271#272# 261#	GSHP_RAT DYL5_CHLRE BSAA_BACSU REHY_TORRU PHOS_HUMAN REHY_ORYSA THIO_BPT4 TDX2_BRUMA BTUE_ECOLI	Plasma Glutathione Peroxidase (1.11.1.9) Dynein , Flagellar Outer Arm - C.Reinhardtii Glutathione Peroxidase Homolog Bsa Rehydrin - Tortula Ruralis (Moss) Phosducin (33 Kd Phototransducing Protein) Rab24 Protein - Oryza Sativa (Rice) Thioredoxin (Bacteriophage T4) Thioredoxin Peroxidase 2 Vitamin B12 Transport Periplasmic Protein Btue
10	28	7.3.6	342# 376#336# 341#356# 343# 361# 361#376# 336# 203#	BRAZ_PENBA SCKK_TITSE AF2B_SINAL DEFA_ZOPAT DMYC_DROME SCX5_BUTEU SCX3_LEIQH SIA1_SORBI	Brazzein - Pentadiplandra Brazzeana Neurotoxin Ts-Kapa (Tsk) - (Brazilian Scorpion) Cysteine-Rich Antifungal Protein 2b (Afp2b) Defensin, Isoforms B And C - Zophobas Atratus Drosomycin Precursor (Cysteine-Rich Peptide) Insectotoxin I5a - (Lesser Asian Scorpion) Leuropeptide Iii - (Scorpion) Small-Pr Inhibitor Of Insect Alpha-Amylases
7	34	4.79.3	310# 311# 231# 312# E3.1.- 314#	AB18_PEA DRR3_PEA MPAA_CORAV L18B_LUPLU RNS2_PANGI SAM2_SOYBN	Aba-Responsive Protein Abr18 - Garden Pea Disease Resistance Response Protein Pi49 Major Pollen Allergen Cor A 1, - Eu. Hazel Protein L1r18b (Lpr10.1b) Ribonuclease 2 (3.1.-) - Panax Ginseng Stress-Induced Protein Sam22
7	43	1.26.1	184# 381#564#184# 185# 187# 186# 188#	CSF2_SHEEP IL4_RAT LIF_HUMAN PRL_ANGAN PLF3_MOUSE SOMA_PAROL	Colony-Stimulating Factor Interleukin-4 (B-Cell Igg Diff. Factor) Leukemia Inhibitory Factor (Lif) Prolactin Precursor (Pr) - Proliferin 3 Mitogen-Regulated Somatotropin (Growth Hormone)

Table 3, Most versatile superfamily combinations in multi-domain proteins

Note, the combination with the greatest number of different functions is that of 1.95.1 and 7.33.1. Careful examination reveals that all the proteins with this combination are DNA-binding and most of them act as various hormone receptors. (In particular, HNF4_DROME and NR42_HUMAN also have transcription activator functions. Note that these two proteins are considerably longer than the others in this group and are not covered completely by structural matches: a large C-terminal and a large N-terminal portion are left uncovered respectively.)

No. func	No. prot	Sfam comb.	Function	Swissprot ID	Swissprot function
22	176	1.95.1/7.33.1	29# 10# 31#32# 29#30# 32# 556#564#35# 576# 36# 37# 38#	THB_RANCA HNF4_DROME EAR2_MOUSE ECR_MANSE ERBA_AVIER NGFI_XENLA NR42_HUMAN PPAT_HUMAN RXRG_CHICK TLL_DROVI	Thyroid Hormone Receptor Beta Transcription Factor HNF-4 Homolog V-Erba Related Protein Ear-2 Ecdysone Receptor (Ecdysteroid Receptor) Erba Oncogene Protein Nerve Growth Factor Induced Protein I-B Immediate-Early Response Protein Not Peroxisome Proliferator Activated Receptor Retinoic Acid Receptor RXR-Gamma Tailless Protein
8	54	3.4.1/4.48.1	E1.8.2 E1.8.1 E1.6.4 E1.16.1 E1.6.99	DHSU_CHRVI DLDH_ZYMMO TYTR_TRYCR MERA_STRLI NAOX_MYCPN	Sulfide Dehydrogenase (1.8.2.-) Dihydrolipoamide Dehydrogenase (1.8.1.4) Trypanothione Reductase (1.6.4.8) (Tr) Mercuric Reductase (1.16.1.1) Probable NADH Oxidase (1.6.99.3) (Noxase)
8	23	1.78.1/2.1.1	19# 20# 21#427# 22# 23# 24# 546#25#	ARYB_MANSE CRPI_PERAM HCY_PALVU HEXA_BLADI JSP1_TRINI LSP2_DROME SSP1_BOMMO	Arylphorin Beta Subunit-(Tobacco Hornworm) Allergen Cr-Pi Precursor - (Am. Cockroach) Hemocyanin - (European Spiny Lobster) Hexamerin Precursor - (Tropical Cockroach) Acidic Juvenile Hormone-Suppressible Protein Larval Serum Protein 2 Precursor (LSP-2) Sex-Specific Storage-Protein 1

