UNIVERSITY OF CALIFORNIA, SAN DIEGO

Evolution of the Oligopeptide Transporter (OPT) Family

A Thesis submitted in partial satisfaction of the requirements for the degree Master of Science

in

Biology

by

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Parts of this Thesis are being prepared for publication. The thesis author will be the primary investigator and author of this paper.

ABSTRACT OF THE THESIS

Evolution of the Oligopeptide Transporter (OPT) Family

by

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Professor Milton H. Saier, Jr., Chair

The Oligopeptide Transporter (OPT) family of peptide and iron-siderophore transporters includes members in both prokaryotes and eukaryotes but with restricted distribution in the latter domain. All functionally characterized peptide transporters segregate from the iron-siderophore transporters on a phylogenetic tree. Prokaryotic members derive from many different phyla, but they belong only to the iron-siderophore subdivision. This fact suggests, but does not prove, that this family arose in prokaryotes, and that the peptide transporters arose from iron-siderophore transporters in eukaryotes. Eukaryotic members are found only in fungi and plants with a single slime mold homologue clustering with the fungal proteins, suggestive of horizontal transfer from a fungus.

OPT family proteins have 16, or occasionally 17 transmembrane spanning α -helical segments. We provide statistical evidence that the 16 TMS topology arose via three sequential duplication events followed by a gene fusion event for proteins with a seventeenth TMS. 2 TMSs \rightarrow 4 TMSs \rightarrow 8 TMSs \rightarrow 16 TMSs \rightarrow 17 TMSs. The seventeenth C-terminal TMS, which probably arose just once, is found in a restricted phylogenetic group of these homologues. Analyses for orthology revealed that a few phylogenetic clusters consist exclusively of orthologs, but most have undergone intermixing, suggestive of horizontal transfer. The results suggest that in this family, horizontal gene transfer was frequent among prokaryotes, rare among eukaryotes and totally absent between prokaryotes and eukaryotes as well as between plants and fungi. These observations provide evidence concerning the pathway taken for the evolution of this family. They also provide guides for future structural and functional analyses.

Parts of the Abstract are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Introduction

Transport proteins found in eukaryota, bacteria, and archaea can be grouped into four distinct classes as described in the Transporter Classification Database, TCDB (Saier, 2000a; Saier et al., 2006; Saier et al., 2009). The first class is composed of channels/pores which catalyze facilitated diffusion (by an energy-independent process) through a transmembrane aqueous pore or channel without a carrier-mediated mechanism. These channels/pores do not exhibit stereospecificity but may be specific for a particular molecular species or class of molecules (Saier, 2000b). The electrochemical potential-driven transporters, comprising the second class, utilize a carrier-meditated process to catalyze uniport (a single species is transported by facilitated diffusion in a process that is not coupled to the utilization of an energy source), antiport (two or more species are transported in the opposite direction in a tightly coupled process which utilizes chemiosmotic energy), and/or symport (two or more species are transported together in the same direction in a tightly coupled process which also utilizes chemiosmotic energy) (Saier, 2000c; Busch & Saier, 2004). In contrast to channels of class 1, carriers of class 2 are usually stereospecific. The third class consists of the primary active transporters which utilize a primary source of energy (chemical, electrical, and/or solar) to drive active transport of a solute against a concentration gradient (Saier, 2000a). Group translocators of the fourth class utilize a primary energy source to chemically alter a substrate as it is transported across a membrane; this alteration makes the transported substance impermeable to the membrane, thus localizing it to a new location (Mitchell and Moyle, 1958; see TCDB, www.tcdb.org).

The oligopeptide transporter (OPT; TC# 2.A.67) family is a member of the second class of transport proteins, the electrochemical potential-driven transporters. All functionally characterized members of this family catalyze uptake of their solutes by a cation:solute symport mechanism (Hauser *et al.*, 2001; Lubkowitz, 2006; Yen *et al.*, 2001). Members consist of transporters specific for oligopeptides (3-8 amino acids) found in fungi, plants, slime molds, and prokaryotes (Yen *et al.*, 2001). These proteins occur within many phyla of bacteria and archaea. The OPT family is not to be confused with the proton-dependent oligopeptide transporter (POT or PTR; TC# 2.A.17) family (Paulsen and Skurray, 1994), the peptide transporters (PepT) of the ATP binding cassette (ABC; TC# 3.A.1.5) superfamily (Saier, 2000a; Busch & Saier, 2004), or the peptide-acetyl-CoA transporters (PAT) of the major facilitator superfamily (MFS; TC# 2.A.1.25) (Pao *et al.*, 1998).

The OPT family can be broken down into two clades, the peptide transporters and the yellow stripe-type iron-complex transporters (YS) (Lubkowitz, 2006; Yen *et al.*, 2001). Peptide transporter homologues primarily transport oligopeptides, glutathione, glutathione conjugates, and various other glutathione derivatives (Kaur *et al.*, 2009; Lubkowitz *et al.*, 1998). Characterized YS homologues, on the other hand, mediate the uptake of metal-chelating phytosiderophores, including iron-nicotinamide, and complexes of iron with secondary amino acids such as mugineic acid and deoxymugineic acid (Kaur *et al.*, 2009).

The biochemical and physiological characteristics of several OPT homologues have been studied (Lubkowitz, 2006; Osawa *et al.*, 2006; Stacey *et al.*, 2008; Thakur *et al.*, 2008). Two highly conserved motifs (NPG and KIPPR) have been found among all

or most OPT family proteins (Koh *et al.*, 2002). The two generalized transport reactions known to be catalyzed by functionally characterized members of the OPT family are:

- 1) Oligopeptide (out) + nH^+ (out) \rightarrow Oligopeptide (in) + nH^+ (in)
- 2) Fe^{3+} -phytosiderophore (out) + nH^+ (out) $\rightarrow Fe^{3+}$ -phytosiderophore (in) + nH^+ (in).

The transport of oligopeptides plays an important role in nitrogen storage and mobilization, quorum sensing in bacteria, bacterial differentiation, sexual induction in Gram-positive bacteria, yeast mating, and pheromone and hormone sensing in animals. One of the yeast homologues is the sexual differentiation process (ISP4) protein of *Schizosaccharomyces pombe*. In yeast, OPT family homologues transport oligopeptides which are commonly tri-, tetra-, and/or pentapeptides (Wiles *et al.*, 2006). Recently, it has been found that high-affinity *S. cerevisiae* and *S. pombe* glutathione transporters, Hgt1p and OPT1, respectively, belong to the OPT family (Dworeck *et al.*, 2009; Kaur *et al.*, 2009). These proteins appear to be the sole or dominant glutathione transporters in these species. Both OPT1 and Hgt1p localize to the plasma membrane (Dworeck *et al.*, 2009).

In *C. albicans*, eight OPT genes have been identified encoding putative oligopeptide transporters. Almost all are represented by polymorphic alleles (Reuss and Morschhäuser, 2006). OPT 1,2,3 Δ triple mutants were found to have a severe growth defect which could be rescued by reintroduction of a single copy of OPT1, OPT2, or OPT3. The various oligopeptide transporters differ in their substrate preferences as shown by the ability of strains expressing specific OPT genes to grow on peptides of defined length and sequence (Reuss and Morschhäuser, 2006).

In plants, many OPTs appear to be plasma membrane-embedded proteins that import substrates from the apoplasm (the aqueous phase of the cell wall) and the external environment. They may play a role in plant growth and development (Lubkowitz, 2006). Unlike many other OPTs which function in long-distance transport of peptides or metals, YS1, an Fe³⁺-phytosiderophore uptake system of *Zea mays*, is known to translocate substrates from the rhizosphere (the region of the soil that is directly influenced by root secretions and associated with soil microbes) (Yen *et al.*, 2001; Curie *et al.*, 2001). The expression of the YS1 gene is increased in roots and shoots under iron deficient conditions (Curie *et al.*, 2001). When YS1 is expressed in mutant yeast lacking its native iron uptake system, it is able to correct the defect, specifically in Fe³⁺-phytosiderophore-containing media.

In *Arabidopsis*, nine OPT paralogues have been identified. All of them show highly significant sequence similarity to OPTs found in *C. albicans* (e.g., CaOpt1p), *S. pombe* (e.g., Isp4p), and *S. cerevisiae* (e.g., Opt1p and Opt2p) (Koh *et al.*, 2002). Of the OPT homologues found in *Arabidopsis*, seven of them mediate the transport of tetra- and pentapeptides while two are believed to transport glutathione and its conjugates (Cagnac *et al.*, 2004). For example, Cagnac *et al.* (2004), showed that AtOPT6 can mediate uptake of glutathione derivatives and metal complexes, which led them to suggest that it may also be involved in stress resistance. OPT homologues found in rice (*Oryza sativa*) and Indian mustard (*Brassica juncea*) have also been described as glutathione derivative transporters (Cagnac *et al.*, 2004).

Bacterial and archaeal homologues of the OPT family have yet to be studied, but as shown here, they are prevalent throughout the prokaryotic world. Currently, little

information is available concerning the detailed mechanistic characteristics of these OPT family members (Kaur et al., 2009). A high-resolution 3-dimensional x-ray structure of an OPT family homologue has yet to be solved. We therefore carried out detailed bioinformatics analyses of these transporters. We show that the family is far more widespread than previously recognized and demonstrate the evolutionary relationships of the members of this family to each other. Most surprising, we found that these 16 TMS proteins arose from a 2 TMS precursor-encoding genetic element which duplicated three times sequentially. It was inferred from the fact that the first and third repeats, as well as the second and fourth repeats are substantially more similar to each other than are any other pairs of repeats were, that the two last duplication events $4 \rightarrow 8$ and $8 \rightarrow 16$, were separated from each other by a substantial period of time. Although this finding is in principle similar to the origin of animal Na⁺ and Ca²⁺ channel proteins of the voltage gated ion channel (VIC; TC# 1.A.1) family, where a 6 TMS precursor twice duplicated to give 24 TMS proteins (Nelson et al., 1999), this is the first demonstration of such an event occurring from a 2 TMS element and involving three successive intragenic duplication events. Other findings further characterize the greatly expanded superfamily of these secondary active transporters.

Parts of the Introduction are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Methods

PSI-BLAST (Altschul *et al.*, 1997) searches were performed to screen the National Center for Biotechnology Information (NCBI) non-redundant protein database using *Candida albicans* Opt1 (gi# 74582040), *Schizosaccharomyces pombe* Isp4 (gi# 19859374), *Saccharomyces cerevisiae* Opt1 (gi# 731969), *Zea mays* YS1 (gi# 75168533), and *Myxococcus xanthus* EspB (gi# 75421577). The corresponding TinySeq XML format (NCBI) of these proteins was obtained and modified using the script MakeTable5 (Yen *et al.*, 2009) to generate a FASTA file for all of the sequences, and a table containing each protein's abbreviation, description, organismal source, size, gi number, organismal kingdom or phylum, and organismal domain. MakeTable5 (Yen *et al.*, 2009) was also used to remove protein sequences with greater than 90% sequence identity to an included protein. Redundant and partial sequences were removed so that only full length, representative OPT family homologues were further analyzed.

Multiple alignments of homologous proteins and the construction of phylogenetic trees were generated using the CLUSTAL X program (Thompson *et al.*, 1997) followed by the TreeView program (Zhai *et al.*, 2002) with default settings. The WHAT (Zhai and Saier, 2001a) and TMHMM (Käll *et al.*, 2007) programs were used to perform topological analyses on single protein sequences. The AveHAS program (Zhai and Saier, 2001b) with default settings was used to generate average hydropathy, amphipathicity, and similarity plots for multiply aligned sequences. Internal homologous repeat segments in all OPT proteins examined were statistically compared using the IC(Faa2) program (Yen *et al.*, 2009). Segments giving the best comparison scores were further examined using the GAP program with default settings and 500 random shuffles with comparison

scores expressed in standard deviations (S.D.) (Devereux *et al.*, 1984). A value of 10 S.D. corresponds to a probability of 10^{-24} that the observed degree of similarity occurred by chance (Dayhoff *et al.*, 1983). To optimize, the non-aligned segments were removed, numbers of identities were maximized, and numbers of gaps were minimized, maintaining a length of at least 60 residues. The comparison score was then determined again as before. 10 S.D., for a stretch of at least 60 amino acyl residues, corresponding to a typical, average sized protein domain, is deemed sufficient to establish homology (Saier, 1994; Saier *et al.*, 2009; Yen *et al.*, 2009).

The GGSEARCH (http://fasta.bioch.virginia.edu/gasta_www2/fasta_list2.shtml), HMMER (http://hmmer.janelia.org; Eddy, 2008) and SAM (Yen *et al.*, 2009; Wang *et al.*, 2009) programs were subsequently used to provide confirmatory evidence for homology. The halves, quarters and eighths of these homologues, which showed significant sequence similarity using IC/GAP (Table 3) were subsequently used to generate a profile and a database for each program.

The **hmmbuild** program was first used to build an HMM profile for each 8 TMS or 4 TMS segment. This profile was then calibrated using the **hmmcalibrate** program to obtain more accurate e-values. The resulting calibrated profile was then used to search a corresponding 8 TMS or 4 TMS segment database (FASTA formatted sequence file) with the **hmmsearch** program. The resulting output file showed the domain and alignment annotation for each sequence. HMMER commands used were:

hmmbuild <a lignment file> hmmcalibrate <a lignment file> hmmsearch <a lignment file> hmmsearch <a lignment file> file> hmmsearch <a lignment file> hmmsearch <a lignment

The same essential procedures were used for SAM and GGSEARCH. Using the SAM program, the sequence files from the halves and quarters were first trained to build models. The models were subsequently used to search against a database consisting of the corresponding untrained halves and quarters. The SAM commands used were:

buildmodel <model name> -train <training set> -randseed0 hmmscore <output> -I <model file> -db <target sequence file? -sw 2 -

calibrate 1

GGSEARCH of the FASTA package from the University of Virginia (http://fasta.bioch.virginia.edu/fasta_www2?fasta_www.cgi?rm=select&pgm=gnw) was similarly used to compare the 8 TMS halves and the 4 TMS quarters.

Parts of the Methods are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Chapter 1: Phylogenetic Analysis of OPT Family Members

The 325 proteins included in this study are listed alphabetically in Table 2 and according to cluster and position in the phylogenetic tree (Figure 1) in Table 1. The dendogram corresponding to the tree shown in Figure 1 can be viewed in Figure 2. The tree shown in Figure 1 reveals five clusters subdivided as follows. Cluster 1 includes three sub-clusters, 1A - 1C; clusters 2 and 3 have two sub-clusters each, A and B; cluster 4 includes seven sub-clusters labeled, 4A - 4G; cluster 5 has been subdivided into four sub-clusters, 5A - 5D (Figure 1).

The analysis presented in Table 1 reveals the organismal types and size distributions of these proteins according to sub-cluster. Thus, for example, sub-cluster 1A (56 proteins) and 1B (48 proteins) are derived exclusively from fungi, but sub-cluster 1C (27 proteins) is derived exclusively from plants. Sub-cluster 1C is also more distantly related to 1A and 1B than these two latter sub-clusters are to each other (Figure 1). The average sizes of the proteins in sub-clusters 1A - 1C are 825 ± 103 amino acids (aas), 893 ± 41 aas, and 761 ± 105 aas, respectively. These size differences are statistically significant and suggest fundamental differences between these three groups of proteins. The plant proteins on average are 11% smaller than the fungal proteins. This corresponds to the same average size differences observed between plant and fungal homologues of several other ubiquitous families of transporters as reported by Chung *et al.* (2001).

The variations in size within each of these sub-clusters are also of considerable interest. For example, in sub-cluster 1A, the four proteins, Ncr6, Cgl3, Ssc1 and Gze5, cluster tightly together and are roughly 250 aas larger than most of the other homologues. BLAST searches revealed that the extra amino acids in these proteins are at the N-

termini, do not comprise a domain recognized by the Conserved Domain Database (CDD), and although probably homologous, are very diverse in sequence. Another protein of even greater size is Cci3 with 1292 aas. This protein also exhibits a long N-terminal extension that proved to similarly represent a CDD non-recognizable domain. It showed similarity to only a few other fungal proteins. Finally, two moderately large fungal proteins, Cne3 and Uma1, have 961 - 985 aas. The extension again proved to be at the N-terminus, and these sequences showed little similarity to other protein sequences in the NCBI database. When these large homologues were removed from the list of subcluster 1A proteins, the average size proved to be 790 ± 30 aas. Thus, we conclude that the basic size of these proteins is about 790 aas, and all of the larger homologues have extra N-terminal hydrophilic extensions.

The variation in size within sub-cluster 1B is minimal. Several proteins have sizes within the range 900 - 967 aas, but one protein, Yli7, contains 1032 aas. This protein was also examined and proved to have an N-terminal extension that was not homologous to anything in the NCBI Database. When this protein was removed from sub-cluster 1B proteins, the average size was 890 ± 36 aas.

