Integration of phenotypic metadata and protein similarity in Archaea using a spectral bipartitioning approach

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ABSTRACT
In order to simplify and meaningfully categorize large sets of protein sequence data, it is commonplace to cluster proteins based on the similarity of those sequences. However, it quickly becomes clear that the sequence flexibility allowed a given protein varies significantly among different protein families. The degree to which sequences are conserved not only differs for each protein family, but also is affected by the phylogenetic divergence of the source organisms. Clustering techniques that use similarity thresholds for protein families do not always allow for these variations and thus cannot be confidently used for applications such as automated annotation and phylogenetic profiling. In this work, we applied a spectral clustering approach (1,2) to cluster proteins in this network. This method clusters proteins based on the topology of the entire network rather than based on individual similarities between proteins. Since there is no need to apply arbitrary identity thresholds, spectral clustering is highly suitable for the study of families of proteins which are the result of evolutionary processes such as sequence conservation, sequence divergence and duplication (1).

INTRODUCTION
Clustering is a commonly used method for partitioning large data sets into meaningful groups, i.e. to categorize data by the characteristics they share. The distance from each data point in the set to every other data point is expressed by some measure—typically the amino acid sequence when clustering proteins. Similarity of sequence is then often used to infer some commonality among proteins, whether it be a common evolutionary function, shared domain structure or shared function.

Before proteins are clustered into groups of proteins that share sequence characteristics, it is helpful to view the relationship between proteins as a network where proteins are represented as nodes, and the similarity between proteins are represented as edges between nodes. In this work, we used a spectral clustering approach (1,2) to cluster proteins in this network. This method clusters proteins based on the topology of the entire network rather than on individual similarities between proteins. Since there is no need to apply arbitrary identity thresholds, spectral clustering is highly suitable for the study of families of proteins which are the result of evolutionary processes such as sequence conservation, sequence divergence and duplication (1).

Proteins can also be clustered at different levels; at the most basic level, we could cluster proteins that have only a general similarity, such as for instance a shared domain. These clusters would reflect the distribution of domains among families of proteins. On a deeper level, we could cluster proteins that are orthologous but do not share a function, and thereafter orthologous proteins of the same function. Additionally, we could also separate orthologous proteins by effects of species divergence. For instance, within a small phylogenetic space (e.g. within a genus), a cluster of orthologous proteins that share the same function can be difficult to separate according to phylogeny, while separation could be easier as the phylogenetic distance increases (e.g. across different phyla). The phylogenetic separation is not necessarily the last in order, since different protein families have varying sequence conservation. It is, for instance, conceivable that some proteins with strongly conserved sequences (e.g. cell machinery) would not be easily separable at the phylum level.

By applying a spectral clustering algorithm which successively divides clusters of proteins into two child clusters

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(bipartitioning), we here attempt to assemble a hierarchy of clusters which may reflect these various levels of clustering. This approach partitions clusters of proteins until no further topology-based partition is possible. As a result, different clusters may exhibit very different levels of protein identity depending on the degree of sequence conservation, and different clusters may reflect varying aspects of the evolutionary processes involved.

Taking the Archaea as an example, the phylogenetic distance separating the two main phyla (the Crenarchaeota and the Euryarchaeota) is greater than the distance between two families within the Crenarchaeota (e.g. the Thermoproteaceae and the Sulfolobaceae). For a typical group of orthologous proteins, their sequence similarity would be lower between the phyla than between the families. For clustering methods based on similarity thresholds (see ref. 1 for a discussion of methods), the sequence divergence between the phyla may obscure the similarities in the proteins. In contrast, a topology-based clustering method could group all the members of a protein family, such as the phosphokinases, then partition the crenarchaeal phosphokinases from their euryarchaeal counterparts, and perhaps even further partition the phosphokinases in the Thermoproteaceae from those of the Sulfolobaceae.