Table 4

Superfamilies with the same function in single- and multi-domain proteins as determined from their keyword combination or first 3 components of their EC numbers.

Sfam	function	Singedomain proteins		Multidomain proteins	
		Swiss. ID	Swissprot function	Swiss. ID	Swissprot function
1.81.1	E3.2.1	GUNY_ERWCH	Endoglucanase (3.2.1.4)	AMYG_NEUCR	Glucoamylase Precursor (3.2.1.3)
2.66.2	E3.5.1	URE2_YERPS	Urease Beta (3.5.1.5)	URE1_HELPY	Urease Alpha Subunit (3.5.1.5)
3.17.2	E6.3.5	NADE_MYCPN	NAD(+) Synthetase (6.3.5.1)	GUAA_YEAST	GMP Synthase (6.3.5.2)
3.37.1	E3.1.3	PTP2_NPVOP	Protein-Tyrosine Phosphatase 2 (3.1.3.48)	PTNB_RAT	Protein-Tyrosine Phosphatase (3.1.3.48)
3.67.1	E4.2.1	TRPB_VIBPA	Tryptophan Synthase (4.2.1.20)	TRP_YEAST	Tryptophan Synthase (4.2.1.20)
4.19.1	E5.2.1	FKB1_METJA	Peptidylprolyl Cis-Trans Isomerase (5.2.1.8)	FKB7_WHEAT	70 Kd Peptidylprolyl Isomerase (5.2.1.8)
4.2.1	E3.2.1	LYCV_BPP2	Lysozyme (3.2.1.17)	CHIX_PEA	Endochitinase Precursor (3.2.1.14)
4.29.1	85#	RS5_ACYKS	30s Ribosomal Protein S5	RS5_TREPA	30s Ribosomal Protein S5
4.52.1	E3.4.24	SNPA_STRCS	Extracellular Neutral Protease (3.4.24.-)	BMPH_STRPU	Collagenase 3 Precursor (3.4.24.-)
4.6.1	E3.5.1	URE3_YERPS	Urease Gamma (3.5.1.5)	URE1_HELPY	Urease Alpha Subunit (3.5.1.5)
5.10.1	E2.7.7	KANU_STAAU	Kanamycin Nucleotidyltransferase (2.7.7.-)	DPOB_XENLA	Dna Polymerase Beta (2.7.7.7)
5.4.1	161#	AMPH_ECOLI	Penicillin-Binding Protein Amph	PBPX_STREN	Penicillin-Binding Protein 2x Pbp2x

Table 5

Examples of superfamilies present in both single- and multi-domain proteins, carrying out different functions. A/ Single-domain proteins, B/ Multi-domain proteins.