Sub-cluster 1C includes proteins with sizes that vary between 689 – 771 aas with one exception, Osa16. This plant protein shows a long C-terminal hydrophilic extension of about 530 aas. CDD recognized this domain as a member of the pepsin (protease) superfamily. It makes physiological sense that a protease would be fused to a peptide transporter, and thus it appears likely that this fusion is not artifactual. Two programs, TMHMM (Krogh *et al.*, 2001) and HMMTOP (Tusnády and Simon, 2001), were used to determine the orientation of this protein in the membrane. Both programs indicated that

the protease domain is located to the cytoplasmic side of the membrane. In fact these programs showed agreement that most 16 TMS members of the OPT family have both their N- and C-termini on the inside. Excluding Osa16, the average size for all remaining proteins in this sub-cluster is 742 ± 20 aas.

Clusters 2 (11 proteins) and 3 (16 proteins) are close together on the phylogenetic tree, and both derive exclusively from fungi. Both clusters can be subdivided into two sub-clusters where these sub-clusters in cluster 2 are deep branching while those in cluster 3 are not. Cluster 3 has an average size of 788 ± 30 aas, and all proteins occur within the range 746 - 860 aas. Cluster 2 is of even greater size uniformity except for one protein (Ncr4), which is about twice as large (1619 aas) as the others. The OPT family homology region begins at about residue 920 with the expected ~16 TMSs, while the first 900 residues exhibit characteristics of a water-soluble protein. A BLAST search against the NCBI Database of this region retrieved fungal peptidases from the peptidase S41 family. It was therefore clear that Ncr4 is the second OPT family protein identified which has a fused protease domain. However in contrast to Osa16, which had a Cterminal pepsin fusion, Ncr4 has an N-terminal peptidase S41 homologue fusion. Again, the two programs, TMHMM and HMMTOP, were used to estimate the orientation of this protein in the membrane. Surprisingly, and contrary to results of most other members of the OPT family, these two programs predicted that the N-terminus of Ncr4 was on the outside. We therefore examined the distribution of lysine and arginine residues within the transmembrane domain of this protein as well as all members present in the multiple alignment shown in Supplementary Figure S1 which can be viewed on our website (www.biology.ucsd.edu/~msaier/supmat/OPT). In both cases the results clearly

suggested that the N-termini are on the cytoplasmic side of the membrane. The mistake made by the two programs may have resulted from incorrect assignments of four cytoplasmic regions that the programs considered transmembrane. Once again, fusion of a peptidase with a peptide transporter makes excellent physiological sense. As expected based on topological and charge distribution analyses, the cytoplasmic peptidase would hydrolyze the peptides brought in by the transporter in a sequential or coupled process (Saier *et al.*, 2005; Merdanovic *et al.*, 2005; Black and DiRusso, 2007).

Cluster 4 (84 proteins) and Cluster 5 (83 proteins) are the two largest clusters of OPT family members (about half of the total proteins included) as shown in the top half of the tree in Figure 1. While cluster 4 can be conveniently divided into seven subclusters, we have divided cluster 5 into 4 sub-clusters. All cluster 4 proteins are derived from prokaryotes, very few of which are derived from archaea (two in sub-cluster 4A, one in sub-cluster 4B, two in sub-cluster 4F, and one in sub-cluster 4G). Only sub-cluster 4F lacks bacterial homologues. Within each of these sub-clusters there is little size variation; thus the average sizes of sub-clusters 4A- 4D vary between 642 – 665 aas. By contrast, the proteins in sub-clusters 4E – 4G are much smaller (average sub-cluster size of 529 – 553 aas). Not even a single protein within these seven sub-clusters is substantially outside of its sub-cluster size range. The difference in size between these two groups of sub-clusters, about 110 residues, proved to be due to a C-terminal extension present in every one of the former proteins but lacking in the latter as well as the loss of several short sequences within the loop regions between transmembrane domains of the latter. This 110 aa extension approved to be unrelated to anything else in the NCBI nr-protein databank.

Cluster 5 is much more divergent with respect to organismal type and size, but each of the four sub-clusters exhibits a surprising degree of uniformity. Thus, sub-cluster 5A (15 proteins) derives exclusively from δ - and γ -proteobacteria, and these proteins exhibit an average size of 589 ± 29 aas; no protein is strikingly outside of this range. Sub-cluster 5B (27 proteins) derives from fungi with one exception, a protein from the slime mold *Dictyostelium discoideum*. The average size is 742 ± 45 aas, and two *Aspergillus* proteins are substantially larger than the others (Afu3, 843 aas and Aor6, 851 aas). Examination of the multiple alignment revealed that these proteins have neither N-or C-terminal extensions. Instead, they both have internal insertions, one near their N-termini that immediately proceed TMS 1. This insert is found only in these two proteins. The other insert is near the C-termini of these proteins, immediately preceding the last TMS. Homologous sequences are found in a few other proteins, mostly from species of *Aspergillus*. Neither of these 40 residue inserts shows appreciable sequence similarity with other proteins in the NCBI Protein Database.

Sub-cluster 5C (4 proteins) derives from three β -proteobacteria and one δ -proteobacterium. The average size is 606 ± 20 aas, similar to that of sub-cluster 5A, also derived from proteobacteria. These proteins are much shorter than the eukaryotic proteins of sub-clusters 5B and 5D. Sub-cluster 5D (37 proteins) is derived exclusively from plants and has an average size of 697 ± 40 aas. Only one protein is substantially larger than the others; this protein is Osa13 (882 aas). It has an approximately 150 residue C-terminal hydrophilic extension found in no other member of this sub-cluster. This region of the protein showed a low degree of sequence similarity with chloride

transporters of the ClC family (TC# 2.A.49). However the significance of this observation is questionable.

One member of each sub-cluster was used as the query sequence to search TCDB using TC-BLAST. All sub-clusters in clusters 1-3 (lower half of the tree) proved to bring up peptide transporters, while all of the sub-clusters from clusters 4 and 5 brought up the iron-complex transporters. The phylogenetic segregation between these two functional types is so considerable that one must conclude that in general, function correlates remarkably well with phylogeny.

Parts of Chapter 1 are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Chapter 2: Orthologous Relationships within Sub-clusters of the OPT Family Tree

The phylogenetic tree for the 16S/18S rRNAs is shown in Figure 3. The bacteria appear at the top of this tree, the archaea in the small cluster on the right hand side, and the eukaryotes at the bottom. Every genus included in our study of OPT family members is represented in this tree with the exceptions of *Acidobacteria*, *Ashbya*, *Cryptococcus*, and *Thlaspi*. The tree shows that all of the γ - and β -proteobacteria cluster most closely together followed by the α -, δ -, and ϵ -proteobacteria on the upper-left hand side of the tree. Surprisingly, in this tree the ϵ -proteobacteria cluster loosely with the bacteriodetes, distantly from the other proteobacteria. The cluster on the upper-right hand side of the tree includes a single member of the acidobacteria, a single cluster of actinobacterial rRNAs and two distinct clusters of firmicutes. The eukaryotic branch of the tree shows the slime mold *Dictyostelium* closer to the center of the tree, with the fungal and plant RNAs clustering more closely to each other but much more distinctly from the slime mold at the bottom of the tree.

Comparing the protein tree (Figure 1) with the RNA tree (Figure 3) we see that in some, but not other cases, orthologous relationships are difficult to establish. This is true for the large Cluster 1. For example, sub-cluster 1C can be sub-divided in five sub-sub-clusters, all but one of which contain paralogues from a single organism. In the largest sub-sub-cluster, for example, we find five paralogues from *Vitis vinifera*, two from *Oryza sativa* of the Indica group, and two from *Arabidopsis thaliana*. The only sub-sub-cluster that lacks paralogues is the uppermost sub-sub-cluster with four proteins from four different organisms. Based on the comparison between Figures 1 and 3, only in this sub-sub-cluster are the results consistent with orthology.

In the adjacent sub-sub-cluster, where we find three proteins, one from rice (*Oryza*) and two from thale cress (*Arabidopsis*), it appears that the two thale cress proteins arose by gene duplication after these two organisms diverged from each other. The same situation is observed for the next sub-sub-cluster where three *Arabidopsis* proteins cluster tightly together with a single *V. vinifera* being the outlier. We interpret these results to mean that after *Arabidopsis* diverged from *Vitis*, two gene duplication events in the former organism gave rise to the three paralogues, Ath9, Ath16, and Ath17. Similar observations were made for sub-clusters 1A and 1B.

Cluster 2 shows relationships which suggest orthology. Thus, in both trees, we find the proteins and rRNAs from *Neosartorya*, *Aspergillus* and *Sclerotinia* clustering together, *Candida*, *Lodderomyces* and *Pichia* clustering together, and *Neurospora* and *Botryotinia* clustering together. Even within each of these three groups, the phylogenetic order in both trees is the same. We conclude that Cluster 2 probably represents a collection of pure orthologs with no evidence for paralogues or horizontal gene transfer. This observation suggests that these proteins all serve a single unified function in all of these organisms.

In contrast to Cluster 2, Cluster 3 contains a number of non-adjacent paralogues, and also shows clear non-orthologous relationships. The obvious paralogues include two proteins each from *Gibberella zeae* and *Ustilago maydis* in two different sub-clusters that are shared by this pair of paralogues from these two organisms. Additionally, based on the comparison between Figures 1 and 3, Uma4 from *Ustilago maydis* does not show orthologous relationships with the other members of this sub-cluster. Furthermore, the two *Neurospora crassa* proteins, Ncr5 and Ncr7, are two paralogues within the same sub-

sub-cluster. On the other hand, the three *Aspergillus* proteins and the one from *Neosartorya fisceri* form a sub-sub-cluster on the protein tree as well as the RNA tree, and the same is true for the two *Schizophyllum* and *Laccaria* proteins and RNAs which form a distinct sub-sub-cluster in both trees. The relationships of all of these proteins are similar to the corresponding relationships in the rRNA tree and are therefore consistent with orthology.

The prokaryotic proteins were similarly analyzed. Starting with sub-cluster 4A, we find seven distinct sub-sub-clusters. Progressing in the clockwise direction, sub-sub-cluster 1 includes proteins from α - and β -proteobacteria as well as actinobacteria. As a single β -proteobacterial protein is flanked by α -proteobacterial proteins, it is possible that this one β -proteobacterial protein (Neu1) was obtained by horizontal transfer. However, the α -proteobacterial proteins do not show orthologous relationships. The actinobacterial proteins show relationships consistent with orthology.

Sub-sub-cluster 2 is derived exclusively from *Campylobacter* species. Sub-sub-cluster 3 contains β -proteobacterial proteins with a single outlier (Pae1) from a γ -proteobacterium. The members of this small sub-sub-cluster could be orthologous. However, in sub-sub-clusters 4, 6, and 7 orthology is not possible. For example, in sub-sub-cluster 4, *Haemophilus* and *Actinobacillus* proteins are interspersed, while in sub-sub-cluster 7, γ -proteobacterial and archaeal proteins are interspersed. It would appear that the precursor of the two archaeal proteins were obtained from γ -proteobacteria via horizontal transfer, but this remains speculative.

Analyses of sub-clusters 4B through 4G allowed us to come to similar conclusions. Thus for example, sub-cluster 4B contains proteins from highly divergent

organisms including δ -proteobacteria, acidobacteria, firmicutes, and archaea; sub-cluster 4C includes proteins from two different bacterial phyla, the bacteriodetes and the acidobacteria; sub-cluster 4E includes just two proteins from two different bacterial phyla; sub-cluster 4G contains proteins from firmicutes, β - and γ -proteobacteria, and an archaeon. It seems clear that in all of these sub-clusters, horizontal gene transfer was rampant during the evolution of these proteins.

The four Cluster 5 sub-clusters (A – D) were similarly analyzed. Sub-cluster 5A, derived from δ - and γ -proteobacteria, includes paralogues with little indication of orthology. Sub-cluster 5B derives from fungi with the exception of one slime mold protein. It also exhibits relationships suggestive of horizontal gene transfer (especially the slime mold protein, Ddi1, which probably derived from a fungus) as well as distant paralogues from three different genuses. Even the small sub-cluster 5C shows signs of the existence of horizontal gene transfer since the δ -proteobacterial protein (Sau3) is unexpectedly closely related to the β -proteobacterial proteins. Finally, sub-cluster 5D, shows many paralogous proteins (*e.g.*, at least 12 probable *Oryza sativa* (Japonica group) paralogues and at least seven *A. thaliana* paralogues). In this case, it is difficult to know if horizontal gene transfer has occurred, as all of these proteins could have arisen by vertical transmission from multiple precursor paralogues in the primordial plant.

Parts of Chapter 2 are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Chapter 3: Topological Analyses of OPT Family Proteins

Figure 4 shows the average hydropathy (top) and average similarity (bottom) plots for all 325 members of the OPT family included in this study. This plot reveals 16 peaks of hydropathy that in general correspond to peaks of similarity. The first four TMSs (labeled 1-4) cluster loosely together. TMSs 4 and 5 are separated by a substantial hydrophilic loop, but again, the next four TMSs (5-8) cluster together. Between TMSs 8 and 9 is an even larger hydrophilic loop, but the remaining eight TMSs cluster tightly together. It is interesting to note that peak 3 and also peak 11 appear to divide into two small peaks, possibly due to a problem of misalignment. In fact, there appears to be a gap within the region designated as peak 3 and a smaller gap within the region designated as peak 11. Based on the appearance of this plot it seemed possible that TMSs 1 - 8 are repeated in TMSs 9 - 16. Further, the clustering pattern suggested the possibility that these proteins might have arisen from a 4 TMS precursor peptide that duplicated twice to give the present day 16 TMS proteins. In this regard, it should be noted that in all four apparent quadrants, the first two TMSs (1-2, 5-6, 9-10, and 13-14) are always close together, while the subsequent two TMSs in each quadrant are separated by greater distances. The possibility that these 16 TMS proteins arose by a quadruplication event will be demonstrated below. It is worthy of note, that following TMS 16, is a poorly conserved region that exhibits moderate hydrophobicity.

When the individual sub-clusters shown in Figure 1 were analyzed for average hydropathy and average similarity as shown in Figure 4 for all members of the family, we found that almost all sub-clusters exhibited the typical 16 TMS topology. However, the proteins within sub-clusters 4A-4D appeared to have a seventeenth transmembrane

segment that was not part of the C-terminal 4 TMS repeat. Also, in these four subclusters, TMS 13 showed only moderate hydrophobicity as revealed by the AveHAS program. The origin of TMS 17 in these proteins is unknown but could have arisen as a result of a gene fusion event. The long N-terminal and C-terminal hydrophilic extensions have been discussed above and proved to be homologues of functionally recognizable proteases in only two cases.

Parts of Chapter 3 are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Chapter 4: Establishment of Internal Repeats in OPT Family Proteins

As noted above, most members of the OPT family contain 16 putative TMSs, although a few appear to have 17 TMSs, the extra one being at the C-terminus of each of these proteins. In order to confirm TMS assignment and establish the evolutionary origins of these proteins, we conducted analyses of potential internal repeats. Although initially analyzed assuming different numbers of TMSs per repeat unit, we were able to show with relative ease that these proteins include an 8 TMS duplication. Thus, when using the IC/GAP programs to compare the first halves of these proteins with the second halves, comparison scores of up to 12.6 standard deviations (S.D.) were obtained (see Table 3 and Figure 5). This value is substantially greater than required to establish homology (Saier, 1994; Yen *et al.*, 2009; Wang *et al.*, 2009; Matias *et al.*, 2010).

We next examined the possibility that the 8 TMS halves themselves arose by an earlier intragenic duplication event from a 4 TMS precursor. The results from these analyses are also presented in Table 3, and the alignment upon which the best comparison score was based is shown in Figure 6. In Table 3, we summarize the results obtained using the IC and GAP programs with 500 random shuffles and default settings. All four quarters of these proteins were compared with each other. Only the top two scores are reported, and these were averaged. For all comparisons, values in excess of 10 S.D. were obtained, clearly indicating homology. However the best scores were obtained when A vs C and B vs D were compared (12.2 S.D. and 13.2 S.D., respectively). The fact that higher values were obtained for these two comparisons than for any of the others is strong evidence that these two duplication events, giving rise to the 16 TMS proteins, were separated by a substantial period of evolutionary time. Thus, we suggest that the

primordial 4 TMS-encoding genetic element duplicated once to give the 8 TMS precursor, and then later, the second duplication occurred giving rise to the 16 TMS proteins. Alternatively, segments A and C may share a structure/function that is substantially different from the structure/function shared by segments B and D (see Discussion section).