Using this spectral bipartitioning clustering approach, we created successive protein similarity clusters for the protein sequences from 53 archaeal genomes. We then assessed how well the clusters distinguished by this method correlated with phylogeny (3) and functional annotation. Likewise, by including phenotypic metadata with each protein, such as the metabolism and habitat of its source organism, metadata co-occurrence profiles analogous to the phylogenetic profiles could be generated, thus adding further value to the protein clustering.

We also developed an intuitive, graphical online tool (available at: http://coal.jgi-psf.org/) where these spectral bipartitioning clusters (SBCs) can be analyzed and explored further.

**MATERIALS AND METHODS**

Data were collected from IMG (4) for all 53 currently available sequenced archaeal genomes (34 Euryarchaeota, 15 Crenarchaeota, two Thaumarchaeota, one Korarchaeota and one Nanoarchaeota) and bidirectional best hits were calculated between the genomes for all protein sequences using BLAST (5) (blastp, default parameters and BLOSUM62 matrix). An initial low-end cutoff (the only hard cutoff in this study) was set at 30% protein identity and e-values < 10−6 as a reasonable limit of functional relevance. IMG proteins were assigned to COGs using reverse position-specific BLAST on position-specific scoring matrices provided by CDD (6).

**Clustering**

We have applied the spectral clustering procedure described previously (1,2) to the set of archaeal proteins. The proteins are represented as nodes in a connected undirected graph with edges that carry weights based on node-to-node similarity according to the protein identity. The clustering procedure is analogous to a random walk of a particle moving over the nodes of the graph. At each transition, the particle moves to an adjacent node with probabilities corresponding to the weights of the edges. The amount of time the particle spends in a given subgraph will determine whether this is indeed a cluster of its own or not.

Preliminary to clustering, our data were partitioned into disjoint block-matrices. Eigenvalues and eigenvectors were calculated using an ARPACK library specifically designed for large-scale symmetric eigenproblems where the eigenvectors of the largest eigenvalues are desired.

The largest eigenvalue will equal one since all block matrices are stochastic (i.e. row sums to one). The magnitude of the second largest eigenvalue will determine how fast the corresponding Markov chain will approach its stationary distribution (7). In this approach, if the second largest eigenvalue is >0.8, we further divide the block matrix into two new blocks using K-means clustering of eigenvectors 1 and 2. The eigenvalue cutoff of 0.8 therefore reflects the topology of the protein similarity graph, and not the actual similarity scores.

Metadata were added in the form of COGs (8), arCOGs (9), phenotype (10) and phylogeny (4). The phenotypic data were roughly categorized as habitat, metabolism, oxygen usage and preferred temperature range. For the habitat mappings, we intentionally used broad categories. For instance, ‘marine’ is a catch-all phrase for e.g. deep sea vents and seawater. While this may seem overly insensitive in some cases, it is arguably the most manageable form of grouping habitats.

**Consistency**

The calculated consistency scores (see below) for each SBC reflect the degree of homogeneity of each type of metadata for that cluster. For instance, a cluster is said to be fully consistent with respect to phylogeny if all its member proteins belong to the same phylogenetic group. Similarly, a cluster would be consistent with respect to its oxygen requirements if all member proteins belong to species that were annotated as ‘aerobic’.

Given that the consistency of an SBC (c) for a given metadata or phylogeny type (k) is inversely proportional to the degree of entropy (E), the consistency score $c_k$ can be calculated as follows:

$$c_k = 1 - E_k = \sum_{i=1...m} \frac{P(k_i) \ln(P(k_i))}{\ln(m)} + 1 \text{ for } m > 1$$