Part A

Sfam	Funct #	Swissprot ID	Swissprot function
1.25.1	352#	FTN2_HAEIN	Ferritin Like Protein 2
	183#	NIGY_DESVH	Nigerythrin
	E1.17.4	RIR4_YEAST	(Ribonucleotide Reductase) (1.17.4.1)
	192#	NLP_HAEIN	Ner-Like Protein Homolog
1.4.3	196#	H1A_PLADU	Histone H1A, Sperm
1.81.2	E2.5.1	PFTB_PEA	Farnesyltransferase Beta Su (2.5.1.-)
2.45.1	226#	ERBP_RAT	Epididymal-Retinoic Acid Binding Protein
	227#	FAB3_CAEEL	Fatty Acid-Binding Protein Homolog 3
	228#412#	NGAL_MOUSE	Neutrophil Gelatinase-Assoc. Lipocalin
	229#	NP4_RHOPR	Nitrophorin 4 Precursor
	E5.3.99	PGHD_HUMAN	Prostaglandin-H2 D-Isomerase (5.3.99.2)
	230#421#	VNS1_MOUSE	Vesomeral Secretory Protein I
2.5.1	231#	MPA3_AMBEL	Pollen Allergen AMB A 3 (AMB A Iii)
	232#427#	SOXE_SULAC	Sulfocyanin (Blue Copper Protein)
3.14.2	373#	RRF1_DESVH	Rrf1 Protein
3.29.1	E6.3.4	PURA_CAEEL	Adenylosuccinate Synthetase (6.3.4.4)
	E2.7.4	KTHY_YEAST	Thymidylate Kinase (2.7.4.9)
	D259#	VA57_VACCV	Guanylate Kinase Homolog
	E2.7.1	KITH_VZVW	Thymidine Kinase (2.7.1.21)
3.47.1	275#	MBL_BACSU	MBL Protein
	276#	MREB_BACSU	Rod Shape-Determining Protein Mreb
3.48.1	E3.1.3	PPA5_YEAST	Repressible Acid Phosphatase (3.1.3.2)
3.81.1	D281#	AMIC_PSEAE	Aliphatic Amidase Expression-Regulator
	282#	LUXP_VIBHA	LUXP Protein Precursor
4.103.1	E2.4.2	TOX1_BORPE	Pertussis Toxin Su 1 (2.4.2.-)
4.105.1	291#	LECC_POLMI	Lectin - Polyandrocarpa Misakiensis
4.11.5	295#	TERP_PSESP	Terpredoxin
4.19.1	E5.2.1	FKB1_METJA	Pept-Prolyl Cis-Trans Isomerase (5.2.1.8)
6.5.1	E3.6.1	ATPL_VIBAL	ATP Synthase (3.6.1.34) (Lipid-Binding)
	540#325#	BACT_HALVA	Sensory Rhodopsin II (Sr-li)
7.35.4	E1.9.3	COXB_RAT	Cytochrome C Oxidase (1.9.3.1) (Via*)
	345#	DESR_DESGI	Desulforedoxin (Dx)
7.7.1	349#	TAP_ORNMO	Tick Anticoagulant Peptide

Table 5B

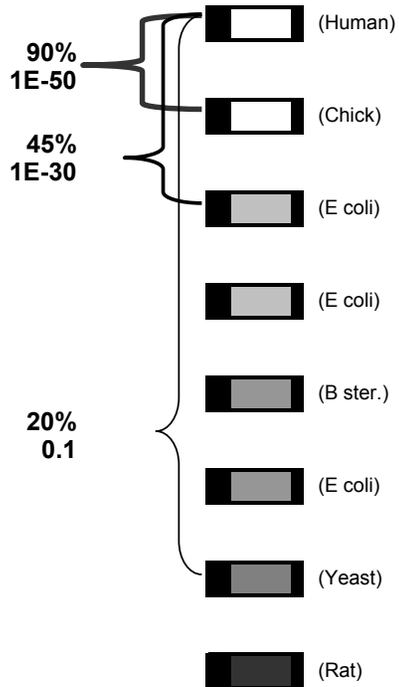
Sfam Comb.	Funct #	Swissprot ID	Swissprot function
1.25.1/7.35.4	104#	RUBY_METJA	Putative Rubrerythrin
1.32.1/3.81.1	11#	PURR_HAEIN	Purine Nucleotide Synthesis Repressor
	12#	DEGA_BACSU	Degradation Activator
	581#11#	SCRR_STRMU	Sucrose Operon Repressor
	582#11#	REGA_CLOAB	Transcription Regulatory Protein Rega
1.4.3/3.14.2	10#	SKN7_YEAST	Transcription Factor Skn7 (Pos9 Protein)
	11#	VIRG_AGRT5	Virg Regulatory Protein
	13#	RGX3_MYCTU	Sensory Transduction Protein REGX3
	190#	PFER_PSEAE	Transcriptional Activator Protein Pfer
	366#	PETR_RHOCA	Petr Protein
2.45.1/7.7.1	203#153#	HC_RAT	Alpha-1-Microglobulin / Trypsin Inhibitor
2.5.1/6.5.1	E1.9.3	COX2_ZOOAN	Cytochrome C Oxidase li (1.9.3.1)
3.29.1/3.48.1	E2.7.1	F26_RANCA	6-Phosphofructo-2-Kinase (2.7.1.105)
3.47.1/5.17.1	1#	YED0_YEAST	Heat Shock Protein 70 Homolog YEL030w
	1#83#	GR73_MAIZE	Ig-Binding Protein