As the final step we examined the possibility that within each of the 4 TMS quadrants of these proteins we could detect two 2 TMS repeat sequences. Much to our surprise and delight, this possibility could be demonstrated. As shown in Table 3 and Figure 7, comparing the first 2 TMSs with the second 2 TMSs of the first of these four 4 TMS repeats gave a maximal value of 8.9 S.D., insufficient to establish homology. However, when comparing the two 2 TMS segments of the second of these four repeats, we were able to get comparison scores in excess of 10 S.D., thus establishing homology. In this case the alignment giving this value included all of TMS 5 compared to TMS 7. When the same was done with the third of these four repeats, a maximal value of 8.6 S.D. was obtained. The same procedure with the fourth of these four repeats did not give values above 7 S.D. Thus, applying the superfamily principle, the values obtained clearly indicate that these proteins arose from an initial 2 TMS precursor. We therefore conclude that members of the OPT superfamily arose in three steps; duplication of 2 TMSs to give 4, duplication of 4 TMSs to give 8, and the last duplication to give 16 TMSs. The addition of a seventeenth TMS to a small fraction of these proteins presumably occurred as a result of a late gene fusion event in just one phylogenetic cluster of these proteins.

Parts of Chapter 4 are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Chapter 5: Use and Evaluation of Programs to Detect Similarity and Establish Homology

To confirm the results obtained using the IC/GAP programs, three other programs capable of identifying sequence similarity between repeat segments were used. These programs were GGSEARCH, HMMER, and SAM (Table 4). All three programs substantiated the conclusions obtained with IC/GAP. For example with GGSEARCH, when the two halves were compared, a value of 1.7 e⁻⁸ was obtained. The best value resulting from the use of the HMMER program was 4 e⁻⁴. When SAM was used, the best value was 4 e⁻³. All of these values confirm our conclusion of homology.

When the four quarters of the OPT family proteins were compared, again the best values were usually obtained when segments A were compared with segments C, and segments B were compared with segments D. Thus, when using GGSEARCH, the values obtained for these two comparisons were 8.6 e⁻⁶ and 3.9 e⁻⁸. When using HMMER, the best values were 0.03 and 0.006. With SAM, the best values were 0.002 and 0.001, respectively (Table 4). As revealed by the data in Table 4, only in two instances were values obtained in the other comparisons comparable to these. We conclude on the basis of all of these results that 1) the four 4 TMS quarters of OPT family proteins are all homologous and therefore derive from a common origin, 2) the first and third 4 TMS segments are more similar to each other than they are to the second and fourth TMS segments, and the second and fourth TMS segments are more similar to each other than they are to the first and third segments. On this basis, we suggest that they probably arose by two distinct intergenic duplication events separated by a substantial period of evolutionary time. The possibility of greater restriction between A and C, and B and D

due to common structure and function cannot be eliminated, but in an analogous situation where a 6 TMS voltage-gated ion channel has four 6 TMS repeats, this last possibility seemed unlikely (Nelson *et al.*, 1999).

Three additional superfamilies were examined with the four programs used above in order to evaluate their relative abilities to detect distant phylogenetic relationships.

The superfamilies include 1) the CRAC/CDF superfamily (Matias *et al.*, 2010), 2) the Drug/Metabolite Transporter (DMT) superfamily (Tran and Saier, 2004; Jack *et al.*, 2001), and 3) the Bile acid/Arsenite/Riboflavin Transporter (BART) superfamily (Mansour *et al.*, 2007). The data are presented in Table 5.

The first two entries in Table 5 present comparisons between the CDF family and the CRAC (Orai) family. The first entry compares the complete sequences of both proteins, while the second entry compares TMSs 3-4 in the CDF protein with TMSs 1-2 in the Orai homologue. These are the regions showing the greatest sequence similarity. These comparisons using the IC/GAP program set gave 14 S.D., a value far in excess of what is required to establish homology. GGSEARCH also gave values sufficient to strongly suggest homology (4.9 e⁻³ and 5.4 e⁻⁵) for the full-length sequence, and 1.6 e⁻¹⁸ and 9.4 e⁻⁵ for the CDF TMSs 3-4 compared with Orai TMSs 1-2. According to the HMMER website, e-values smaller than 0.1 are significant. According to this criterion, one value obtained with this program was borderline (0.09). Finally, SAM gave on value (0.02) that was suggestive of homology.

The DMT superfamily was next examined (Table 5). When two members of a single family within this superfamily were compared, all four programs predicted homology. The same was true for members of two distinct families within this

superfamily (SLC35A1 with PfCRT) and the degrees of sensitivity detected by the last three programs were GGSEARCH (G) > SAM (S) > HMMER (H).

For the BART superfamily, three different comparisons were run: the first between two families of known transport function, and the second two of unknown function. In the first comparison the sensitivities of the three programs was G>H>S. In the second and third comparisons, the order was G>H>S where S did not give significant e-values.

When considering all distantly related comparisons (Table 5), five showed G>H>S, two showed G>S>H, and one showed H>S>G. Thus, while we consider IC/GAP is the gold standard for establishing homology, we suggest that of the three remaining programs, for the purpose of detecting sequence similarities, GGSEARCH is better than HMMER, which is better than SAM (the most-time consuming program to use). However since SAM was better than HMMER in two cases, and HMMER was better than GGSEARCH in one case, we conclude that the use of all three of these programs is superior to the use of any one or two of them when time and effort are not limiting. We recommend IC/GAP and GGSEARCH as the two most sensitive programs for detection of significant sequence similarity between distantly related homologues. It should be noted that if one program detects significant sequence similarity and any number of programs do not, the first program, giving positive results, is to be trusted over those that give negative results because only the first program has correctly aligned the sequences being compared.

Parts of Chapter 5 are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

Discussion

In this paper, we have described the OPT family of peptide and iron-siderophore uptake transporters and have defined the evolutionary pathway by which these proteins arose. This pathway is illustrated in Figure 8. A genetic element encoding a 2 TMS precursor duplicated to give 4 TMSs, this duplicated again to give 8 TMSs, and this also duplicated to give the final 16 TMS topology. In few instances has it been possible to trace back the evolutionary history as far as we have done for the OPT family (Saier, 2003). Furthermore, in no other instance has this particular pathway been demonstrated for any other family of transport proteins (Saier, 2003 and unpublished observations).

We could demonstrate greater similarities between TMSs 1-4 and TMSs 9-12, as well as TMSs 5-8 and TMSs 13-16 than for other quadrant comparisons, suggesting that there was a reasonable period of evolutionary time between these two last duplication events. However the fact that similar maximal values were obtained for the 8 TMS halves, the 4 TMS quarters, and the 2 TMS eighths suggests that all three of these duplication events happened in a relatively short period of evolutionary time. These two apparent inconsistencies could be resolved if the first and third quadrants serve a common structure/function that differs from that of quadrants 2 and 4. This may prove to be the explanation for the observed relative degrees of sequence similarity.

The same has been suggested for members of the Mitochondrial Carrier Family which underwent triplication of a primordial 2 TMS encoding genetic element (Kuan and Saier 1993a, b). This family of proteins appears to have undergone rapid intragenic and extragenic duplication events giving rise not only to the 6 TMS porters but also to the main functional types or subfamilies within a short period of time (Kuan and Saier

1993a). Interestingly, in the mitochondrial carriers, the third thirds of these proteins diverged in sequence more than the first two thirds (Kuan and Saier 1993a). The explanation for this observation is not yet clear.

Many transporters have been shown to arise from a 2 TMS precursor, but in no case has it been possible to demonstrate three sequential duplication events (Sawhney *et al.*, 2010). Other families in which a 2 TMS element duplicated to give 4 TMSs include the Voltage-gated Ion Channel (VIC; TC# 1.A.1) Family, the C-subunits of F-type ATPases (F-ATPase; TC# 3.A.2) which both duplicated and triplicated, and the YiaAB Family (TC# 9.B.44) (Saier, 2003). Other examples are reported in Sawhney *et al.* (2010).

A surprising observation was that all members of the OPT family have either 16 or 17 TMSs. The vast majority have 16 TMSs, while a smaller fraction (sub-clusters 4A – 4D in the phylogenetic tree shown in Figure 1) have 17 TMSs. In fact no 17 TMS protein was found outside of sub-clusters 4A – 4D and only 17 TMS proteins were found in these four sub-clusters. The extra TMS at the C-terminus of these proteins most probably arose only once during the evolution of this family. The only additional variation resulted from the fusion of these integral membrane proteins with soluble domains, two of which could be recognized on the basis of homology searches. In these two cases the fused domains proved to correspond to two different families of peptidases. Since the transporters were predicted to function in peptide uptake, and since the peptidase domains were predicted to be localized to the cytoplasmic side of the membrane, the fusion of these two catalytic proteins made physiological sense. The peptidase domain probably hydrolyzes the peptide upon entry into the cell, possibly in a

tightly or loosely coupled process. If tightly coupled, this could be a novel example of group translocation where chemical modification of the substrate is coupled to its transporter (Herbert *et al.*, 2003; Hirsch *et al.*, 1998; Merdanovic *et al.*, 2005; Saier *et al.*, 2005).

Uniformity of topology is found in some families while others show tremendous variation. For example all recognized proteins in the Mitochondrial Carriers Family (TC# 2.A.29) have 6 TMSs, and no exception has yet been reported (Kuan and Saier 1993a, and unpublished results). Another example is the largest superfamily of secondary carriers, the Major Facilitator Superfamily (TC# 2.A.1). All recognized members of this superfamily have either 12 or 14 TMSs, where the extra 2 TMSs in the 14 TMS proteins are present in the center between the two 6 TMS repeat units, and they occur only in three of the 70 currently recognized MFS families. This situation is to be contrasted with families that show tremendous topological variations. These include the integral membrane cytochrome c biogenesis proteins of the Heme Handling Protein Family (TC# 9.B.14) (Lee et al., 2007) and the SdpI Family of receptor/signal transduction proteins (TC# 9.A.32) (Povolotsky et al., 2010). In both of these cases the family includes proteins having a wide variety of topological types with numbers of TMSs ranging anywhere from three to twelve. Further, they can have segments inverted in some of the proteins relative to other members of the same family. In the SdpI family, this is understood because the different 3 TMS segments within these proteins probably serve distinct functions (Povolotsky *et al.*, 2010).

OPT family members were found in both eukaryotes and prokaryotes. The vast majority of the eukaryotic proteins were derived from fungi (sub-clusters 1A, 1B, and 5B

as well as clusters 2 and 3) and plants (sub-clusters 1C and 5D). The only exception is a single slime mold homolog found in sub-cluster 5B, a cluster otherwise entirely derived from fungi. We hypothesize that this one homolog from *Dictyostelium discoideum* was acquired by horizontal transfer from a fungus, a suggestion that is not surprising since slime molds eat other microorganisms (Eichinger *et al.*, 2005). Otherwise, we have obtained no evidence for horizontal transfer between fungi and plants. In view of the fact that homologs of these proteins are found in many bacterial and archaeal phyla, it is surprising that these proteins are not found within the animal kingdom or any of the unicellular eukaryotes except for slime molds.

Prokaryotic homologs of the OPT family are found in sub-clusters 4A - 4G as well as 5A and 5C. In contrast to the situation with eukaryotes, apparent horizontal transfer between prokaryotic phyla has been rampant. For example, in sub-cluster 4A, proteins are derived from four of the five orders of proteobacteria, the only exception being the δ -proteobacteria. However this sub-cluster also contains proteins from actinobacteria and even euryarchaeota. Similarly sub-cluster 4B includes proteins from δ -proteobacteria, firmicutes, acidobacteria and euryarchaeota. Sub-cluster 4C has representation of proteins only from bacteriodetes and acidobacteria. Sub-cluster 4D is one of the few "pure" prokaryotic sub-clusters where all of the proteins derive from firmicutes. Sub-cluster 4E includes just two proteins, one from firmicutes and one from actinobacteria. Sub-cluster 4F similarly has two proteins, but they are derived from euryarchaeota. Sub-cluster 4G, a small sub-cluster of seven proteins, is exceptionally diverse having members from firmicutes, β - and γ -proteobacteria, and euryarchaeota.

cluster 5C has representation only from β - and δ -proteobacteria. These observations can be interpreted to suggest that horizontal transfer has occurred in all but two of the prokaryotic sub-clusters identified in this study.

In summary, we have characterized the large OPT family of peptide and iron-siderophore uptake porters. We have shown that, based on functionally characterized members of this family, all of the iron-siderophore transporters (Clusters 4 and 5) segregate from all of the peptide transporters (Clusters 1 – 3). Assuming this functional assignment to be correct, then all peptide transporters of this family are found in a restricted group of eukaryotes, the plants and fungi. By contrast, the iron-siderophore members of this family are found in many phyla of prokaryotes as well as the fungi, plants, and a single slime mold. This distribution is consistent with the suggestion that the primordial transporters of this family were prokaryotic iron-siderophore transporters, that these were transmitted to eukarotes, and that the peptide transporters arose just once in the eukaryotic domain from the former functional type.

While this suggestion appears most reasonable, we still have no certain explanation as to why members of this family appear to be lacking in the animal kingdom as well as almost all eukaryotic protists. If further genome sequencing reveals the presence of these homologs in other types of eukaryotes, this will raise the question of whether these resulted from horizontal gene transfer from fungi, plants or slime molds. This may be an important question, since in this study, we have found very little evidence for horizontal transfer between eukaryotic phyla. Future functional analyses and further sequencing efforts are likely to provide eventual answers to these questions. We hope

that the analyses reported here will provide useful guides for molecular biological and bioinformatic analyses of this interesting family of transporters.

Parts of the Discussion are being prepared for publication. The Thesis author will be the primary investigator and author of this paper.

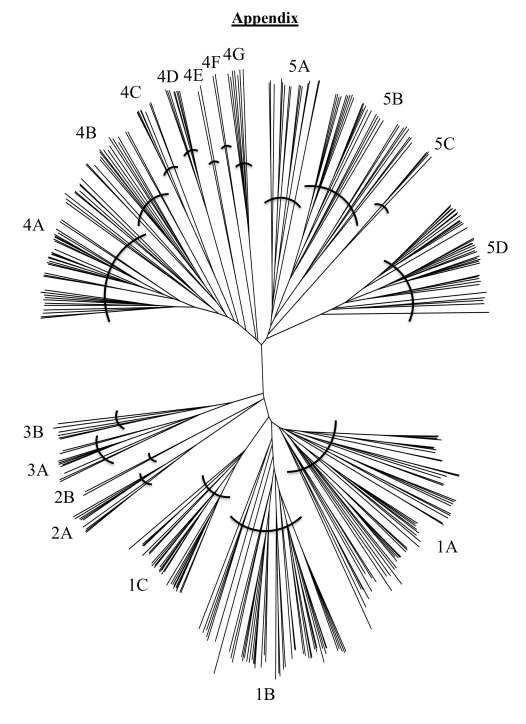


Figure 1. Phylogenetic tree of 325 OPT superfamily proteins based on the ClustalX multiple alignment show in Figure S1, and drawn using the FigTree program. Clusters 1-5 are labeled with their respective sub-clusters. Sub-clusters 1A-3B are putative peptide transporters while sub-clusters 4A-5D are likely to be iron-siderophore transporters. Protein abbreviations are presented in Table 1 together with the characteristics of these proteins.

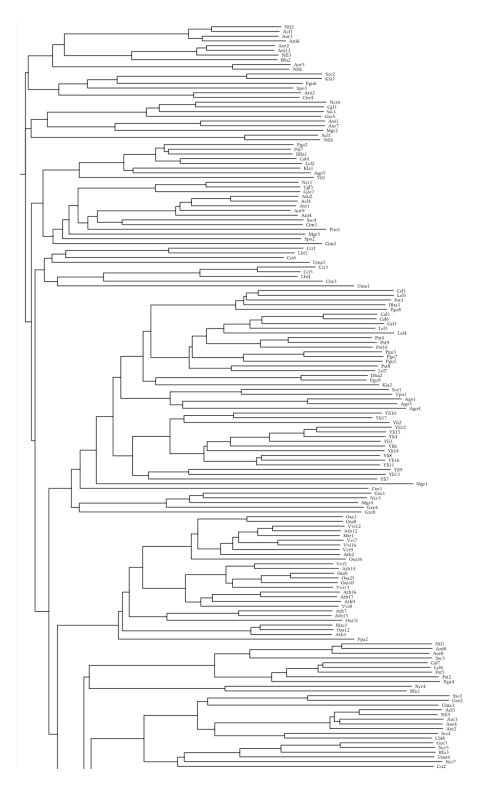


Figure 2. Dendogram of all 325 OPT family proteins included in this study corresponding to the phylogenetic tree shown in Figure 1.