$$c_k = 1 \text{ for } m = 1$$

where $k$ is the metadata or phylogeny type (e.g. ‘phylum’), $i$ specifies each of the possible character states for $k$ (e.g. Crenarchaeota, Euryarchaeota), $P(k_i)$ is the proportion of SBC member proteins with character state $k_i$, and $m$ is the number of different character states observed in the SBC. Consistency scores are bounded by 0 and 1, with 1 being a fully consistent SBC. However, scoring was modified for the function metadata type when member proteins lacked annotation and thus could not be assigned.
We found that under these conditions the proportion of proteins, with five characters occurring at an equal rate. We simulated 10,000 SBCs of 10 proteins, with five characters occurring at an equal rate. We found that under these conditions the proportion of SBCs with a consistency score $c$ of $>0.7$ was very low ($10^{-4}$). When one character is more dominant, such as for phyla where 34 genomes are of the same character, it is far easier to find fully consistent SBCs by chance. Through simulation, we found that 43% of the SBCs were fully consistent ($c=1$) as compared to the observed 76%. However, in case the significance of the phylogenetic consistencies is questioned, it is easy to quality check the measure by studying the next level of phylogeny. At the class level, the expected frequency of fully consistent SBCs drops to 1.1% compared to the observed frequency 59%. For some phenotypic characters, false positives may still be a problem. For instance, 39% of SBCs are consistent with oxygen usage, whereas we expect 15% by chance. While fully consistent SBCs are significantly overrepresented, we still suggest that the protein annotation be consulted as a quality check to see if indeed the functions may be relevant to the oxygen usage of the organism.

**RESULTS**

**Clusters generated by spectral bipartitioning**

The cluster architecture produced by spectral bipartitioning is inherently hierarchical since the clusters generated at each level are, in turn, bipartitioned so long as they fulfill the specified topological requirements (see ‘Materials and Methods’ section). ‘Root’ clusters are at the top level of the hierarchy and are designated by integers (e.g., 0 or 1 or 2). Partitioning of root cluster 1, for example, would yield two subclusters assigned the identifiers 1.0 and 1.1. Likewise, further partitioning of 1.1 would yield 1.1.0 and 1.1.1, and so on. Clusters that do not have any subclusters (i.e. clusters that cannot be partitioned further) are termed ‘leaf’ clusters. In some cases, these designations overlap since a root SBC that cannot be partitioned is also a leaf SBC.

From a total number of 122,452 archaeal proteins, we selected those proteins with reciprocal best hits in other genomes, resulting in a sequence similarity matrix of 95,803 proteins. Thus, there are no SBCs composed of paralogs from a single species or proteins unique to one species, ensuring that correlations between protein similarity, function and metadata focus on proteins that are present in at least two species. From this matrix we identified 8,463 independent root SBCs with 11 ± 61 members. Successive bipartitioning of these root SBCs produced 10,247 additional SBCs for a total of 18,710. These 18,710 SBCs included 13,586 leaf SBCs (7 ± 10 members), some of which are also root SBCs and others which were generated by partitioning.

**Using consistency scores to assess correlations**

We assessed how well the protein clustering correlates with (i) the functional annotations of the member proteins; (ii) the phylogeny of the source organisms; and (iii) four lifestyle-related phenotypic traits of the source organisms. The phenotypic metadata for habitat, oxygen preference, temperature range and metabolism were extracted from the Genomes Online Database (GOLD) (10) for all 53 archaeal genomes (Supplementary Table S1). Consistency scores were calculated for all leaf SBCs (see ‘Materials and Methods’ section). Scores range from 0 to 1. A high score indicates that one character state dominates the cluster; likewise, a low score denotes that the cluster is heterogeneous with respect to that type of data. When considering temperature preference metadata, for example, a cluster with only hyperthermophilic members would be termed fully consistent and would have a consistency score of 1. When calculating consistency scores for the functional annotations, we ignored a cluster if any members lacked a COG assignment (i.e. we set the consistency score to 0).

Figure 1 shows the distribution of consistency scores for leaf SBCs with respect to functional annotations and all four types of phenotypic metadata. Clusters were highly consistent for functional annotation, mostly due to the relaxed similarity requirements for COG assignments. Leaf SBCs also tend to be consistent with respect to temperature preference, oxygen usage and metabolism, but significantly less so with respect to habitat. However, it is likely that most of the fully consistent SBCs would be composed of proteins from closely related species, therefore reflecting the phylogeny rather than the phenotype of organisms. Some of the most interesting insights stem from those clusters that include members from distant archaebacteria. For example, when investigating aerobicity, we would pay most attention to those few clusters where the member proteins reside in distantly related aerobic organisms, since this may indicate ‘lateral gene transfer’ (LGT) rather than a vertical inheritance.