Figure 1, Schematic Illustrating Annotation Transfer

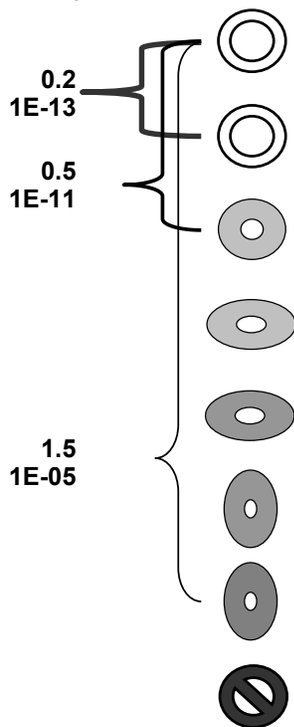
The figure illustrates the process of annotation transfer a group of hypothetical TIM barrel proteins. The leftmost panel represents sequence comparisons between an idealized barrel domains from a number of organisms. Moving rightwards, the next panel shows analogous results for structural comparison, and panel after that, functional comparison. The rightmost panel represents sequence comparisons between idealized multi-domain proteins that match over a single domain, the subject of much of this paper.

Sequence Similarity (%ID, P-seq)

Single Domain Proteins



Structural Similarity (RMS, P-str)

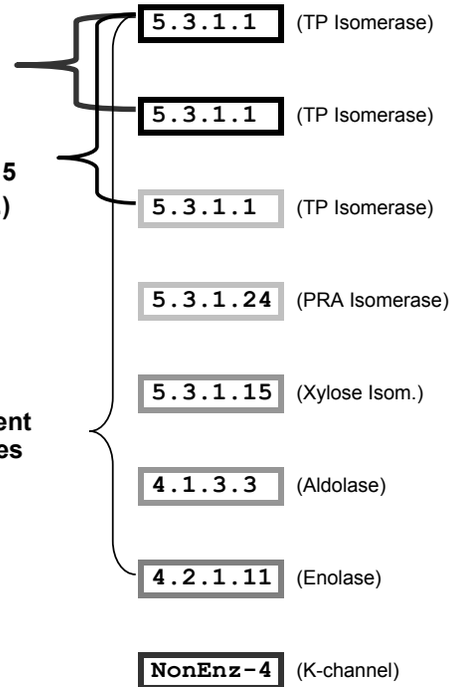


Functional Similarity (%Same-Func.)

Same Exact

Both Class 5 (isom.)

Different Classes



Sequence Similarity (%ID, P-seq)