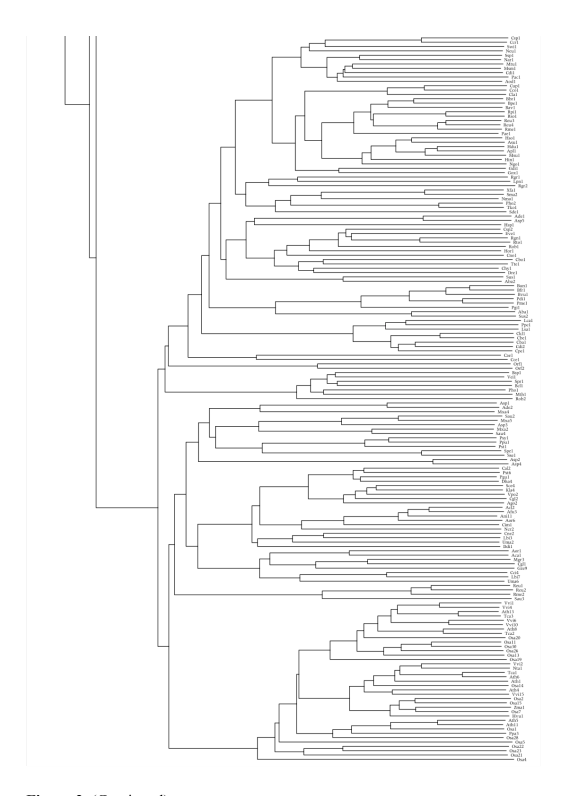


Figure 2. (Continued)

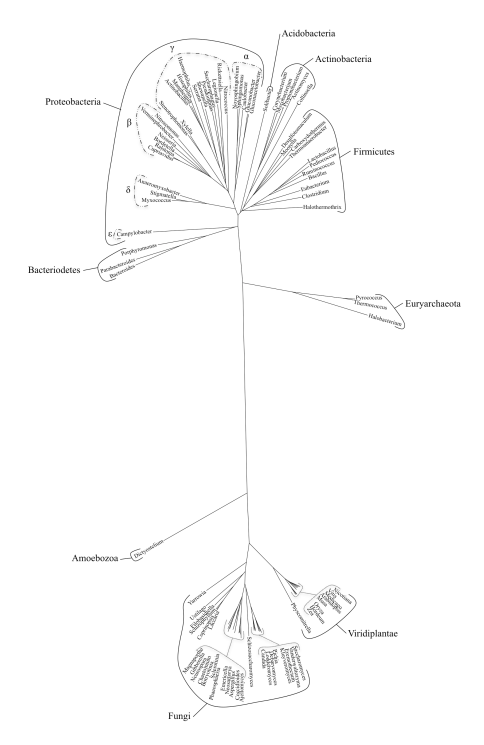


Figure 3. Phylogenetic tree of 16S/18S rRNAs from all genuses represented in this study with the exceptions of *Acidobacteria*, *Ashbya*, *Cryptococcus*, and *Thlaspi*. All bacterial rRNAs appear at the top of the tree; the eukaryotic rRNAs are at the bottom of the tree, and the three archaeal homologues are positioned on the central branch on the right hand side of the tree. The phylum/kingdom is indicated for each of the clusters while the genus is shown at the end of each branch.

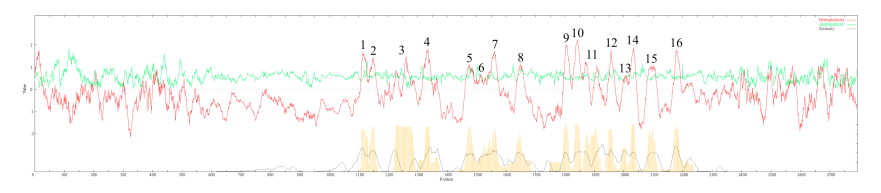


Figure 4. Average hydropathy, amphipathicity, and similarity plots for the 325 OPT superfamily proteins included in this study. The majority of OPT proteins contain 16 TMSs which correspond to the 16 conserved peaks labeled 1 - 16. The central portion of this plot includes all 16 peaks of hydrophobicity which comprise the transporter domain. Functional assignments for the N-terminal and C-terminal hydrophilic domains are discussed in the text.

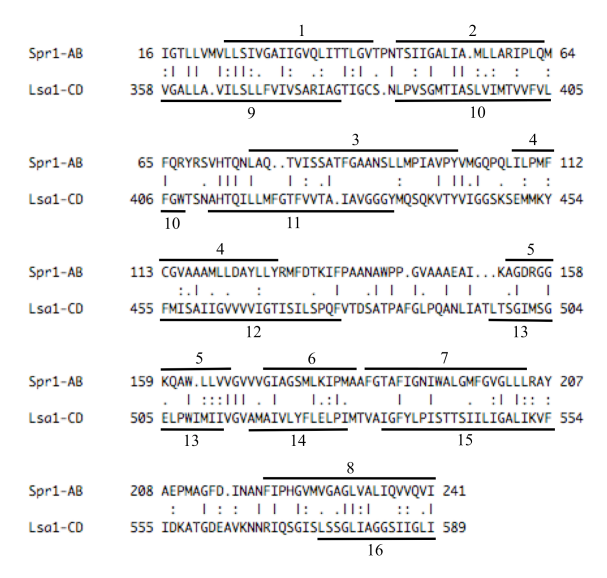


Figure 5. Alignment of OPT TMSs 1-8 of Spr1 (*Serratia proteamaculans*, GI# 157369266) with OPT TMSs 9-16 of Lsa1 (*Lactobacillus sakei*, GI# 81427933). The IC program was used to identify the two internal segments exhibiting the greatest statistical similarity. The GAP program was used to generate the alignment with default settings and 500 random shuffles. Numbers at the beginning and end of each line indicate the residue numbers in the proteins. The | represents an identity, the : represents close similarity, and the . represents a more distant similarity. This convention of presentation is used in Figures 6 and 7. In all three figures, positions of the TMSs were predicted using the TMHMM program. This alignment gave a comparison score of 12.6.

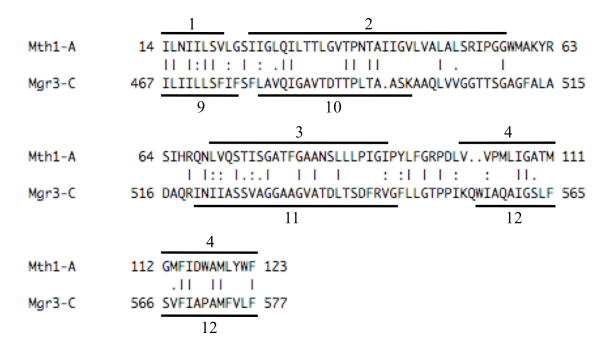


Figure 6. Alignment of OPT TMSs 1-4 of Mth1 (*Moorella thermoacetica*, GI# 83589078) with OPT TMSs 9-12 of Mgr3 (*Magnaporthe grisea*, GI# 39955178). This alignment gave a comparison score of 11.9.

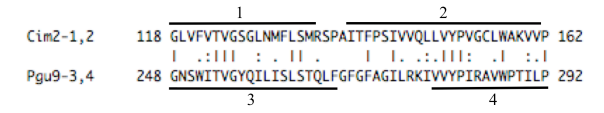


Figure 7. Alignment of OPT TMSs 1-2 of Cim2 (*Coccidioides immitis*, GI# 119190959) with OPT TMSs 3-4 of Pgu9 (*Pichia guilliermondii*, GI# 146422868). This alignment gave a comparison score of 8.7.

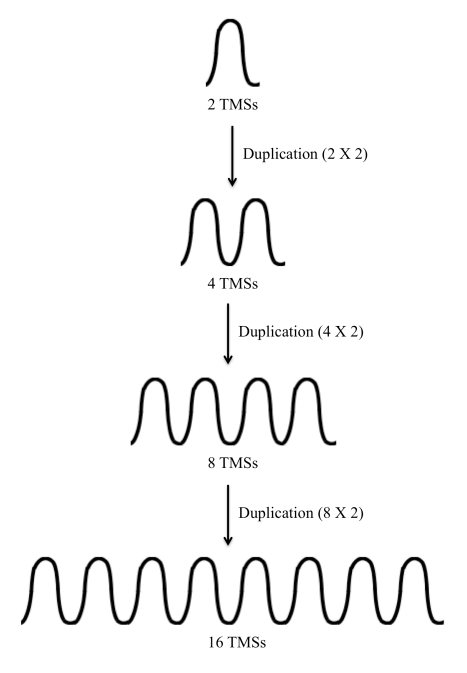


Figure 8. Proposed pathway for the evolutionary appearance of present-day OPT family proteins. Evidence is presented that the ultimate precursor of the 16 (and sometimes 17) TMS proteins was a 2 TMS hairpin structure (top). This then duplicated three times: first to give the 4 TMS intermediate; second to give the 8 TMS intermediate, and last to give the present day 16 TMS proteins. Evidence is presented that the duplication of 4 TMSs to give 8 TMSs occurred substantially before the duplication of 8 TMSs that gave rise to the 16 TMS permeases. In the 17 TMS proteins, the extra TMS is at the C-termini of these homologues.

Table 1. OPT protein sequences included in this study. Proteins are listed based on position in the phylogenetic tree (Figure 1; clockwise direction) according to cluster and sub-cluster. The average sizes of the members of each sub-cluster are presented below the list of these proteins.

Sub-Cluster 1A (56 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Nfi2	Neosartorya fischeri NRRL 181	119471104	Fungi	Eukaryota	757
Acl5	Aspergillus clavatus NRRL 1	121709515	Fungi	Eukaryota	761
Aor3	Aspergillus oryzae	83768538	Fungi	Eukaryota	751
Ani6	Aspergillus niger CBS 513.88	145241488	Fungi	Eukaryota	859
Aor2	Aspergillus oryzae	83768389	Fungi	Eukaryota	765
Ani12	Aspergillus niger CBS 513.88	145251507	Fungi	Eukaryota	771
Nfi3	Neosartorya fischeri NRRL 181	119471211	Fungi	Eukaryota	770
Bfu2	Botryotinia fuckeliana B05.10	154313655	Fungi	Eukaryota	779
Aor5	Aspergillus oryzae	83768732	Fungi	Eukaryota	770
Nfi6	Neosartorya fischeri NRRL 181	119491377	Fungi	Eukaryota	768
Sce2	Saccharomyces cerevisiae YJM789	151943695	Fungi	Eukaryota	799
Kla3	Kluyveromyces lactis NRRL Y-1140	50307929	Fungi	Eukaryota	793
Pgu6	Pichia guilliermondii ATCC 6260	146419361	Fungi	Eukaryota	754
Spo3	Schizosaccharomyces pombe	63054465	Fungi	Eukaryota	851
Ani2	Aspergillus niger CBS 513.88	67540564	Fungi	Eukaryota	778
Cne4	Cryptococcus neoformans var. neoformans B-3501A	134113154	Fungi	Eukaryota	797
Ncr6	Neurospora crassa OR74A	164422675	Fungi	Eukaryota	1094
Cgl3	Chaetomium globosum CBS 148.51	116193201	Fungi	Eukaryota	1027
Ssc1	Sclerotinia sclerotiorum 1980	156039822	Fungi	Eukaryota	1055
Gze5	Gibberella zeae PH-1	46125699	Fungi	Eukaryota	1060
Ani1	Aspergillus niger CBS 513.88	67516837	Fungi	Eukaryota	792
Aor7	Aspergillus oryzae	83770544	Fungi	Eukaryota	778
Mgr2	Magnaporthe grisea 70-15	39944474	Fungi	Eukaryota	783
Acl1	Aspergillus clavatus NRRL 1	121699197	Fungi	Eukaryota	788
Nfi4	Neosartorya fischeri NRRL 181	119477757	Fungi	Eukaryota	772
Pgu2	Pichia guilliermondii ATCC 6260	146416527	Fungi	Eukaryota	784
Pst7	Pichia stipitis CBS 6054	150864787	Fungi	Eukaryota	782
Dha1	Debaryomyces hansenii CBS767	50413511	Fungi	Eukaryota	776
Cal4	Candida albicans	68485275	Fungi	Eukaryota	783

Lel2	Lodderomyces elongisporus NRRL YB-4239	149235877	Fungi	Eukaryota	804
Kla1	Kluyveromyces lactis NRRL Y-1140	50307527	Fungi	Eukaryota	794
Ago5	Ashbya gossypii ATCC 10895	45201069	Fungi	Eukaryota	796
Yli1	Yarrowia lipolytica CLIB122	50542874	Fungi	Eukaryota	836
Ncr1	Neurospora crassa OR74A	9368956	Fungi	Eukaryota	801
Cgl5	Chaetomium globosum CBS 148.51	116198757	Fungi	Eukaryota	871
Gze7	Gibberella zeae PH-1	46134295	Fungi	Eukaryota	799
Afu2	Aspergillus fumigatus Af293	70999364	Fungi	Eukaryota	792
Acl4	Aspergillus clavatus NRRL 1	121705906	Fungi	Eukaryota	793
Ate1	Aspergillus terreus NIH2624	115397517	Fungi	Eukaryota	788
Aor9	Aspergillus oryzae	83775779	Fungi	Eukaryota	768
Ani4	Aspergillus nidulans FGSC A4	67901220	Fungi	Eukaryota	794
Ssc4	Sclerotinia sclerotiorum 1980	156049297	Fungi	Eukaryota	827
Cim3	Coccidioides immitis RS	119194107	Fungi	Eukaryota	812
Pno1	Phaeosphaeria nodorum SN15	160705030	Fungi	Eukaryota	845
Mgr5	Magnaporthe grisea 70-15	145614314	Fungi	Eukaryota	849
Spo2	Schizosaccharomyces pombe	19115899	Fungi	Eukaryota	785
Cim2	Coccidioides immitis RS	119190959	Fungi	Eukaryota	810
Cci1	Coprinopsis cinerea okayama7#130	116500528	Fungi	Eukaryota	757
Lbi5	Laccaria bicolor S238N-H82	164641826	Fungi	Eukaryota	730
Cci6	Coprinopsis cinerea okayama7#130	116510327	Fungi	Eukaryota	772
Uma5	Ustilago maydis 521	71020527	Fungi	Eukaryota	807
Cci3	Coprinopsis cinerea okayama7#130	116506493	Fungi	Eukaryota	1292
Cci5	Coprinopsis cinerea okayama7#130	116509020	Fungi	Eukaryota	771
Lbi4	Laccaria bicolor S238N-H82	164640879	Fungi	Eukaryota	757
Cne3	Cryptococcus neoformans var. neoformans JEC21	58268358	Fungi	Eukaryota	961
Uma1	Ustilago maydis 521	71012856	Fungi	Eukaryota	985
		A viama ma Di	uatain Cira Ctan	dand Daviation (ass), 925 1	102

Average Protein Size \pm Standard Deviation (aas): 825 ± 103

Sub-Cluster 1B (48 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Cal1	Candida albicans	2367386	Fungi	Eukaryota	945
Lel5	Lodderomyces elongisporus NRRL YB-4239	149237448	Fungi	Eukaryota	919
Pst3	Pichia stipitis CBS 6054	126139203	Fungi	Eukarvota	917

Table 1. (Continued)

Dha3	Debaryomyces hansenii CBS767	50419775	Fungi	Eukaryota	907
Pgu8	Pichia guilliermondii ATCC 6260	146421835	Fungi	Eukaryota	881
Cal5	Candida albicans SC5314	87045969	Fungi	Eukaryota	929
Cal6	Candida albicans	87045975	Fungi	Eukaryota	904
Cal3	Candida albicans SC5314	68476729	Fungi	Eukaryota	921
Lel3	Lodderomyces elongisporus NRRL YB-4239	149236581	Fungi	Eukaryota	862
Lel4	Lodderomyces elongisporus NRRL YB-4239	149236916	Fungi	Eukaryota	967
Pst4	Pichia stipitis CBS 6054	146280790	Fungi	Eukaryota	891
Pst9	Pichia stipitis CBS 6054	150866640	Fungi	Eukaryota	913
Pst10	Pichia stipitis CBS 6054	150951233	Fungi	Eukaryota	911
Pgu3	Pichia guilliermondii ATCC 6260	146416529	Fungi	Eukaryota	922
Pgu7	Pichia guilliermondii ATCC 6260	146420005	Fungi	Eukaryota	944
Pgu5	Pichia guilliermondii ATCC 6260	146419149	Fungi	Eukaryota	922
Pst8	Pichia stipitis CBS 6054	150866635	Fungi	Eukaryota	907
Lel7	Lodderomyces elongisporus NRRL YB-4239	149246151	Fungi	Eukaryota	924
Dha2	Debaryomyces hansenii CBS767	50417315	Fungi	Eukaryota	850
Pgu9	Pichia guilliermondii ATCC 6260	146422868	Fungi	Eukaryota	849
Kla2	Kluyveromyces lactis NRRL Y-1140	50307927	Fungi	Eukaryota	869
Sce1	Saccharomyces cerevisiae	6325452	Fungi	Eukaryota	877
Vpo1	Vanderwaltozyma polyspora DSM 70294	156838884	Fungi	Eukaryota	892
Ago1	Ashbya gossypii ATCC 10895	45185387	Fungi	Eukaryota	890
Ago3	Ashbya gossypii ATCC 10895	45187474	Fungi	Eukaryota	885
Ago4	Ashbya gossypii ATCC 10895	45198503	Fungi	Eukaryota	877
Yli10	Yarrowia lipolytica CLIB122	50551841	Fungi	Eukaryota	876
Yli17	Yarrowia lipolytica CLIB122	50557248	Fungi	Eukaryota	767
Yli2	Yarrowia lipolytica CLIB122	50543154	Fungi	Eukaryota	896
Yli12	Yarrowia lipolytica CLIB122	50553458	Fungi	Eukaryota	884
Yli15	Yarrowia lipolytica CLIB122	50555966	Fungi	Eukaryota	882
Yli4	Yarrowia lipolytica CLIB122	50545932	Fungi	Eukaryota	886
Yli3	Yarrowia lipolytica CLIB122	50545745	Fungi	Eukaryota	872
Yli6	Yarrowia lipolytica CLIB122	50548489	Fungi	Eukaryota	883
Yli14	Yarrowia lipolytica CLIB122	50555666	Fungi	Eukaryota	883
Yli8	Yarrowia lipolytica CLIB122	50549187	Fungi	Eukaryota	882
Yli16	Yarrowia lipolytica CLIB122	50556388	Fungi	Eukaryota	948