Additional factors need to be borne in mind when interpreting consistency scores for the four metadata types that are lifestyle-related phenotypes (temperature preference, oxygen usage, metabolism and habitat). It can be difficult to distinguish the similarities due to phenotype from those resulting from phylogeny. Since, for example, only a small
percentage of proteins may be directly associated with thermal adaptation, one would expect that the majority of the consistent clusters found would be simply due to phylogeny. A leaf SBC could be consistent with respect to oxygen usage because the member proteins are all specifically related to the aerobic lifestyle, or, it could be consistent simply because the member proteins all reside in very closely related organisms, all of which happen to be aerobic. This situation is illustrated by SBCs 1267.0 and 1093, all members of which are aerobes. Although both clusters are fully consistent with respect to oxygen usage, the interpretation of their consistency differs. SBC 1267.0 is composed of three closely related Halobacteria, while SBC 1093 includes both thermoprotei (Crenarchaeota) and halobacteria (Euryarchaeota). Although the clustering of SBC 1093 correlates with oxygen usage, that for SBC 1267.0 appears to be an artifact reflecting predominantly inheritance effects.

Correlating clusters with protein functional annotations

Through subsequent partitioning of a root cluster, it is conceivable that the subclusters would at some point correspond to orthologous proteins, sharing a function. To test how well SBCs correlate with functional annotations, we used the Clusters of Orthologous Genes [COG, (8)] and the Archaeal COG [arCOG, (9)] assignments as benchmarks and calculated the consistency within SBCs. The vast majority of both root and leaf SBCs fall into one of two categories: either all members are assigned to the same COG, or one or more members lack COG assignments and thus the SBC is scored as having ‘zero’ COGs (Table 1). Similarly, the arCOGs (7) were created analogously to COGs, but based exclusively based on 41 Archaea. Due to the differences in methodology between SBCs and (ar)COGs, and since SBCs are based on similarity and not necessarily orthology, we do not expect the two methods to completely overlap. Thus, SBCs may, depending on level, be more specific or more general then (ar)COGs. For instance, the root cluster

| Table 1. The distribution of the number (N) of COGs, arCOGs and Pfams associated with individual SBCs |
|----------|----------|----------|----------|----------|----------|
| N        | Root SBC  | Leaf SBC |
|----------|----------|----------|----------|----------|----------|
| COG      | arCOG    | Pfam     | COG      | arCOG    | Pfam     |
|----------|----------|----------|----------|----------|----------|
| 0        | 4248     | 5144     | 4212     | 4984     | 6643     |
| 1        | 3682     | 2576     | 3373     | 7512     | 5920     |
| 2        | 376      | 466      | 616      | 881      | 857      |
| 3        | 100      | 110      | 156      | 157      | 121      |
| 4        | 24       | 59       | 52       | 34       | 32       |
| 5        | 14       | 32       | 21       | 15       | 9        |
| 6        | 8        | 16       | 11       | 1        | 3        |
| 7        | 2        | 14       | 6        | 2        | 0        |
| 8        | 2        | 9        | 3        | 0        | 1        |
| 9        | 5        | 10       | 3        | 0        | 0        |
| More     | 11       | 27       | 10       | 0        | 0        |

The members of most SBCs are all assigned to the same functional cluster (i.e. COG, arCOG, Pfam), but some SBCs contain members from, for instance, several COGs, and the members of ~4000 SBCs have no associated COGs. (Leaf SBCs also include those root SBCs that could not be partitioned.)