Multi Domain Proteins

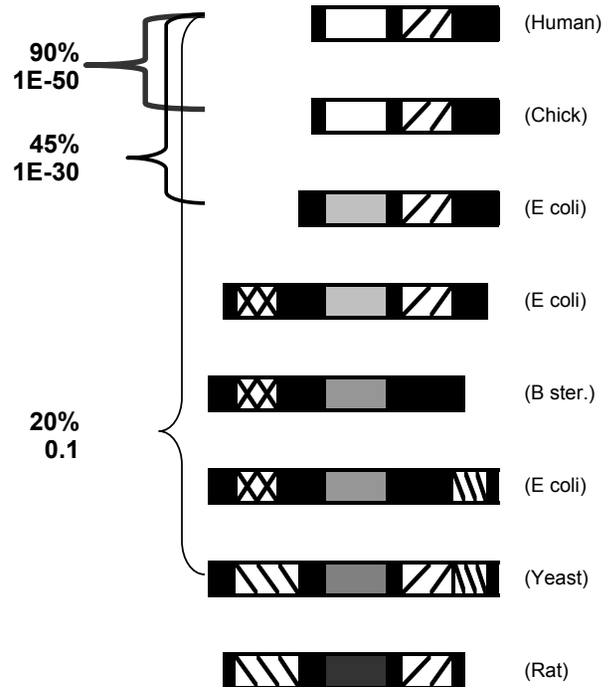


Figure 2, Distribution of Multidomain Combinations amongst the Genomes

The figure shows the occurrence of multidomain fold combinations in a number of genomes, indicating its great variability. Each row indicates a particular combination of scop fold pairs (using scop 1.39), where a fold pair is defined as two distinct folds occurring in tandem in a protein. Each column represents a different genome, using the four letter codes in the PartsList system (Qian *et al.*, 2001), viz: Aaeo, Aquifex aeolicus; Aful, Archaeoglobus fulgidus; Bbur, Borrelia burgdorferi; Bsub, Bacillus subtilis; Cele, Caenorhabditis elegans; Cpne, Chlamydia pneumoniae; Ctra, Chlamydia trachomatis; Ecol, Escherichia coli; Hinf, Haemophilus influenzae Rd; Hpyl, Helicobacter pylori; Mthe, Methanobacterium thermoautotrophicum; Mjan, Methanococcus jannaschii; Mtub, Mycobacterium tuberculosis; Mgen, Mycoplasma genitalium; Mpne, Mycoplasma pneumoniae; Phor, Pyrococcus horikoshii; Rpro, Rickettsia prowazekii; Scer, Saccharomyces cerevisiae; Syne, Synechocystis sp.; Tpal, Treponema pallidum. The numbers in each intersection cell indicate the number of times the fold pairs occur in a genome. Only the 20 most common fold pair combinations are shown here; the remainder are shown on the website (partslist.org/func). If a cell is greater than 6, it is shaded black, between 3 and 6, gray, and below 3, white. The blank spaces show instances in which one of the pairs do not occur in the organism at all (indicated by a value of -1 in the data table on the website). The fold assignments are done in a fashion consistent with those in PartsList and associated systems (Qian *et al.*, 2001; Lin *et al.*, 2000; Gerstein, 1997; Harrison *et al.*, 2001; Drawid *et al.*, 2001).

FOLD PAIRS

fold 1	fold 2	aful	mjan	mthe	phor	scer	cele	aaeo	syne	ecol	bsub	mtub	hinf	hpyl	mgen	mpne	bbur	tpal	ctra	cpne	rpro
3.29	2.32	4	3	4	3	12	14	6	7	8	4	6	7	5	3	3	4	5	3	4	4
2.29	4.61	1	1	1	2	6	3	2	4	5	4	4	3	3	3	4	1	2	3	3	2
4.1	4.34	1	1	1	1	5	3	1	3	1	1	2	1	1	1	1	2	2	1	1	1
1.28	3.29	1	1	2	1	1	1	1	1	2	1	1	2	1	1	1	1	1	1	1	1
3.4	4.48	4	1	1	2	3	4	1	2	4	3	5	2		2	2	1	1	1	1	2
3.22	4.42	1	1	1	0	4	5	3	4	5	4	4	3	4	1	1	2	2	3	3	1
2.32	4.1	1	1	1	1	4	2	1	3	1	1	2	1	1	1	1	2	2		1	1
2.32	2.33	2	1	1	1	1	2	2	1	2	1		2	1	1	1	1	1	1	1	1
4.32	3.1	1	1	1	2	5	1	1	1	4	5	1	1	1	1	1	1	1	1	1	0
3.23	4.89	3	3	3	0	9	10	6	5	6	8	7	2	4	0	0	1	1	2	2	2
3.47	5.17	0	0	1	0	12	10	1	3	3	1	1	2	1	1	1	2	1	1	1	2
4.72	5.13	1	0	0	0	1	3	1	1	2	2	1	2	1	2	2	2	1	2	2	2
3.22	4.1	1	1	1	0	3	3	2	1	1	2	1	1	0	1	1	1	0	1	1	1
3.5	3.1	1	1	2	1			1	1	5	1		1	1	1	1	1	1	1	1	1
4.61	3.42	2	2	2	2	2	1	1	1	1	1	1	1	0	2	2	1	1	1	0	0
1.76	3.3	1	0	1	1	2	1	1	1	1	2	1	1	1	0	0	0	1	1	1	1
4.29	4.1	1		1	1	2	1	1	1	1	1		1		0	1	1	1	1	1	1
2.32	4.34					2	1	1	2	1	2	2	1	2	1	1	1	1	1	1	1
3.22	1.79	1	1	1	0	3	1	2	2	2	4	3	2	1	0	0	1	1	1	1	0
3.52	2.34	0	0	0	0	0	1	2	1	2	1	1	1	1	1	1	1	1	1	1	1

Figure 3, Distribution of proteins amongst broad structural and functional classes

The distribution of the matches among the 7 structural and 2 functional classes in single- and multi-domain proteins. The single-domain and multi-domain matches each total to 100%, independently of each other. The horizontal axis indicates the 7 scop classes, which are from 1 to 7: all-alpha, all-beta, alpha/beta, alpha+beta, multi-domain, membrane, and small protein.