Table 1. (Continued)

Yli11	Yarrowia lipolytica CLIB122	50553314	Fungi	Eukaryota	879
Yli9	Yarrowia lipolytica CLIB122	50549349	Fungi	Eukaryota	903
Yli13	Yarrowia lipolytica CLIB122	50555622	Fungi	Eukaryota	874
Yli7	Yarrowia lipolytica CLIB122	50549017	Fungi	Eukaryota	1032
Mgr1	Magnaporthe grisea 70-15	39941802	Fungi	Eukaryota	926
Cne1	Cryptococcus neoformans var. neoformans JEC21	58259793	Fungi	Eukaryota	812
Gze1	Gibberella zeae PH-1	46115170	Fungi	Eukaryota	874
Ncr3	Neurospora crassa OR74A	85093666	Fungi	Eukaryota	864
Mgr4	Magnaporthe grisea 70-15	145602334	Fungi	Eukaryota	870
Gze4	Gibberella zeae PH-1	46124369	Fungi	Eukaryota	851
Gze8	Gibberella zeae PH-1	46136533	Fungi	Eukaryota	839

Average Protein Size \pm Standard Deviation (aas): 893 \pm 41

Sub-Cluster 1C (27 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Osa3	Oryza sativa Indica Group	41053195	Viridiplantae	Eukaryota	755
Osa8	Oryza sativa Japonica Group	74267416	Viridiplantae	Eukaryota	751
Vvi12	Vitis vinifera	157355114	Viridiplantae	Eukaryota	744
Ath12	Arabidopsis thaliana	41352045	Viridiplantae	Eukaryota	729
Mtr1	Medicago truncatula	124359202	Viridiplantae	Eukaryota	729
Vvi7	Vitis vinifera	157338674	Viridiplantae	Eukaryota	757
Vvi16	Vitis vinifera	157359604	Viridiplantae	Eukaryota	739
Vvi9	Vitis vinifera	157338676	Viridiplantae	Eukaryota	740
Ath2	Arabidopsis thaliana	15218799	Viridiplantae	Eukaryota	734
Osa16	Oryza sativa Indica Group	115459700	Viridiplantae	Eukaryota	1278
Vvi5	Vitis vinifera	157335739	Viridiplantae	Eukaryota	689
Ath14	Arabidopsis thaliana	67460718	Viridiplantae	Eukaryota	766
Osa9	Oryza sativa Japonica Group	90265681	Viridiplantae	Eukaryota	763
Osa25	Oryza sativa Japonica Group	125540410	Viridiplantae	Eukaryota	766
Osa10	Oryza sativa Indica Group	90265683	Viridiplantae	Eukaryota	771
Vvi13	Vitis vinifera	157355237	Viridiplantae	Eukaryota	690
Ath16	Arabidopsis thaliana	79518939	Viridiplantae	Eukaryota	741
Ath17	Arabidopsis thaliana	145359208	Viridiplantae	Eukaryota	736
Ath9	Arabidopsis thaliana	18402162	Viridiplantae	Eukaryota	733

Vvi8	Vitis vinifera	157338675	Viridiplantae	Eukaryota	731
Ath7	Arabidopsis thaliana	15238763	Viridiplantae	Eukaryota	755
Ath15	Arabidopsis thaliana	79484897	Viridiplantae	Eukaryota	753
Osa31	Oryza sativa Japonica Group	125583075	Viridiplantae	Eukaryota	733
Mac1	Musa acuminata	102140021	Viridiplantae	Eukaryota	748
Osa12	Oryza sativa Japonica Group	115440825	Viridiplantae	Eukaryota	757
Ath3	Arabidopsis thaliana	15234254	Viridiplantae	Eukaryota	737
Ppa2	Physcomitrella patens subsp. patens	162689084	Viridiplantae	Eukaryota	733

Average Protein Size ± Standard Deviation (aas): 761 ± 105

Average Protein Size \pm Standard Deviation (aas): 778 \pm 21

Sub-Cluster 2A (9 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Nfi1	Neosartorya fischeri NRRL 181	119467402	Fungi	Eukaryota	788
Ani8	Aspergillus niger CBS 513.88	145243688	Fungi	Eukaryota	799
Aor8	Aspergillus oryzae	83772997	Fungi	Eukaryota	793
Ssc3	Sclerotinia sclerotiorum 1980	156046206	Fungi	Eukaryota	812
Cal7	Candida albicans	87045979	Fungi	Eukaryota	747
Lel6	Lodderomyces elongisporus NRRL YB-4239	149246053	Fungi	Eukaryota	765
Pst5	Pichia stipitis CBS 6054	150864397	Fungi	Eukaryota	765
Pst2	Pichia stipitis CBS 6054	126139089	Fungi	Eukaryota	771
Pgu4	Pichia guilliermondii ATCC 6260	146417045	Fungi	Eukaryota	760

Sub-Cluster 2B (2 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)	
Ncr4	Neurospora crassa OR74A	85107500	Fungi	Eukaryota	1094	
Bfu1	Botryotinia fuckeliana B05.10	154292901	Fungi	Eukaryota	767	
		Average Protein Size \pm Standard Deviation (aas): 931 \pm 231				

Sub-Cluster 3A (10 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Ssc2	Sclerotinia sclerotiorum 1980	156046040	Fungi	Eukaryota	790
Gze2	Gibberella zeae PH-1	46115236	Fungi	Eukaryota	789
Uma3	Ustilago maydis 521	71016547	Fungi	Eukarvota	797

Acl3	Aspergillus clavatus NRRL 1	121701255	Fungi	Eukaryota	775
Nfi5	Neosartorya fischeri NRRL 181	119488556	Fungi	Eukaryota	757
Ani3	Aspergillus nidulans FGSC A4	67542049	Fungi	Eukaryota	746
Aor4	Aspergillus oryzae	83768691	Fungi	Eukaryota	774
Ate2	Aspergillus terreus NIH2624	115401822	Fungi	Eukaryota	780
Sco1	Schizophyllum commune	6716399	Fungi	Eukaryota	777
Lbi8	Laccaria bicolor S238N-H82	164643810	Fungi	Eukaryota	749

Average Protein Size \pm Standard Deviation (aas): 773 \pm 17

Sub-Cluster 3B (6 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Gze3	Gibberella zeae PH-1	46120458	Fungi	Eukaryota	782
Ncr5	Neurospora crassa OR74A	85113749	Fungi	Eukaryota	788
Bfu3	Botryotinia fuckeliana B05.10	154321612	Fungi	Eukaryota	829
Uma4	Ustilago maydis 521	71019889	Fungi	Eukaryota	860
Ncr7	Neurospora crassa OR74A	164423970	Fungi	Eukaryota	793
Cci2	Coprinopsis cinerea okayama7#130	116504373	Fungi	Eukaryota	824

Average Protein Size \pm Standard Deviation (aas): 813 ± 30

Sub-Cluster 4A (41 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Csp1	Caulobacter sp. K31	113935253	Alphaproteobacteria	Bacteria	662
Ccr1	Caulobacter crescentus CB15	16126881	Alphaproteobacteria	Bacteria	666
Swi1	Sphingomonas wittichii RW1	148555886	Alphaproteobacteria	Bacteria	658
Neu1	Nitrosomonas eutropha C91	114332234	Betaproteobacteria	Bacteria	676
Ssp1	Sphingomonas sp. SKA58	94496206	Alphaproteobacteria	Bacteria	655
Nar1	Novosphingobium aromaticivorans DSM 12444	87199977	Alphaproteobacteria	Bacteria	650
Mtu1	Mycobacterium tuberculosis H37Rv	15609532	Actinobacteria	Bacteria	667
Msm1	Mycobacterium smegmatis str. MC2 155	118470017	Actinobacteria	Bacteria	663
Cdi1	Corynebacterium diphtheriae NCTC 13129	38232950	Actinobacteria	Bacteria	658
Pac1	Propionibacterium acnes KPA171202	50842040	Actinobacteria	Bacteria	662
Aod1	Actinomyces odontolyticus ATCC 17982	154508464	Actinobacteria	Bacteria	666
Cup1	Campylobacter upsaliensis RM3195	57506152	Epsilonproteobacteria	Bacteria	657
Cco1	Campylobacter coli RM2228	57168345	Epsilonproteobacteria	Bacteria	668

Cla1	Campylobacter lari RM2100	57241657	Epsilonproteobacteria	Bacteria	661
Bbr1	Bordetella bronchiseptica RB50	33602645	Betaproteobacteria	Bacteria	693
Bpe1	Bordetella petrii DSM 12804	163856141	Betaproteobacteria	Bacteria	689
Bav1	Bordetella avium 197N	115422286	Betaproteobacteria	Bacteria	677
Rpi1	Ralstonia pickettii 12J	121528839	Betaproteobacteria	Bacteria	684
Rso1	Ralstonia solanacearum GMI1000	17548014	Betaproteobacteria	Bacteria	683
Reu3	Ralstonia eutropha JMP134	113869213	Betaproteobacteria	Bacteria	679
Reu4	Ralstonia eutropha H16	116696492	Betaproteobacteria	Bacteria	679
Rme1	Ralstonia metallidurans CH34	94312045	Betaproteobacteria	Bacteria	676
Pae1	Pseudomonas aeruginosa UCBPP-PA14	116051974	Gammaproteobacteria	Bacteria	678
Hso1	Haemophilus somnus 2336	32029457	Gammaproteobacteria	Bacteria	668
Asu1	Actinobacillus succinogenes 130Z	152977801	Gammaproteobacteria	Bacteria	670
Hdu1	Haemophilus ducreyi 35000HP	33152874	Gammaproteobacteria	Bacteria	669
Apl1	Actinobacillus pleuropneumoniae L20	126209177	Gammaproteobacteria	Bacteria	668
Msu1	Mannheimia succiniciproducens MBEL55E	52424073	Gammaproteobacteria	Bacteria	668
Hin1	Haemophilus influenzae R2866	53733327	Gammaproteobacteria	Bacteria	662
Ngo1	Neisseria gonorrhoeae FA 1090	59802215	Betaproteobacteria	Bacteria	672
Gdi1	Gluconacetobacter diazotrophicus PAl 5	162148874	Alphaproteobacteria	Bacteria	659
Gox1	Gluconobacter oxydans 621H	58038663	Alphaproteobacteria	Bacteria	648
Rgr1	Rickettsiella grylli	160871957	Gammaproteobacteria	Bacteria	654
Lpn1	Legionella pneumophila str. Corby	148360634	Gammaproteobacteria	Bacteria	666
Rgr2	Rickettsiella grylli	160872420	Gammaproteobacteria	Bacteria	669
Xfa1	Xylella fastidiosa Ann-1	71899907	Gammaproteobacteria	Bacteria	653
Sma2	Stenotrophomonas maltophilia R551-3	126466290	Gammaproteobacteria	Bacteria	654
Nmo1	Nitrococcus mobilis Nb-231	88812607	Gammaproteobacteria	Bacteria	655
Pho2	Pyrococcus horikoshii OT3	14590884	Euryarchaeota	Archaea	626
Tko1	Thermococcus kodakarensis KOD1	57641714	Euryarchaeota	Archaea	624
Sde1	Saccharophagus degradans 2-40	90020298	Gammaproteobacteria	Bacteria	672
		A		1.4 (). ((5 + 14	

Average Protein Size ± Standard Deviation (aas): 665 ± 14

Sub-Cluster 4B (16 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Ade1	Anaeromyxobacter dehalogenans 2CP-C	86156672	Deltaproteobacteria	Bacteria	690
Asp5	Anaeromyxobacter sp. Fw109-5	163767022	Deltaproteobacteria	Bacteria	706

Hsp1	Halobacterium sp. NRC-1	16120189	Euryarchaeota	Archaea	655	
Csp2	Clostridium sp. L2-50	160894507	Firmicutes	Bacteria	632	
Eve1	Eubacterium ventriosum ATCC 27560	154484314	Firmicutes	Bacteria	649	
Rgn1	Ruminococcus gnavus ATCC 29149	154504363	Firmicutes	Bacteria	631	
Rto1	Ruminococcus torques ATCC 27756	153813838	Firmicutes	Bacteria	633	
Rob1	Ruminococcus obeum ATCC 29174	153810748	Firmicutes	Bacteria	632	
Hor1	Halothermothrix orenii H 168	89210028	Firmicutes	Bacteria	636	
Cno1	Clostridium novyi NT	118445126	Firmicutes	Bacteria	679	
Cbo1	Clostridium botulinum F str. Langeland	153941447	Firmicutes	Bacteria	651	
Tte1	Thermoanaerobacter tengcongensis MB4	20806685	Firmicutes	Bacteria	647	
Chy1	Carboxydothermus hydrogenoformans Z-2901	78045182	Firmicutes	Bacteria	640	
Dre1	Desulfotomaculum reducens MI-1	134300485	Firmicutes	Bacteria	656	
Sus1	Solibacter usitatus Ellin6076	116620777	Acidobacteria	Bacteria	674	
Aba2	Acidobacteria bacterium Ellin345	94971229	Acidobacteria	Bacteria	675	
	Average Protein Size \pm Standard Deviation (aas): 655 ± 23					

Sub-Cluster 4C (8 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Bun1	Bacteroides uniformis ATCC 8492	160890502	Bacteroidetes	Bacteria	663
Bfr1	Bacteroides fragilis YCH46	53713327	Bacteroidetes	Bacteria	662
Bvu1	Bacteroides vulgatus ATCC 8482	150005284	Bacteroidetes	Bacteria	663
Pdi1	Parabacteroides distasonis ATCC 8503	150008072	Bacteroidetes	Bacteria	665
Pme1	Parabacteroides merdae ATCC 43184	154492906	Bacteroidetes	Bacteria	666
Pgi1	Porphyromonas gingivalis W83	34540265	Bacteroidetes	Bacteria	659
Aba1	Acidobacteria bacterium Ellin345	94969462	Acidobacteria	Bacteria	664
Sus2	Solibacter usitatus Ellin6076	116622365	Acidobacteria	Bacteria	667

Average Protein Size \pm Standard Deviation (aas): 664 \pm 3

Sub-Cluster 4D (8 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Lcal	Lactobacillus casei ATCC 334	116495639	Firmicutes	Bacteria	641
Ppe1	Pediococcus pentosaceus ATCC 25745	116491982	Firmicutes	Bacteria	639
Lsa1	Lactobacillus sakei subsp. sakei 23K	81427933	Firmicutes	Bacteria	645
Ckl1	Clostridium kluyveri DSM 555	153954672	Firmicutes	Bacteria	639

Table	e 1.	(Continued)
IUDI	· • •	(Communaca)

Cbe1	Clostridium beijerinckii NCIMB 8052	150016123	Firmicutes	Bacteria	640
Cba1	Clostridium bartlettii DSM 16795	164687644	Firmicutes	Bacteria	648
Cdi2	Clostridium difficile 630	126699006	Firmicutes	Bacteria	642
Cpe1	Clostridium perfringens str. 13	18310260	Firmicutes	Bacteria	638
•	1 0 0	Average Prote	ein Size ± Standard De	viation (aas): 642 ± 3
		8		`	,
Sub-Cluster	4E (2 Proteins)				
Abberviation	· · · · · · · · · · · · · · · · · · ·	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Cae1	Collinsella aerofaciens ATCC 25986	139438467	Actinobacteria	Bacteria	558
Cce1	Clostridium cellulolyticum H10	118726871	Firmicutes	Bacteria	537
			ein Size ± Standard De	viation (aas): 548 ± 15
					,
Sub-Cluster	4F (2 Proteins)				
Abberviation	n Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Orf1	uncultured methanogenic archaeon RC-I	147920129	Euryarchaeota	Archaea	553
Orf2	uncultured methanogenic archaeon RC-I	147920131	Euryarchaeota	Archaea	552
	C	Average Prote	ein Size ± Standard De	viation (aas): 553 ± 1
		8		`	,
Sub-Cluster	4G (7 Proteins)				
Abberviation	n Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Bsp1	Bacillus sp. B14905	126653239	Firmicutes	Bacteria	524
Vei1	Verminephrobacter eiseniae EF01-2	121610237	Betaproteobacteria	Bacteria	524
Spr1	Serratia proteamaculans 568	157369266	Gammaproteobacteria	Bacteria	524
Bcl1	Bacillus clausii KSM-K16	56962356	Firmicutes	Bacteria	526
Pho1	Pyrococcus horikoshii OT3	14590271	Euryarchaeota	Archaea	527
Mth1	Moorella thermoacetica ATCC 39073	83589078	Firmicutes	Bacteria	519
Rob2	Ruminococcus obeum ATCC 29174	153812663	Firmicutes	Bacteria	558
		Average Prote	ein Size ± Standard De	viation (aas): 529 ± 13
Sub-Cluster	5A (15 Proteins)				
Abberviation	n Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Asp1	Anaeromyxobacter sp. K	153003141	Deltaproteobacteria	Bacteria	540
4 1 2	inter only to oue ter sp. 11				
Ade2	Anaeromyxobacter dehalogenans 2CP-C	86158243	Deltaproteobacteria	Bacteria	540
Ade2 Mxa4	, 1			Bacteria Bacteria	540 592