![Figure 1. Distribution of leaf cluster consistency scores for functional annotation and four phenotypic metadata types. A score of 1 indicates that all proteins in the cluster share the same character state (e.g., all belong to hyperthermophiles). A score of <1 means that the cluster includes members with different character states (e.g., some from hyperthermophiles and some from mesophiles). If any cluster member lacked a functional annotation, we defined that cluster's consistency score for function metadata as 0.](https://academic.oup.com/nar/article-abstract/37/7/2096/1018854/0)
corresponds to the initiation factor 2 subunit family [pfam01008, (11)], while subsequent child clusters distinguish the orthologous clusters COG1184 and COG0182. Further partitioning results in separating COG1184 based on differences in Halobacterial versus other archaeal versions of the protein.

Since COGs (and to a lesser extent arCOGs) are typically large collections of proteins, which may be only distantly related, one would expect leaf SBCs to be subsets of COGs. Although most leaf SBCs were found to be subsets of COGs, some do contain more than one COG. While root clusters might be expected to contain several COGs prior to partitioning, it is more surprising to find leaf clusters with multiple COGs or arCOGs. There are several explanations for this finding. In some instances, although the proteins in a leaf SBC are assigned to different COGs, they are so similar that the algorithm cannot partition the SBC further. This appears to be the case in SBC 1376 which contains two COGs: COG0555 (Na+/proline symporter) and COG4149 (Na+/panthothenate symporter). In this case, the SBC has only one arCOG, so the latter approach consolidates the functional classification and is consistent with protein similarities. Other times, two COGs within a leaf cluster have the same annotation, thus suggesting that these COGs could be merged. For instance, see SBC 113.0.0 where member COGs COG0668 and COG3264 are both annotated as small-conductance mechanosensitive channels. Another situation is illustrated by the finding of both COG1111 and COG1948 in SBC 222.0. Here, the presence of the ERCC4 domain in both helicases (COG1111) and nucleases (COG1948) blurs the distinction between them.

Sometimes, the resolution of the spectral bipartitioning algorithm partitions a single COG into two or more subclusters. For instance, root SBC 623 contains COG1958. Partitioning of that cluster revealed that there are two subclasses of COG1958 (arCOG00998), one of which is conserved in both phyla and another which is unique to the Thermoprotei within the Crenarchaeota. These observations suggest that duplication of the original gene in the Thermoprotei was followed by a functional adaptation within the Thermoprotei class.

In summary, the overall agreement between (ar)COGs and SBCs suggest that a spectral bipartitioning approach is able to not only cluster proteins based on effects of shared domains and phylogenetic divergence, but is also able to reconstruct clusters of orthologous proteins.

**Correlating leaf clusters with phenotypic metadata**

We found that 41%, 39% and 32% of leaf clusters are fully consistent with respect to thermal preference, oxygen usage and metabolism, respectively, but only 22% with respect to habitat. One reason for the lower value for habitat is that it is more difficult to classify habitats into non-overlapping groups. Also, a habitat assigned based on where an archaeon was observed in nature may not be its optimal habitat. Generally, one must also be aware of false positives (see ‘Materials and methods’ section), and the protein function annotation should always be consulted.

Of particular interest are those fully consistent SBCs that contain at least five members and include representatives of different phyla. These cases are quite rare; accounting for only 0.3% to 1.4% of the fully consistent clusters for the various metadata types. When interpreting these clusters, special care must be taken to distinguish between the effects of phenotype and phylogeny. For instance, a protein family unique to the thermoprotei will appear to correlate strongly with thermophilic temperature preference, but most likely this would be due to the recent divergence of these species. To identify protein families truly associated with thermophilic adaptation, one looks for clusters that span large phylogenetic distances, i.e. that have members from different phyla. By restricting further analyses to such phylogenetically inconsistent clusters, we would expect to identify the clusters with the strongest correlations with phenotype, but would likely miss others. In the following sections, we discuss some specific SBCs that are consistent with respect to a phenotypic metadata type and also are phylogenetically inconsistent at the phylum level, and offer possible interpretations of these observations.