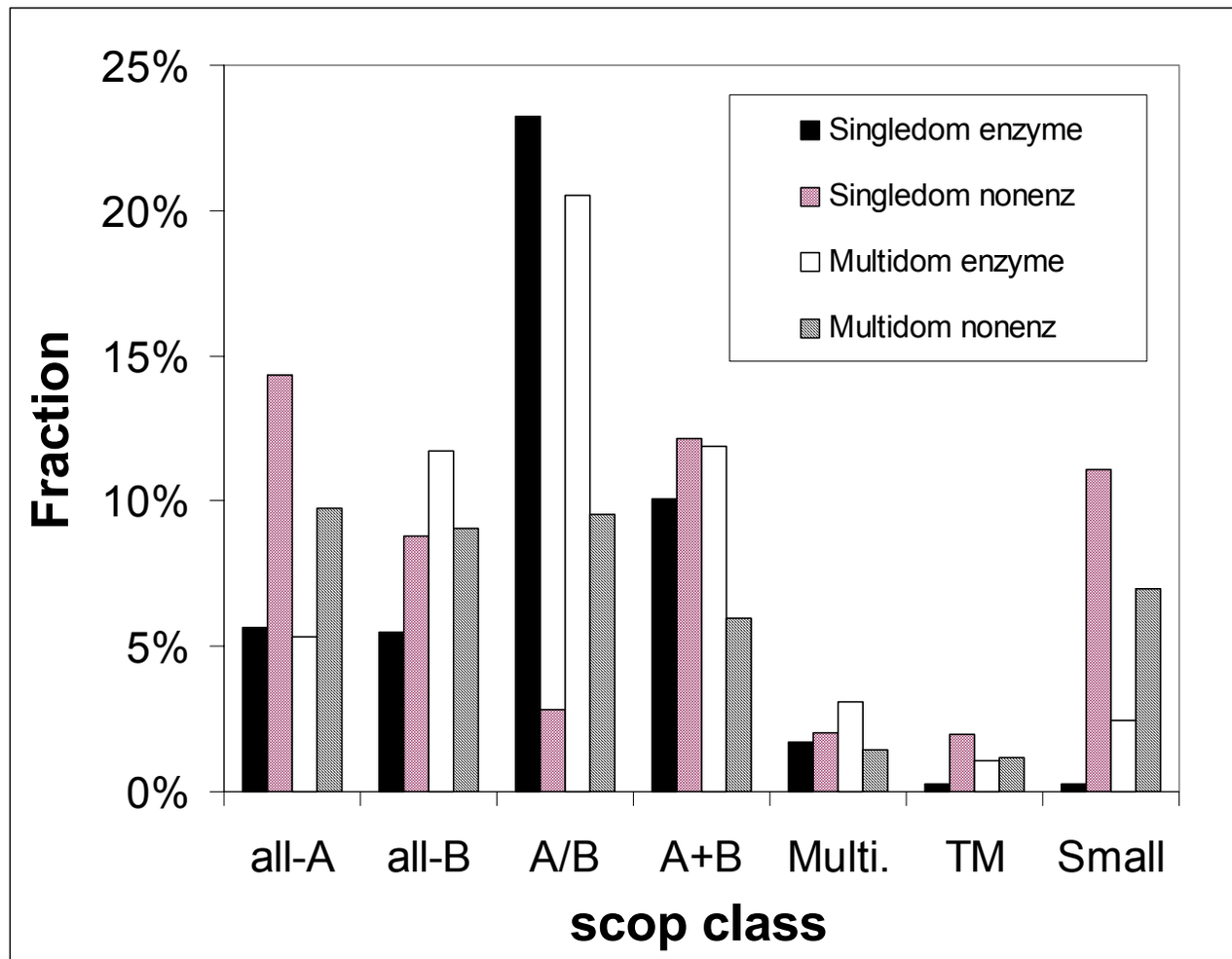


Figure 4, Divergence in function with respect to sequence similarity

Relative number of matching domains with multiple functions, as the function of e-value threshold. Diamonds represent single-domain proteins, squares multi-domain ones (matching just for a single domain), respectively. The first value on the x-axis starts at 4 (corresponding to an e-value= 10^{-4}).