Sau2	Stigmatella aurantiaca DW4/3-1	115377255	Deltaproteobacteria	Bacteria	637
Mxa5	Myxococcus xanthus DK 1622	108763588	Deltaproteobacteria	Bacteria	631
Asp3	Anaeromyxobacter sp. Fw109-5	153005805	Deltaproteobacteria	Bacteria	605
Mxa2	Myxococcus xanthus DK 1622	108762092	Deltaproteobacteria	Bacteria	606
Sau4	Stigmatella aurantiaca DW4/3-1	115378283	Deltaproteobacteria	Bacteria	625
Psy1	Pseudomonas syringae pv. syringae B728a	66044430	Gammaproteobacteria	Bacteria	581
Ppu1	Pseudomonas putida W619	119857963	Gammaproteobacteria	Bacteria	585
Pst1	Pseudomonas stutzeri A1501	126134803	Gammaproteobacteria	Bacteria	570
Spe1	Shewanella pealeana ATCC 700345	157963678	Gammaproteobacteria	Bacteria	577
Sse1	Shewanella sediminis HAW-EB3	157373494	Gammaproteobacteria	Bacteria	576
Asp2	Anaeromyxobacter sp. K	153003206	Deltaproteobacteria	Bacteria	583
Asp4	Anaeromyxobacter sp. Fw109-5	163766993	Deltaproteobacteria	Bacteria	583
	Average Protein Size ± Standard Deviation (aas): 589 ± 29				

Sub-Cluster 5B (27 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Cal2	Candida albicans SC5314	68475797	Fungi	Eukaryota	718
Pst6	Pichia stipitis CBS 6054	150864483	Fungi	Eukaryota	722
Pgu1	Pichia guilliermondii ATCC 6260	146416523	Fungi	Eukaryota	658
Dha4	Debaryomyces hansenii CBS767	50423315	Fungi	Eukaryota	723
Sce4	Saccharomyces cerevisiae YJM789	162453039	Fungi	Eukaryota	725
Kla4	Kluyveromyces lactis NRRL Y-1140	50311091	Fungi	Eukaryota	732
Vpo2	Vanderwaltozyma polyspora DSM 70294	156848856	Fungi	Eukaryota	733
Cgl2	Candida glabrata CBS 138	116182960	Fungi	Eukaryota	724
Ago2	Ashbya gossypii ATCC 10895	45185483	Fungi	Eukaryota	704
Acl2	Aspergillus clavatus NRRL 1	121699721	Fungi	Eukaryota	800
Afu3	Aspergillus fumigatus Af293	71002356	Fungi	Eukaryota	843
Ani11	Aspergillus nidulans FGSC A4	145249626	Fungi	Eukaryota	754
Aor6	Aspergillus oryzae	83770379	Fungi	Eukaryota	851
Cim1	Coccidioides immitis RS	119186699	Fungi	Eukaryota	797
Ncr2	Neurospora crassa	85075374	Fungi	Eukaryota	738
Cne2	Cryptococcus neoformans var. neoformans JEC21	58265596	Fungi	Eukaryota	740
Lbi3	Laccaria bicolor S238N-H82	164637207	Fungi	Eukaryota	646
Uma2	Ustilago maydis 521	71016340	Fungi	Eukaryota	740

Ddi1	Dictyostelium discoideum AX4	66802892	Slime Mold	Eukaryota	777		
Aor1	Aspergillus oryzae	83766128	Fungi	Eukaryota	725		
Acal	Ajellomyces capsulatus NAm1	154279250	Fungi	Eukaryota	759		
Mgr3	Magnaporthe grisea 70-15	39955178	Fungi	Eukaryota	740		
Cgl1	Chaetomium globosum CBS 148.51	50287709	Fungi	Eukaryota	753		
Gze9	Gibberella zeae PH-1	46138015	Fungi	Eukaryota	743		
Cci4	Coprinopsis cinerea okayama7#130	116509017	Fungi	Eukaryota	726		
Lbi7	Laccaria bicolor S238N-H82	164643762	Fungi	Eukaryota	706		
Uma6	Ustilago maydis 521	71023771	Fungi	Eukaryota	751		
		Average Protein Size \pm Standard Deviation (aas): 742 ± 45					

Sub-Cluster 5C (4 Proteins)

Abberviation	Organism	GenBank Index#	Kingdom	Domain 1	Protein Size (aas)
Reu1	Ralstonia eutropha H16	73539143	Betaproteobacteria	Bacteria	592
Reu2	Ralstonia eutropha JMP134	73542650	Betaproteobacteria	Bacteria	593
Rme2	Ralstonia metallidurans CH34	94314714	Betaproteobacteria	Bacteria	634
Sau3	Stigmatella aurantiaca DW4/3-1	115377807	Deltaproteobacteria	Bacteria	606
		Average Protein Size \pm Standard Deviation (aas): 606 ± 20			

Sub-Cluster 5D (37 Proteins)

Suo-Ciusii 3D (37 Troicius)								
Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)			
Vvi1	Vitis vinifera	147765903	Viridiplantae	Eukaryota	665			
Vvi4	Vitis vinifera	147843808	Viridiplantae	Eukaryota	665			
Ath13	Arabidopsis thaliana	42568235	Viridiplantae	Eukaryota	688			
Tca3	Thlaspi caerulescens	82468795	Viridiplantae	Eukaryota	693			
Vvi6	Vitis vinifera	157335740	Viridiplantae	Eukaryota	713			
Vvi10	Vitis vinifera	157354855	Viridiplantae	Eukaryota	713			
Ath8	Arabidopsis thaliana	15241078	Viridiplantae	Eukaryota	724			
Tca2	Thlaspi caerulescens	82468793	Viridiplantae	Eukaryota	716			
Osa20	Oryza sativa Japonica Group	115466102	Viridiplantae	Eukaryota	708			
Osa11	Oryza sativa Indica Group	115435562	Viridiplantae	Eukaryota	771			
Osa30	Oryza sativa Indica Group	125562004	Viridiplantae	Eukaryota	717			
Osa26	Oryza sativa Japonica Group	125549198	Viridiplantae	Eukaryota	724			
Osa13	Oryza sativa Japonica Group	115455379	Viridiplantae	Eukaryota	882			

Table 1. (Continued)

Osa19	Oryza sativa Japonica Group	115462865	Viridiplantae	Eukaryota	694
Vvi2	Vitis vinifera	147778971	Viridiplantae	Eukaryota	677
Nta1	Nicotiana tabacum	126567465	Viridiplantae	Eukaryota	675
Tca1	Thlaspi caerulescens	82468791	Viridiplantae	Eukaryota	672
Ath6	Arabidopsis thaliana	15238761	Viridiplantae	Eukaryota	675
Ath1	Arabidopsis thaliana	15218331	Viridiplantae	Eukaryota	664
Osa14	Oryza sativa Indica Group	115459506	Viridiplantae	Eukaryota	716
Ath4	Arabidopsis thaliana	15236800	Viridiplantae	Eukaryota	673
Vvi15	Vitis vinifera	157356740	Viridiplantae	Eukaryota	661
Osa2	Oryza sativa Japonica Group	38347209	Viridiplantae	Eukaryota	674
Osa15	Oryza sativa Japonica Group	115459698	Viridiplantae	Eukaryota	726
Zma1	Zea mays	162460137	Viridiplantae	Eukaryota	682
Osa7	Oryza sativa Japonica Group	57834124	Viridiplantae	Eukaryota	672
Hvu1	Hordeum vulgare subsp. vulgare	84453180	Viridiplantae	Eukaryota	678
Ath5	Arabidopsis thaliana	15236912	Viridiplantae	Eukaryota	670
Ath11	Arabidopsis thaliana	25083021	Viridiplantae	Eukaryota	677
Osa1	Oryza sativa Indica Group	28144882	Viridiplantae	Eukaryota	678
Ppa3	Physcomitrella patens subsp. patens	162697041	Viridiplantae	Eukaryota	661
Osa28	Oryza sativa Japonica Group	125553884	Viridiplantae	Eukaryota	724
Osa5	Oryza sativa Japonica Group	49387869	Viridiplantae	Eukaryota	708
Osa22	Oryza sativa Indica Group	116309354	Viridiplantae	Eukaryota	717
Osa23	Oryza sativa Japonica Group	116310949	Viridiplantae	Eukaryota	683
Osa21	Oryza sativa Japonica Group	115466104	Viridiplantae	Eukaryota	679
Osa4	Oryza sativa Japonica Group	42409160	Viridiplantae	Eukaryota	686

Average Protein Size \pm Standard Deviation (aas): 697 \pm 40

Table 2. OPT protein sequences included in this study. Proteins are listed in alphabetical order according to genus and species. Protein abbreviation, GenBank Index#, Kingdom, Domain, and protein size are also presented in the list of proteins.

Abberviation	Organism	GenBank Index#	Kingdom	Domain	Protein Size (aas)
Abal	Acidobacteria bacterium Ellin345	94969462	Acidobacteria	Bacteria	664
Aba2	Acidobacteria bacterium Ellin345	94971229	Acidobacteria	Bacteria	675
Apl1	Actinobacillus pleuropneumoniae L20	126209177	Gammaproteobacteria	Bacteria	668
Asu1	Actinobacillus succinogenes 130Z	152977801	Gammaproteobacteria	Bacteria	670
Aod1	Actinomyces odontolyticus ATCC 17982	154508464	Actinobacteria	Bacteria	666
Aca1	Ajellomyces capsulatus NAm1	154279250	Fungi	Eukaryota	759
Adel	Anaeromyxobacter dehalogenans 2CP-C	86156672	Deltaproteobacteria	Bacteria	690
Ade2	Anaeromyxobacter dehalogenans 2CP-C	86158243	Deltaproteobacteria	Bacteria	540
Asp3	Anaeromyxobacter sp. Fw109-5	153005805	Deltaproteobacteria	Bacteria	605
Asp4	Anaeromyxobacter sp. Fw109-5	163766993	Deltaproteobacteria	Bacteria	583
Asp5	Anaeromyxobacter sp. Fw109-5	163767022	Deltaproteobacteria	Bacteria	706
Asp1	Anaeromyxobacter sp. K	153003141	Deltaproteobacteria	Bacteria	540
Asp2	Anaeromyxobacter sp. K	153003206	Deltaproteobacteria	Bacteria	583
Ath1	Arabidopsis thaliana	15218331	Viridiplantae	Eukaryota	664
Ath2	Arabidopsis thaliana	15218799	Viridiplantae	Eukaryota	734
Ath3	Arabidopsis thaliana	15234254	Viridiplantae	Eukaryota	737
Ath4	Arabidopsis thaliana	15236800	Viridiplantae	Eukaryota	673
Ath5	Arabidopsis thaliana	15236912	Viridiplantae	Eukaryota	670
Ath6	Arabidopsis thaliana	15238761	Viridiplantae	Eukaryota	675
Ath7	Arabidopsis thaliana	15238763	Viridiplantae	Eukaryota	755
Ath8	Arabidopsis thaliana	15241078	Viridiplantae	Eukaryota	724
Ath9	Arabidopsis thaliana	18402162	Viridiplantae	Eukaryota	733
Ath11	Arabidopsis thaliana	25083021	Viridiplantae	Eukaryota	677
Ath12	Arabidopsis thaliana	41352045	Viridiplantae	Eukaryota	729
Ath13	Arabidopsis thaliana	42568235	Viridiplantae	Eukaryota	688
Ath14	Arabidopsis thaliana	67460718	Viridiplantae	Eukaryota	766
Ath15	Arabidopsis thaliana	79484897	Viridiplantae	Eukaryota	753
Ath16	Arabidopsis thaliana	79518939	Viridiplantae	Eukaryota	741
Ath17	Arabidopsis thaliana	145359208	Viridiplantae	Eukaryota	736
Ago1	Ashbya gossypii ATCC 10895	45185387	Fungi	Eukaryota	890
Ago2	Ashbya gossypii ATCC 10895	45185483	Fungi	Eukaryota	704

 Table 2. (Continued)

Ago3	Ashbya gossypii ATCC 10895	45187474	Fungi	Eukaryota	885
Ago4	Ashbya gossypii ATCC 10895	45198503	Fungi	Eukaryota	877
Ago5	Ashbya gossypii ATCC 10895	45201069	Fungi	Eukaryota	796
Acl1	Aspergillus clavatus NRRL 1	121699197	Fungi	Eukaryota	788
Acl2	Aspergillus clavatus NRRL 1	121699721	Fungi	Eukaryota	800
Acl3	Aspergillus clavatus NRRL 1	121701255	Fungi	Eukaryota	775
Acl4	Aspergillus clavatus NRRL 1	121705906	Fungi	Eukaryota	793
Acl5	Aspergillus clavatus NRRL 1	121709515	Fungi	Eukaryota	761
Afu2	Aspergillus fumigatus Af293	70999364	Fungi	Eukaryota	792
Afu3	Aspergillus fumigatus Af293	71002356	Fungi	Eukaryota	843
Ani3	Aspergillus nidulans FGSC A4	67542049	Fungi	Eukaryota	746
Ani4	Aspergillus nidulans FGSC A4	67901220	Fungi	Eukaryota	794
Ani11	Aspergillus nidulans FGSC A4	145249626	Fungi	Eukaryota	754
Ani1	Aspergillus niger CBS 513.88	67516837	Fungi	Eukaryota	792
Ani2	Aspergillus niger CBS 513.88	67540564	Fungi	Eukaryota	778
Ani6	Aspergillus niger CBS 513.88	145241488	Fungi	Eukaryota	859
Ani8	Aspergillus niger CBS 513.88	145243688	Fungi	Eukaryota	799
Ani12	Aspergillus niger CBS 513.88	145251507	Fungi	Eukaryota	771
Aor1	Aspergillus oryzae	83766128	Fungi	Eukaryota	725
Aor2	Aspergillus oryzae	83768389	Fungi	Eukaryota	765
Aor3	Aspergillus oryzae	83768538	Fungi	Eukaryota	751
Aor4	Aspergillus oryzae	83768691	Fungi	Eukaryota	774
Aor5	Aspergillus oryzae	83768732	Fungi	Eukaryota	770
Aor6	Aspergillus oryzae	83770379	Fungi	Eukaryota	851
Aor7	Aspergillus oryzae	83770544	Fungi	Eukaryota	778
Aor8	Aspergillus oryzae	83772997	Fungi	Eukaryota	793
Aor9	Aspergillus oryzae	83775779	Fungi	Eukaryota	768
Ate1	Aspergillus terreus NIH2624	115397517	Fungi	Eukaryota	788
Ate2	Aspergillus terreus NIH2624	115401822	Fungi	Eukaryota	780
Bcl1	Bacillus clausii KSM-K16	56962356	Firmicutes	Bacteria	526
Bsp1	Bacillus sp. B14905	126653239	Firmicutes	Bacteria	524
Bfr1	Bacteroides fragilis YCH46	53713327	Bacteroidetes	Bacteria	662
Bun1	Bacteroides uniformis ATCC 8492	160890502	Bacteroidetes	Bacteria	663

Table 2. (Continued)