**Thermal preference.** Most member proteins in these selected SBCs are annotated as ‘hypothetical’; those with functional annotations include proteins linked to RNA methyltransferase (SBC 8376.1.0.1.1.1.1.0.0.1.1.0.1). It has been shown (12) that some organisms react to increased temperatures by increasing methylation of RNA, so it is therefore not surprising to find this SBC consistent with a thermal preference. Furthermore, it has been reported that reverse gyrase is the only protein unique to hyperthermophiles (13,14). Reverse gyrase resides in the root SBC 1731 and its two subclusters: 1731.1 that contains only proteins from hyperthermophilic Crenarchaeota of the class Thermoprotei and 1731.0 that includes proteins from all three archaeal phyla. This finding is consistent with previous observations implicating LGT in the evolutionary history of reverse gyrase (14).

Two COGs (COG1857 and COG1688) are key components of a novel putative DNA repair system reported to be essential for thermophiles (15). COG1857 is found in two SBCs (1819.0 and 1819.1) whose members combined include 22 proteins from hyperthermophiles and four proteins from thermophiles. In contrast, COG1688 is found in four root SBCs, suggesting that it includes proteins that are quite dissimilar. Nevertheless, it too is mostly associated with hyperthermophiles. Thus, our results, based on a larger number of sequenced Archaea, support the earlier prediction linking these COGs to thermophiles.

**Oxygen usage.** SBC 1278.0.0 contains 10 members, all annotated as oxygen-sensitive ribonucleoside-triphosphate reductases, from 10 organisms representing three phyla. This enzyme is known to be involved in anaerobic growth in some bacteria (16). Another ribonucleoside-related SBC (1196.0) contains members from the two major phyla and occur only in anaerobic organisms, thus suggesting a correlation in this case with aerobic lifestyle.
Metabolism. Interestingly enough, the only phylogenetically diverse SBCs here are from chemoorganoheterotrophs. Since most chemoorganoheterotrophs are thermophiles, it is difficult to separate the effects of thermal preference and metabolism. For instance, SBC 4682 contains pyrrolidone-carboxylate peptidases, a protein that is adapted for greater thermal stability in hyperthermophiles (17) and that occurs equally among crenarchaea and euryarchaea. In general, however, there are several SBCs related to sugar transport (2078.0.0.1.0, 1924.1.0.0.1.0, 2078.0.1.0, 1924.1.0.0.0.1, 25.1.1.0.0.0.1.0.0, 2695), which is to be expected for heterotrophic archaea since they must transport metabolites.

Habitat. Only eight SBCs were selected here and the members of all eight come from organisms assigned to aquatic or marine habitats. The function of most of their member proteins was annotated as ‘hypothetical’, but one SBC contains transposases (mobile elements by definition) and another contains threonyl-tRNA synthetases that were likely acquired from bacteria (18). This lack of any obvious candidates for proteins required for a particular habitat could be due to any of several factors: the number of proteins unique to and essential for a particular habitat is low, the overwhelming majority of such proteins are transmitted predominantly by vertical inheritance, and/or the habitat classifications are broad and include many microhabitats.

Correlating leaf clusters with phylogeny
To assess how closely the clustering paralleled phylogeny, we calculated the consistency scores for all leaf clusters based on the taxonomic assignments of their members. Figure 2 shows the distribution of the scores obtained for each of five taxonomic levels: phylum, class, order, family and genus. The proportion of fully consistent clusters is greatest at the phylum level (76% or 10 317 of 13 586) and decreases with the decreasing phylogenetic distance, reaching a low of 19% (2556 of 13 586) at the genus level. Thus, as expected, fewer protein classes were found to be unique to a genus than to a higher level such as a class or phylum.

Further interpretations can be derived from the observed consistency scores at the phylum level ($c_P$). However, additional factors need to be considered when interpreting inconsistent clusters. First, when a cluster partitions into two phyla, it might be because the member proteins are unique to one phylum or the other. However, it might instead be because the sequence conservation is relaxed enough that the evolutionary divergence of the phyla becomes the main factor driving the partitioning. Second, strongly conserved SBCs, such as those containing housekeeping proteins, would tend to show low consistency scores due to their ubiquity among archaea. Our data show 3066 conserved leaf SBCs where $c_P$ is $\leq 0.8$ and 10 317 fully consistent SBCs ($c_P = 1$). Thus, about 24% of archaean proteins are conserved to a degree that overshadows the divergence of the archaean phyla, or have been transferred.

LGT has been observed in a variety of organisms, but appears to be especially common in archaea where 50% have acquired one or more protein domains from other organisms (19). Consistency scores can provide an indication of past LGT events. If an SBC acquired a sizeable portion of its members from another phylum, its $c_P$ will be low—in the same range as for highly conserved protein
families. However, if the transfer was recent and is reflected in only a very few members, consistency scores will be high (i.e. \(c_P\) will be close to 1). There are 203 such leaf SBCs in the data set, defined as clusters whose \(c_P\) is between 0.8 and 1. This is to be compared with 10 274 fully consistent SBCs (\(c_P = 1\)) and 2089 SBCs of conserved proteins (\(c_P \leq 0.8\)). Taken together, this suggests that \(\sim\)1% of archaeal proteins have been recently transferred between phyla. The rate of transfer is doubtless much higher, since this value represents only transfers between phyla. At lower taxonomic levels, the number of LGT-candidate SBCs increases. However, these may be false positives because proteins in closely related organisms can be similar enough to cluster together without any LGT being involved.

Possible LGT could also be detected by identifying SBCs that are consistent with respect to functional annotation but inconsistent with respect to phylogeny. A simple example of this would be finding a functionally consistent leaf SBC, which not only contains mostly euryarchaeal proteins but also has one or two members from crenarchaeal organisms. We provide a few examples of this, including their corresponding SBC identifiers and locus tags for individual proteins, in the following paragraphs.

The anaerobic Crenarchaeota *Thermofilum pendens* Hrk 5 is the likely recipient of several transferred proteins, including some from the Euryarchaeota. In SBC 1229.0, its FeoA family protein (Tpen_0974) is fully connected with eight euryarchaeal ferrous iron transport proteins from organisms that are mostly anaerobic methane-producers. *Thermofilum pendens* also has a geranylgeranyl reductase (Tpen_0718) gene which is difficult to separate from euryarchaeal orthologs in SBC 8420.1.1.0. Furthermore, SBC 870 suggests that it acquired a pyruvyl-dependent arginine decarboxylase (Tpen_0872) from anaerobes that share its hot spring environment (20). This is more intriguing since most crenarchaeota do not have a recognizable arginine decarboxylase. Other possible transfers into *T. pendens* include a Rubredoxin-type Fe(Cys)4 protein (Tpen_1457) and a UDP-N-acetylglycosamine 2-epimerase (Tpen_1715), again from euryarchaeal classes such as the Methanobacteria, Methanococci and Methanomicrobia (SBCs 8376.1.0.1.0.1.1.1.0.1.0.1.1.1.1.0.1 and 1255.1).

SBCs 796 and 53.0.1 show that the crenarchaeon *Caldivirga maquilingensis* IC-167 may have obtained genes coding for diaminopimelate decarboxylase (Cmaq_0523) and thiamine-phosphate pyrophosphorylase (Cmaq_0061) from the Euryarchaeota. Likewise, SBC 553.1.0.1.0.1 is composed entirely of adenine deaminases from anaerobic species, all from Euryarchaeota except for the adenine deaminase (Smar_0382) from the crenarchaeon *Staphylothermus marinus* F1.

In some cases, we instead found few Euryarchaeota among many Crenarchaeota. SBC 1306.1.1 contains crenarchaeal phosphohistidine phosphatases and one protein (AF1002) from the euryarchaeon *Archaeoglobus fulgidus* DSM 4304 which is annotated as hypothetical in IMG (4). All the taxa represented in this SBC are aquatic thermophiles or hyperthermophiles, thus suggesting that these organisms may at times share environments.