Bvu1	Bacteroides vulgatus ATCC 8482	150005284	Bacteroidetes	Bacteria	663
Bav1	Bordetella avium 197N	115422286	Betaproteobacteria	Bacteria	677
Bbr1	Bordetella bronchiseptica RB50	33602645	Betaproteobacteria	Bacteria	693
Bpe1	Bordetella petrii DSM 12804	163856141	Betaproteobacteria	Bacteria	689
Bfu1	Botryotinia fuckeliana B05.10	154292901	Fungi	Eukaryota	767
Bfu2	Botryotinia fuckeliana B05.10	154313655	Fungi	Eukaryota	779
Bfu3	Botryotinia fuckeliana B05.10	154321612	Fungi	Eukaryota	829
Cco1	Campylobacter coli RM2228	57168345	Epsilonproteobacteria	Bacteria	668
Cla1	Campylobacter lari RM2100	57241657	Epsilonproteobacteria	Bacteria	661
Cup1	Campylobacter upsaliensis RM3195	57506152	Epsilonproteobacteria	Bacteria	657
Cal1	Candida albicans	2367386	Fungi	Eukaryota	945
Cal4	Candida albicans	68485275	Fungi	Eukaryota	783
Cal6	Candida albicans	87045975	Fungi	Eukaryota	904
Cal7	Candida albicans	87045979	Fungi	Eukaryota	747
Cal2	Candida albicans SC5314	68475797	Fungi	Eukaryota	718
Cal3	Candida albicans SC5314	68476729	Fungi	Eukaryota	921
Cal5	Candida albicans SC5314	87045969	Fungi	Eukaryota	929
Cgl2	Candida glabrata CBS 138	116182960	Fungi	Eukaryota	724
Chy1	Carboxydothermus hydrogenoformans Z-2901	78045182	Firmicutes	Bacteria	640
Ccrl	Caulobacter crescentus CB15	16126881	Alphaproteobacteria	Bacteria	666
Csp1	Caulobacter sp. K31	113935253	Alphaproteobacteria	Bacteria	662
Cgl1	Chaetomium globosum CBS 148.51	50287709	Fungi	Eukaryota	753
Cgl3	Chaetomium globosum CBS 148.51	116193201	Fungi	Eukaryota	1027
Cgl5	Chaetomium globosum CBS 148.51	116198757	Fungi	Eukaryota	871
Cba1	Clostridium bartlettii DSM 16795	164687644	Firmicutes	Bacteria	648
Cbe1	Clostridium beijerinckii NCIMB 8052	150016123	Firmicutes	Bacteria	640
Cbo1	Clostridium botulinum F str. Langeland	153941447	Firmicutes	Bacteria	651
Cce1	Clostridium cellulolyticum H10	118726871	Firmicutes	Bacteria	537
Cdi2	Clostridium difficile 630	126699006	Firmicutes	Bacteria	642
Ckl1	Clostridium kluyveri DSM 555	153954672	Firmicutes	Bacteria	639
Cno1	Clostridium novyi NT	118445126	Firmicutes	Bacteria	679
Cpe1	Clostridium perfringens str. 13	18310260	Firmicutes	Bacteria	638
Csp2	Clostridium sp. L2-50	160894507	Firmicutes	Bacteria	632

Table 2. (Continued)

Cim1	Coccidioides immitis RS	119186699	Fungi	Eukaryota	797
Cim2	Coccidioides immitis RS	119190959	Fungi	Eukaryota	810
Cim3	Coccidioides immitis RS	119194107	Fungi	Eukaryota	812
Cae1	Collinsella aerofaciens ATCC 25986	139438467	Actinobacteria	Bacteria	558
Cci1	Coprinopsis cinerea okayama7#130	116500528	Fungi	Eukaryota	757
Cci2	Coprinopsis cinerea okayama7#130	116504373	Fungi	Eukaryota	824
Cci3	Coprinopsis cinerea okayama7#130	116506493	Fungi	Eukaryota	1292
Cci4	Coprinopsis cinerea okayama7#130	116509017	Fungi	Eukaryota	726
Cci5	Coprinopsis cinerea okayama7#130	116509020	Fungi	Eukaryota	771
Cci6	Coprinopsis cinerea okayama7#130	116510327	Fungi	Eukaryota	772
Cdi1	Corynebacterium diphtheriae NCTC 13129	38232950	Actinobacteria	Bacteria	658
Cne4	Cryptococcus neoformans var. neoformans B-3501A	134113154	Fungi	Eukaryota	797
Cne1	Cryptococcus neoformans var. neoformans JEC21	58259793	Fungi	Eukaryota	812
Cne2	Cryptococcus neoformans var. neoformans JEC21	58265596	Fungi	Eukaryota	740
Cne3	Cryptococcus neoformans var. neoformans JEC21	58268358	Fungi	Eukaryota	961
Dha1	Debaryomyces hansenii CBS767	50413511	Fungi	Eukaryota	776
Dha2	Debaryomyces hansenii CBS767	50417315	Fungi	Eukaryota	850
Dha3	Debaryomyces hansenii CBS767	50419775	Fungi	Eukaryota	907
Dha4	Debaryomyces hansenii CBS767	50423315	Fungi	Eukaryota	723
Dre1	Desulfotomaculum reducens MI-1	134300485	Firmicutes	Bacteria	656
Ddi1	Dictyostelium discoideum AX4	66802892	Slime Mold	Eukaryota	777
Eve1	Eubacterium ventriosum ATCC 27560	154484314	Firmicutes	Bacteria	649
Gze1	Gibberella zeae PH-1	46115170	Fungi	Eukaryota	874
Gze2	Gibberella zeae PH-1	46115236	Fungi	Eukaryota	789
Gze3	Gibberella zeae PH-1	46120458	Fungi	Eukaryota	782
Gze4	Gibberella zeae PH-1	46124369	Fungi	Eukaryota	851
Gze5	Gibberella zeae PH-1	46125699	Fungi	Eukaryota	1060
Gze7	Gibberella zeae PH-1	46134295	Fungi	Eukaryota	799
Gze8	Gibberella zeae PH-1	46136533	Fungi	Eukaryota	839
Gze9	Gibberella zeae PH-1	46138015	Fungi	Eukaryota	743
Gdi1	Gluconacetobacter diazotrophicus PAl 5	162148874	Alphaproteobacteria	Bacteria	659
Gox1	Gluconobacter oxydans 621H	58038663	Alphaproteobacteria	Bacteria	648
Hdu1	Haemophilus ducreyi 35000HP	33152874	Gammaproteobacteria	Bacteria	669

Table 2. (Continued)

Hin1	Haemophilus influenzae R2866	53733327	Gammaproteobacteria	Bacteria	662
Hso1	Haemophilus somnus 2336	32029457	Gammaproteobacteria	Bacteria	668
Hsp1	Halobacterium sp. NRC-1	16120189	Euryarchaeota	Archaea	655
Hor1	Halothermothrix orenii H 168	89210028	Firmicutes	Bacteria	636
Hvu1	Hordeum vulgare subsp. vulgare	84453180	Viridiplantae	Eukaryota	678
Kla1	Kluyveromyces lactis NRRL Y-1140	50307527	Fungi	Eukaryota	794
Kla2	Kluyveromyces lactis NRRL Y-1140	50307927	Fungi	Eukaryota	869
Kla3	Kluyveromyces lactis NRRL Y-1140	50307929	Fungi	Eukaryota	793
Kla4	Kluyveromyces lactis NRRL Y-1140	50311091	Fungi	Eukaryota	732
Lbi3	Laccaria bicolor S238N-H82	164637207	Fungi	Eukaryota	646
Lbi4	Laccaria bicolor S238N-H82	164640879	Fungi	Eukaryota	757
Lbi5	Laccaria bicolor S238N-H82	164641826	Fungi	Eukaryota	730
Lbi7	Laccaria bicolor S238N-H82	164643762	Fungi	Eukaryota	706
Lbi8	Laccaria bicolor S238N-H82	164643810	Fungi	Eukaryota	749
Lca1	Lactobacillus casei ATCC 334	116495639	Firmicutes	Bacteria	641
Lsa1	Lactobacillus sakei subsp. sakei 23K	81427933	Firmicutes	Bacteria	645
Lpn1	Legionella pneumophila str. Corby	148360634	Gammaproteobacteria	Bacteria	666
Lel2	Lodderomyces elongisporus NRRL YB-4239	149235877	Fungi	Eukaryota	804
Lel3	Lodderomyces elongisporus NRRL YB-4239	149236581	Fungi	Eukaryota	862
Lel4	Lodderomyces elongisporus NRRL YB-4239	149236916	Fungi	Eukaryota	967
Lel5	Lodderomyces elongisporus NRRL YB-4239	149237448	Fungi	Eukaryota	919
Lel6	Lodderomyces elongisporus NRRL YB-4239	149246053	Fungi	Eukaryota	765
Lel7	Lodderomyces elongisporus NRRL YB-4239	149246151	Fungi	Eukaryota	924
Mgr1	Magnaporthe grisea 70-15	39941802	Fungi	Eukaryota	926
Mgr2	Magnaporthe grisea 70-15	39944474	Fungi	Eukaryota	783
Mgr3	Magnaporthe grisea 70-15	39955178	Fungi	Eukaryota	740
Mgr4	Magnaporthe grisea 70-15	145602334	Fungi	Eukaryota	870
Mgr5	Magnaporthe grisea 70-15	145614314	Fungi	Eukaryota	849
Msu1	Mannheimia succiniciproducens MBEL55E	52424073	Gammaproteobacteria	Bacteria	668
Mtr1	Medicago truncatula	124359202	Viridiplantae	Eukaryota	729
Mth1	Moorella thermoacetica ATCC 39073	83589078	Firmicutes	Bacteria	519
Mac1	Musa acuminata	102140021	Viridiplantae	Eukaryota	748
Msm1	Mycobacterium smegmatis str. MC2 155	118470017	Actinobacteria	Bacteria	663

Table 2. (Continued)

Mtu1	Mycobacterium tuberculosis H37Rv	15609532	Actinobacteria	Bacteria	667
Mxa2	Myxococcus xanthus DK 1622	108762092	Deltaproteobacteria	Bacteria	606
Mxa4	Myxococcus xanthus DK 1622	108763515	Deltaproteobacteria	Bacteria	592
Mxa5	Myxococcus xanthus DK 1622	108763588	Deltaproteobacteria	Bacteria	631
Ngo1	Neisseria gonorrhoeae FA 1090	59802215	Betaproteobacteria	Bacteria	672
Nfi1	Neosartorya fischeri NRRL 181	119467402	Fungi	Eukaryota	788
Nfi2	Neosartorya fischeri NRRL 181	119471104	Fungi	Eukaryota	757
Nfi3	Neosartorya fischeri NRRL 181	119471211	Fungi	Eukaryota	770
Nfi4	Neosartorya fischeri NRRL 181	119477757	Fungi	Eukaryota	772
Nfi5	Neosartorya fischeri NRRL 181	119488556	Fungi	Eukaryota	757
Nfi6	Neosartorya fischeri NRRL 181	119491377	Fungi	Eukaryota	768
Ncr2	Neurospora crassa	85075374	Fungi	Eukaryota	738
Ncr1	Neurospora crassa OR74A	9368956	Fungi	Eukaryota	801
Ncr3	Neurospora crassa OR74A	85093666	Fungi	Eukaryota	864
Ncr4	Neurospora crassa OR74A	85107500	Fungi	Eukaryota	1094
Ncr5	Neurospora crassa OR74A	85113749	Fungi	Eukaryota	788
Ncr6	Neurospora crassa OR74A	164422675	Fungi	Eukaryota	1094
Ncr7	Neurospora crassa OR74A	164423970	Fungi	Eukaryota	793
Nta1	Nicotiana tabacum	126567465	Viridiplantae	Eukaryota	675
Nmo1	Nitrococcus mobilis Nb-231	88812607	Gammaproteobacteria	Bacteria	655
Neu1	Nitrosomonas eutropha C91	114332234	Betaproteobacteria	Bacteria	676
Nar1	Novosphingobium aromaticivorans DSM 12444	87199977	Alphaproteobacteria	Bacteria	650
Osa1	Oryza sativa Indica Group	28144882	Viridiplantae	Eukaryota	678
Osa3	Oryza sativa Indica Group	41053195	Viridiplantae	Eukaryota	755
Osa10	Oryza sativa Indica Group	90265683	Viridiplantae	Eukaryota	771
Osa11	Oryza sativa Indica Group	115435562	Viridiplantae	Eukaryota	771
Osa14	Oryza sativa Indica Group	115459506	Viridiplantae	Eukaryota	716
Osa16	Oryza sativa Indica Group	115459700	Viridiplantae	Eukaryota	1278
Osa22	Oryza sativa Indica Group	116309354	Viridiplantae	Eukaryota	717
Osa30	Oryza sativa Indica Group	125562004	Viridiplantae	Eukaryota	717
Osa2	Oryza sativa Japonica Group	38347209	Viridiplantae	Eukaryota	674
Osa4	Oryza sativa Japonica Group	42409160	Viridiplantae	Eukaryota	686
Osa5	Oryza sativa Japonica Group	49387869	Viridiplantae	Eukaryota	708

Table 2. (Continued)

Osa7	Oryza sativa Japonica Group	57834124	Viridiplantae	Eukaryota	672
Osa8	Oryza sativa Japonica Group	74267416	Viridiplantae	Eukaryota	751
Osa9	Oryza sativa Japonica Group	90265681	Viridiplantae	Eukaryota	763
Osa12	Oryza sativa Japonica Group	115440825	Viridiplantae	Eukaryota	757
Osa13	Oryza sativa Japonica Group	115455379	Viridiplantae	Eukaryota	882
Osa15	Oryza sativa Japonica Group	115459698	Viridiplantae	Eukaryota	726
Osa19	Oryza sativa Japonica Group	115462865	Viridiplantae	Eukaryota	694
Osa20	Oryza sativa Japonica Group	115466102	Viridiplantae	Eukaryota	708
Osa21	Oryza sativa Japonica Group	115466104	Viridiplantae	Eukaryota	679
Osa23	Oryza sativa Japonica Group	116310949	Viridiplantae	Eukaryota	683
Osa25	Oryza sativa Japonica Group	125540410	Viridiplantae	Eukaryota	766
Osa26	Oryza sativa Japonica Group	125549198	Viridiplantae	Eukaryota	724
Osa28	Oryza sativa Japonica Group	125553884	Viridiplantae	Eukaryota	724
Osa31	Oryza sativa Japonica Group	125583075	Viridiplantae	Eukaryota	733
Pdi1	Parabacteroides distasonis ATCC 8503	150008072	Bacteroidetes	Bacteria	665
Pme1	Parabacteroides merdae ATCC 43184	154492906	Bacteroidetes	Bacteria	666
Ppe1	Pediococcus pentosaceus ATCC 25745	116491982	Firmicutes	Bacteria	639
Pno1	Phaeosphaeria nodorum SN15	160705030	Fungi	Eukaryota	845
Ppa2	Physcomitrella patens subsp. patens	162689084	Viridiplantae	Eukaryota	733
Ppa3	Physcomitrella patens subsp. patens	162697041	Viridiplantae	Eukaryota	661
Pgu1	Pichia guilliermondii ATCC 6260	146416523	Fungi	Eukaryota	658
Pgu2	Pichia guilliermondii ATCC 6260	146416527	Fungi	Eukaryota	784
Pgu3	Pichia guilliermondii ATCC 6260	146416529	Fungi	Eukaryota	922
Pgu4	Pichia guilliermondii ATCC 6260	146417045	Fungi	Eukaryota	760
Pgu5	Pichia guilliermondii ATCC 6260	146419149	Fungi	Eukaryota	922
Pgu6	Pichia guilliermondii ATCC 6260	146419361	Fungi	Eukaryota	754
Pgu7	Pichia guilliermondii ATCC 6260	146420005	Fungi	Eukaryota	944
Pgu8	Pichia guilliermondii ATCC 6260	146421835	Fungi	Eukaryota	881
Pgu9	Pichia guilliermondii ATCC 6260	146422868	Fungi	Eukaryota	849
Pst2	Pichia stipitis CBS 6054	126139089	Fungi	Eukaryota	771
Pst3	Pichia stipitis CBS 6054	126139203	Fungi	Eukaryota	917
Pst4	Pichia stipitis CBS 6054	146280790	Fungi	Eukaryota	891
Pst5	Pichia stipitis CBS 6054	150864397	Fungi	Eukaryota	765

Table 2. (Continued)

Pst6	Pichia stipitis CBS 6054	150864483	Fungi	Eukaryota	722
Pst7	Pichia stipitis CBS 6054	150864787	Fungi	Eukaryota	782
Pst8	Pichia stipitis CBS 6054	150866635	Fungi	Eukaryota	907
Pst9	Pichia stipitis CBS 6054	150866640	Fungi	Eukaryota	913
Pst10	Pichia stipitis CBS 6054	150951233	Fungi	Eukaryota	911
Pgi1	Porphyromonas gingivalis W83	34540265	Bacteroidetes	Bacteria	659
Pac1	Propionibacterium acnes KPA171202	50842040	Actinobacteria	Bacteria	662
Pae1	Pseudomonas aeruginosa UCBPP-PA14	116051974	Gammaproteobacteria	Bacteria	678
Ppu1	Pseudomonas putida W619	119857963	Gammaproteobacteria	Bacteria	585
Pst1	Pseudomonas stutzeri A1501	126134803	Gammaproteobacteria	Bacteria	570
Psy1	Pseudomonas syringae pv. syringae B728a	66044430	Gammaproteobacteria	Bacteria	581
Pho1	Pyrococcus horikoshii OT3	14590271	Euryarchaeota	Archaea	527
Pho2	Pyrococcus horikoshii OT3	14590884	Euryarchaeota	Archaea	626
Reu1	Ralstonia eutropha H16	73539143	Betaproteobacteria	Bacteria	592
Reu4	Ralstonia eutropha H16	116696492	Betaproteobacteria	Bacteria	679
Reu2	Ralstonia eutropha JMP134	73542650	Betaproteobacteria	Bacteria	593
Reu3	Ralstonia eutropha JMP134	113869213	Betaproteobacteria	Bacteria	679
Rme1	Ralstonia metallidurans CH34	94312045	Betaproteobacteria	Bacteria	676
Rme2	Ralstonia metallidurans CH34	94314714	Betaproteobacteria	Bacteria	634
Rpi1	Ralstonia pickettii 12J	121528839	Betaproteobacteria	Bacteria	684
Rso1	Ralstonia solanacearum GMI1000	17548014	Betaproteobacteria	Bacteria	683
Rgr1	Rickettsiella grylli	160871957	Gammaproteobacteria	Bacteria	654
Rgr2	Rickettsiella grylli	160872420	Gammaproteobacteria	Bacteria	669
Rgn1	Ruminococcus gnavus ATCC 29149	154504363	Firmicutes	Bacteria	631
Rob1	Ruminococcus obeum ATCC 29174	153810748	Firmicutes	Bacteria	632
Rob2	Ruminococcus obeum ATCC 29174	153812663	Firmicutes	Bacteria	558
Rto1	Ruminococcus torques ATCC 27756	153813838	Firmicutes	Bacteria	633
Sce1	Saccharomyces cerevisiae	6325452	Fungi	Eukaryota	877
Sce2	Saccharomyces cerevisiae YJM789	151943695	Fungi	Eukaryota	799
Sce4	Saccharomyces cerevisiae YJM789	162453039	Fungi	Eukaryota	725
Sde1	Saccharophagus degradans 2-40	90020298	Gammaproteobacteria	Bacteria	672
Sco1	Schizophyllum commune	6716399	Fungi	Eukaryota	777
Spo2	Schizosaccharomyces pombe	19115899	Fungi	Eukaryota	785

Table 2. (Continued)

Spo3	Schizosaccharomyces pombe	63054465	Fungi	Eukaryota	851
Ssc1	Sclerotinia sclerotiorum 1980	156039822	Fungi	Eukaryota	1055
Ssc2	Sclerotinia sclerotiorum 1980	156046040	Fungi	Eukaryota	790
Ssc3	Sclerotinia sclerotiorum 1980	156046206	Fungi	Eukaryota	812
Ssc4	Sclerotinia sclerotiorum 1980	156049297	Fungi	Eukaryota	827
Spr1	Serratia proteamaculans 568	157369266	Gammaproteobacteria	Bacteria	524
Spe1	Shewanella pealeana ATCC 700345	157963678	Gammaproteobacteria	Bacteria	577
Sse1	Shewanella sediminis HAW-EB3	157373494	Gammaproteobacteria	Bacteria	576
Sus1	Solibacter usitatus Ellin6076	116620777	Acidobacteria	Bacteria	674
Sus2	Solibacter usitatus Ellin6076	116622365	Acidobacteria	Bacteria	667
Ssp1	Sphingomonas sp. SKA58	94496206	Alphaproteobacteria	Bacteria	655
Swi1	Sphingomonas wittichii RW1	148555886	Alphaproteobacteria	Bacteria	658
Sma2	Stenotrophomonas maltophilia R551-3	126466290	Gammaproteobacteria	Bacteria	654
Sau2	Stigmatella aurantiaca DW4/3-1	115377255	Deltaproteobacteria	Bacteria	637
Sau3	Stigmatella aurantiaca DW4/3-1	115377807	Deltaproteobacteria	Bacteria	606
Sau4	Stigmatella aurantiaca DW4/3-1	115378283	Deltaproteobacteria	Bacteria	625
Tte1	Thermoanaerobacter tengcongensis MB4	20806685	Firmicutes	Bacteria	647
Tko1	Thermococcus kodakarensis KOD1	57641714	Euryarchaeota	Archaea	624
Tca1	Thlaspi caerulescens	82468791	Viridiplantae	Eukaryota	672
Tca2	Thlaspi caerulescens	82468793	Viridiplantae	Eukaryota	716
Tca3	Thlaspi caerulescens	82468795	Viridiplantae	Eukaryota	693
Orf1	uncultured methanogenic archaeon RC-I	147920129	Euryarchaeota	Archaea	553
Orf2	uncultured methanogenic archaeon RC-I	147920131	Euryarchaeota	Archaea	552
Uma1	Ustilago maydis 521	71012856	Fungi	Eukaryota	985
Uma2	Ustilago maydis 521	71016340	Fungi	Eukaryota	740
Uma3	Ustilago maydis 521	71016547	Fungi	Eukaryota	797
Uma4	Ustilago maydis 521	71019889	Fungi	Eukaryota	860
Uma5	Ustilago maydis 521	71020527	Fungi	Eukaryota	807
Uma6	Ustilago maydis 521	71023771	Fungi	Eukaryota	751
Vpo1	Vanderwaltozyma polyspora DSM 70294	156838884	Fungi	Eukaryota	892
Vpo2	Vanderwaltozyma polyspora DSM 70294	156848856	Fungi	Eukaryota	733
Vei1	Verminephrobacter eiseniae EF01-2	121610237	Betaproteobacteria	Bacteria	524
Vvi1	Vitis vinifera	147765903	Viridiplantae	Eukaryota	665

Table 2. (Continued)

Vvi2	Vitis vinifera	147778971	Viridiplantae	Eukaryota	677
Vvi4	Vitis vinifera	147843808	Viridiplantae	Eukaryota	665
Vvi5	Vitis vinifera	157335739	Viridiplantae	Eukaryota	689
Vvi6	Vitis vinifera	157335740	Viridiplantae	Eukaryota	713
Vvi7	Vitis vinifera	157338674	Viridiplantae	Eukaryota	757
Vvi8	Vitis vinifera	157338675	Viridiplantae	Eukaryota	731
Vvi9	Vitis vinifera	157338676	Viridiplantae	Eukaryota	740
Vvi10	Vitis vinifera	157354855	Viridiplantae	Eukaryota	713
Vvi12	Vitis vinifera	157355114	Viridiplantae	Eukaryota	744
Vvi13	Vitis vinifera	157355237	Viridiplantae	Eukaryota	690
Vvi15	Vitis vinifera	157356740	Viridiplantae	Eukaryota	661
Vvi16	Vitis vinifera	157359604	Viridiplantae	Eukaryota	739
Xfa1	Xylella fastidiosa Ann-1	71899907	Gammaproteobacteria	Bacteria	653
Yli1	Yarrowia lipolytica CLIB122	50542874	Fungi	Eukaryota	836
Yli2	Yarrowia lipolytica CLIB122	50543154	Fungi	Eukaryota	896
Yli3	Yarrowia lipolytica CLIB122	50545745	Fungi	Eukaryota	872
Yli4	Yarrowia lipolytica CLIB122	50545932	Fungi	Eukaryota	886
Yli6	Yarrowia lipolytica CLIB122	50548489	Fungi	Eukaryota	883
Yli7	Yarrowia lipolytica CLIB122	50549017	Fungi	Eukaryota	1032
Yli8	Yarrowia lipolytica CLIB122	50549187	Fungi	Eukaryota	882
Yli9	Yarrowia lipolytica CLIB122	50549349	Fungi	Eukaryota	903
Yli10	Yarrowia lipolytica CLIB122	50551841	Fungi	Eukaryota	876
Yli11	Yarrowia lipolytica CLIB122	50553314	Fungi	Eukaryota	879
Yli12	Yarrowia lipolytica CLIB122	50553458	Fungi	Eukaryota	884
Yli13	Yarrowia lipolytica CLIB122	50555622	Fungi	Eukaryota	874
Yli14	Yarrowia lipolytica CLIB122	50555666	Fungi	Eukaryota	883
Yli15	Yarrowia lipolytica CLIB122	50555966	Fungi	Eukaryota	882
Yli16	Yarrowia lipolytica CLIB122	50556388	Fungi	Eukaryota	948
Yli17	Yarrowia lipolytica CLIB122	50557248	Fungi	Eukaryota	767
Zma1	Zea mays	162460137	Viridiplantae	Eukaryota	682

Table 3. Comparison of different segments within OPT proteins using the GAP and IC programs. Entry 1 presents comparisons for the first 8 TMS half versus the second 8 TMS half. Entries 2-7 present comparisons for the four 4 TMS quarters compared to each other. Entries 8-10 present comparisons for four representative adjacent 2 TMS hairpin structures.

			Amino			Amino		IC/GAP Score	Average Score	
Comparison	Segment	Protein-1	Acids	TMS	Protein-2	Acids	TMS	(S.D.)	(S.D.)	
1. 1-8 vs 9-16	AB vs CD	Spr1	16 - 241	1 to 8	Lsa1	358 - 589	9 to 16	12.6	12.0	
	AB vs CD	Zma1	51 - 216	1 to 4	Chy1	358 - 505	9 to 12	11.3	12.0	
2. 1-4 vs 5-8	A vs B	Hso1	41 - 139	1 to 3	Sde1	174 - 270	5 to 7	11.9	11.3	
2. 1-4 VS 3-6	A vs B	Ngo1	45 - 143	1 to 3	Sde1	174 - 270	5 to 7	10.7	11.5	
3. 1-4 vs 9-12	A vs C	Zma1	51 - 159	1 to 3	Chy1	358 - 455	9 to 11	12.5	12.2	
3. 1-4 VS 9-12	A vs C	Mth1	14 - 123	1 to 4	Mgr3	467 - 577	9 to 12	11.9	12.2	
4. 1-4 vs 13-16	A vs D	Gze4	139 - 266	1 to 2	Sus1	532 - 662	13 to 14	12.1	11.9	
4. 1-4 VS 13-10	A vs D	Mxa5	54 - 147	1 to 3	Ckl1	512 - 604	13 to 15	11.8	11.9	
5. 5-8 vs 9-12	B vs C	Sco1	327 - 427	7 to 8	Mtu1	366 - 461	11 to 12	12.2	11.6	
3. 3-8 VS 9-12	B vs C	Sco1	320 - 435	6 to 8	Ath5	414 - 531	10 to 12	10.9	11.0	
6. 5-8 vs 13-16	B vs D	Osa28	315 - 421	6 to 8	Asu1	550 - 649	14 to 16	14.1	13.2	
	B vs D	Osa4	202 - 331	6 to 8	Msu1	494 - 621	14 to 16	12.3	13.2	
7. 9-12 vs 13-16	C vs D	Vvi4	370 - 470	9 to 11	Ath9	602 - 706	13 to 15	10.3	10.2	
	C vs D	Pgi1	385 - 469	10 to 11	Ani11	606 - 689	14 to 15	10.1	10.2	
8. 1-2 vs 3-4	A	Cim2	104 - 162	1 to 2	Acl1	176 - 236	3 to 4	9.1	8.9	
6. 1-2 VS 3-4	A	Cim2	118 - 162	1 to 2	Pgu9	248 - 292	3 to 4	8.7	0.9	
9. 5-6 vs 7-8	В	Nfi3	251 - 291	5	Yli4	411 - 450	7	11.5	11	
9. 3-0 VS 1-0	В	Ani11	210 - 260	5	Tko1	244 - 294	7	10.5	11	
10. 9-10 vs 11-12	C	Sus2	351 - 394	9 to 10	Cco1	388 - 431	11 to 12	8.6	8.6	
	C	Asu1	313 - 369	9 to 10	Pdi1	421 - 475	11 to 12	8.5	0.0	

Table 4. Comparison of different segments within OPT proteins using the GGSEARCH, HMMER and SAM programs. The format of presentation is the same as for Table 3.

Composison	Superfamily	Family;		Profile	D	atabase	GGSEARCH	HMMER	SAM		
Comparison	Supertaining	TC#	Protein-1 Acc#		Protein-2	Acc#	(e-value)	(e-value)	(e-value)		
1	OPT AB vs CD	2.A.67.3	Spr1	YP_001477255.1	Lsa1	YP_394932.1	1.7 e ⁻⁸	4.0 e ⁻⁴	0.1		
						YP_001477255.					
	OPT CD vs AB	2.A.67.4	Lsa1	YP_394932.1	Spr1	1	7.7 e ⁻⁷	0.004	0.004		
2	OPT A vs B	2.A.67.4	Ngo1	YP_208927.1	Sde1	YP_526125.1	5.8 e ⁻⁶	0.06	0.5		
2	OPT B vs A	2.A.67.4	Sde1	YP_526125.1	Ngo1	YP_208927.1	3.2 e ⁻⁵	0.2	0.09		
	OPT A vs C	2.A.67.2	Zma1	NP_001104952.1	Chy1	YP_361078.1	8.6 e ⁻⁶	0.03	0.002		
3						NP_001104952.					
	OPT C vs A	2.A.67.4	Chy1	YP_361078.1	Zma1	1	9.2 e ⁻⁶	0.03	0.02		
4	OPT A vs D	2.A.67.1	Gze4	XP_389463.1	Sus1	YP_822933.1	8.0 e ⁻⁴	0.09	2		
4	OPT D vs A	2.A.67.4	Sus1	YP_822933.1	Gze4	XP_389463.1	1.4 e ⁻⁴	0.03	0.2		
5	OPT B vs C	2.A.67.1	Sco1	AAF26618.1	Mtu1	NP_216911.1	3.6 e ⁻²	0.07	0.01		
	OPT C vs B	2.A.67.4	Mtu1	NP_216911.1	Sco1	AAF26618.1	1.9 e ⁻³	0.08	0.003		
						YP_001343430.					
6	OPT B vs D	2.A.67.2	Osa28	CAE02279.2	Asu1	1	3.9 e ⁻⁸	0.006	0.02		
	OPT D vs B	2.A.67.4	Asu1	YP_001343430.1	Osa28	CAE02279.2	3.7 e ⁻⁴	0.007	0.001		
7	OPT C vs D	2.A.67.4	Pgi1	NP_904744.1	Ani11	XP_658304.1	2.4 e ⁻⁴	0.2	2		
,	OPT D vs C	2.A.67.2	Ani11	XP_658304.1	Pgi1	NP_904744.1	2.0 e ⁻⁴	0.05	0.5		

Table 5. Comparison of four different programs for evaluating significance of sequence similarity. These programs are 1) IC/GAP (expressed in standard deviations), 2) GGSEARCH, 3) HMMER and 4) SAM (all three expressed in e-values). The superfamilies compared include the CDF/Orai superfamily (entries 1 and 2), the DMT superfamily (entries 3 and 4) and the BART superfamily (entries 5-7). With the IC/GAP score as the gold standard, GGSEARCH on the average proved better than HMMER which proved better than SAM.

Comparison	Superfamily	Family; TC#	Profile		Database		IC/GAP Score	GGSEARCH	HMMER	SAM
Comparison			Protein-1	Acc#	Protein-2	Acc#	(S.D.)	(e-value)	(e-value)	(e-value)
1	CDF vs Orai	2.A.4.1	PfuCDF	AAL80682	CelOrai	NP_497230	14	0.00034	0.0056	1.4
	Orai vs CDF	1.A.52.1	CelOrai	NP_497230	PfuCDF	AAL80682		5.4 e ⁻⁵	0.033	0.53
	CDF TMS 3-4	1	PfuCDF	AAL80682	CelOrai	NP_497230	- 14 -	0.018	0.018	0.59
	VS	2.A.4.1								
2	Orai TMS 1-2									
2	Orai TMS 1-2							0.00036	0.02	0.19
	vs	1.A.52.1	CelOrai	NP_497230	PfuCDF	AAL80682				
	CDF TMS 3-4									
3	DMT	2.A.7.20	PfCRT	Q86M68	AthCRT	Q8RWL5	16	0	0	0
3	DMT	2.A.7.20	AthCRT	Q8RWL5	PfCRT	Q86M68		0	3.5 e ⁻¹²⁵	1.3 e ⁻¹⁶⁵
4	DMT	2.A.7.12	SLC35A1	Q8BRW7	PfCRT	Q86M68	9	9.9 e ⁻¹⁰	9.9 e ⁻¹⁵	1.0 e ⁻⁵
4	DMT	2.A.7.20	PfCRT	Q86M68	SLC35A1	Q8BRW7		6.9 e ⁻⁹	7.3 e ⁻¹²	1.9 e ⁻⁴
5	BART	P-RFT; 2.A.87.2	YpaA	NP_390186	Ade1	YP_464235	9	0	4.3 e ⁻⁹	6.3 e ⁻⁴
3	BART	Acr3; 2.A.59.1	Ade1	YP_464235	YpaA	NP_390186		0	7.4 e ⁻¹⁶⁸	2.4 e ⁻¹³⁸
6	BART	SHK; 9.B.33	LytS	NP_847838	Rba2	NP_868846	8	5.7	0.02	0.89
	BART	UNK; 2.A.93	Rba2	NP_868846	LytS	NP_847838		-	none	0.38
7	BART	KPSH; 9.B.34	Dge1	YP_604037	Rba2	NP_868846	9	8.5 e ⁻²	0.006	0.49
	BART	UNK; 2.A.93	Rba2	NP_868846	Dge1	YP_604037		0.28	0.6	1.5

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