Frequent genetic input via LGT has been suggested for two groups of thermoacidophilic archaea: Sulfolobus and Thermoplasma (21). Since our sample includes three Sulfolobus and two Thermoplasma archaea, we would expect lower consistency scores (\(c_P \leq 0.8\)) to result from LGT in this case. Thus, contrary to what one might expect, rampant lateral transfer may be more difficult to detect. We found 30 SBCs with members from all five of these thermoacidophiles that also contain fewer than 10 proteins—thus avoiding ubiquitous conserved proteins. Although many of their members are of unknown function, they also include such diverse proteins as rieske-type iron-sulfur proteins, dihydrodipicolinate synthases and cytochrome-related proteins.

### Deep-branching archaea

*Candidatus Korarchaeum cryptofilum* (C. K. cryptofilum; 22) is an unculturable hyperthermophilic Archaeon isolated from hot springs and sediments, and is believed to represent a deep-branching phylum at the base of the two main archaeal phyla Crenarchaeota and Euryarchaeota. The gene content of *C. K. cryptofilum* is also largely hybrid, since it shares a majority of proteins with Crenarchaeota but also scattered components with Euryarchaeota; components often associated with replication/repair mechanisms. The authors suggest that these patterns could have arisen due to several lateral transfers. This can be difficult to determine by phylogenetic consistencies since we have only one representative of the Korarchaeota. For instance, *C. K. cryptofilum* shares a DNA polymerase II protein with euryarchaeal genomes (SBC 270), suggesting either a strong conservation or lateral transfer. In SBC 1067, lysiRNA is more distinct from the bulk of euryarchaeal lysyl-tRNA synthetases, therefore suggesting a vertical inheritance. A CO dehydrogenase maturation factor (SBC 4.0.1.0.1.0.1.0.1.0.1.0.1.1.1.0.1) also seems inseparable from its euryarchaeal counterparts, which may suggest lateral transfer. NADH dehydrogenase subunit A (SBC 31.1.0.1.0.1.1), however, clusters mainly with its crenarchaeal counterparts, although it is slightly more dissimilar to the others. Furthermore, tRNA pseudouridine synthase (SBC 413) belongs to a predominantly crenarchaeal cluster, albeit an outlier in the set.

The presence of *C. K. cryptofilum* in the data set does not seem to play the role of a natural breakpoint between SBCs, i.e. its proteins are not usually intermediates in homology between Crenarchaeota and Euryarchaeota. For instance, SBCs 4.0.0.0.0.1.0.1.0.1.0.1.0.1 can be subpartitioned, but not by virtue of proteins belonging to *C. K. cryptofilum*. Another example is SBC 1122, where the proteins that connect the two child clusters belong to a fused gene (Mbur_2244) from *Methanococoides burtonii* DSM 6242, along with two proteins from the Thaumarchaeota rather than belonging to *C. K. cryptofilum*. Thus, it seems as if the genetic composition of *C. K. cryptofilum* has been influenced by LGT and gene loss, as suggested by Elkins and co-workers (22) in the original paper.
Online tool

All SBCs are publicly available at: http://coal.jgi-psf.org. The search function provided enables you to locate clusters based on numerous parameters, including keywords, consistency scores, number of member proteins and associated phenotypic metadata. Selecting a cluster displays a plot of the sequence orthology of its member proteins, thus visually communicating their relatedness. Furthermore, the user can interactively color the plot to indicate the functional annotation, phylogenetic assignment, or phenotypic metadata associated with each member protein. For example, Figure 3 shows the display for SBC 4 colored to indicate the taxonomic class of the source organisms. For each SBC, the interface also provides a functional summary, a representative member sequence and convenient links to parent clusters and sub-clusters, where available. Proteins belonging to clusters can also be transferred to the IMG (4) gene cart for further analysis.

DISCUSSION

We performed a topology-based soft clustering of the proteins from 53 archaeal genomes and evaluated the suitability of this method in light of the great variation in both sequence conservation and divergence effects between protein families. Integrating functional annotations, phylogeny and associated phenotype with the sequence data allowed us to evaluate the influences of phylogeny and lifestyle on protein families. This methodology provides a valuable framework for biological data mining, of which we report some findings. Our clusters are publicly available for further exploration at: http://coal.jgi-psf.org/
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