Co-expression of statistically over-represented peptides in proteomes: a key to phylogeny?

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Abstract

It is proposed that the co-expression of statistically significant motifs among the sequences of a proteome is a phylogenetic trait. From the co-expression matrix of such motifs in a group of prokaryotic proteomes a suitable definition of a phylogenetic distance is introduced and the corresponding distance matrix between proteomes is constructed. From the distance matrix a phylogenetic tree is inferred, following a standard procedure. The inferred tree is compared with a reference tree deduced from a distance matrix obtained from the alignment of ribosomal RNA sequences. Our results are consistent with the hypothesis that biological evolution manifests itself with a modulation of basic correlations between shared peptides of short length, present in protein sequences. Moreover, the simple procedure we propose confirms that it is possible, sampling entire proteomes, to average the effects of lateral gene transfer and infer reasonable phylogenies.

Key words: Genomics, whole-proteome phylogeny, k-motifs, co-expression matrix.
Each living species is the result of its evolution; this historical assumption is the basic tenet of modern evolutionary biology. Molecular systematics [1] aims at classifying living species by measuring differences in their inherited molecular constituents, not in their phenotypic, macroscopic appearance. Since the classic paper by Zuckerkandl and Pauling [2], rational molecular systematics rests on the analysis of actual sequences, which are, though in an indirect way, the archive of the evolutionary information about biological species (taxa). Differences in nucleotide or amino acid sequences are the objective material elements to start from; that is particularly true in the classification of microscopic, unicellular organisms, where more macroscopic methods, based on ecological or pathological properties of these species, seem to be less fundamental [3]. Carl Woese, more than twenty years ago, has founded the universal molecular classification of living organisms based on ubiquitary co-evolved sequences, like e.g. the RNA of the small ribosomal subunit (SSU rRNA) [4]. The main achievement was the discovery of the fundamental tripartition of the tree-of-life into the branches of Bacteria, Archea and Eukarya [5]. Nowadays molecular phylogeny is a well established discipline, based on probabilistic methods [6]; nevertheless, the existence of lateral gene transfer [7][8] (i.e. mixing up of genes between species, particularly practiced among prokaryotes) has created some problems in the field and led some radical phylogenists to argue against the reliability of single-gene phylogenies [9]. To cope with this problem there has been, since more than ten years ago, a growth of ideas and methods to infer molecular phylogenies not from the analysis of groups of single genes, as classically done, but rather from the analysis of whole genomes and proteomes [10]. This is the subject also of the present work.

Proteomes are far from being a random assembly of peptides. Clustering of aminoacids [11], and strong correlations among genomic [12] and proteomic [13] segments have been clearly demonstrated. These results give meaning to the metaphor of protein sequences viewed as texts written in a still unknown language [14].

Following this view we assume that biological evolution could manifest itself, at the molecular level, through the modulation of significant
sequence elements that can be variously combined in the evolutionary
declination of the language embodied by the proteins of living organisms. It is reasonable to consider as significant those tracts of a protein sequence which exhibit a pronounced deviation from a random assembly of aminoacids. We have then looked for putatively significant elements as short peptide sequences of length $k$ which occur, in a proteome, a number of times larger than expected in a random proteome. In this work we show that these peptides are acted upon by natural selection and display, in different proteomes, statistical correlations able to express evolutionary distances.

A proteome $P$ is a collection of $n_P$ protein sequences, i.e. strings of various lengths made of symbols from an alphabet $A$ of 20 letters: $A = \{\sigma_1, \sigma_2, ..., \sigma_{20}\}$; each $\sigma$ labels one of the different aminoacids a protein is made of. If the proteome contains $N_P$ aminoacids (i.e. letters), then one can compute their relative frequencies: $f(\sigma_i) = n_i/N_P$, $n_i$ being the number of times the $i$-th aminoacid occurs in the proteome and $i = 1, 2, ..., 20$. We define as $k$-peptides sequences of $k$ contiguous letters. The number of all possible $k$-peptides is $20^k$. From the proteome $P$ we can only select $N_P - n_P \cdot (k - 1)$ overlapping $k$-peptides; some of them occur once, others more than once, doing so in the same or in different different proteins.

Denoting with $p_j^{(k)} = \{\sigma_{1j}, \sigma_{2j}, ..., \sigma_{kj}\}$ the $j$-th $k$-peptide we can count the number $N_j^{(o)}$ of times it occurs in the actual proteome. We can also estimate the expected number of occurrences $N_j^{(e)}$ of the $j$-th $k$-peptide in a random proteome (of the same length and with the same number of $k$-peptides), generated by independent random extractions of letters with the constraint of producing, on the average, prescribed relative frequencies $f(\sigma_i)$. That is:

$$N_j^{(e)} = [(N_P - n_P \cdot (k - 1)) \cdot \Pr[p_j^{(k)}]],$$

(1)

where $\Pr[p_j^{(k)}]$, the probability of occurrence of the $j$-th $k$-peptide, can be estimated as $f(\sigma_{1j}) \cdot f(\sigma_{2j}) \cdot ... \cdot f(\sigma_{kj})$, i.e. as the product of the relative frequencies of its component letters in the actual proteome.

For each $k$-peptide of expected occurrence $N^{(e)}$, the probability that
it is observed $N$ times in a random proteome (with the same amino acidic composition and sequences of the same lengths as in the actual proteome) is given by a Poissonian distribution:

$$Pr_{N(o)}[N] = \frac{[N^{(e)}]^{N}}{N!} \cdot \exp[-N^{(e)}]. \quad (2)$$

We define as statistically-relevant the over-expressed $k$-peptides whose observed number of occurrences $N^{(o)}$ is such that:

$$\int_{0}^{N^{(o)}} Pr_{N^{(e)}}[N']dN' \geq 0.95 \quad (3)$$

(i.e. the observed occurrence of the $k$-peptide falls in the upper five percent tail of its Poissonian distribution). Let us call hereafter $k$-motifs the over-expressed $k$-peptides, selected following inequality (3). Analogously, we have defined a test-set of $k$-peptides which, differently from $k$-motifs, are expressed as expected (i.e. $N^{(o)} \approx N^{(e)}$). We call these peptides expected $k$-peptides. $k$-motifs and expected $k$-peptides are differently distributed along protein sequences: $k$-motifs are seldom alone in a protein and in many cases they partly overlap forming longer, potentially significant, tracts. They occur at specific distances one from the other, whereas expected $k$-peptides are isolated and dispersed, without any recurrent clustering. As an example, in fig.1 we show the occurrence of $k$-motifs in an archaeal protein.

The non trivial statistical properties of the $k$-motifs suggest that, among the $k$-peptides present in a proteome, they could display patterns of correlated expression useful to derive phylogenetic distances between taxa.

We have considered eighteen proteomes from the Gene Bank [15]. Ten from Archaea: Aeropyrum pernix, Archaeoglobus fulgidus, Halobacterium spNRC1, Methanococcus jannaschii, Methanobacterium thermautotrophicum, Pyrococcus aerophilum, Pyrococcus abyssi, Pyrococcus furiosus, Sulfolobus solfataricus, Thermoplasma acidophilum and eight from Bacteria: Agrobacterium tumefaciens, Bacillus subtilis, Chlorobium tepidum, Deinococcus radiodurans, Escherichia coli K12, Synechocystis spPCC6803, Thermotoga maritima, Yersinia pestis CO92
We have selected \(k\)-motifs from all these proteomes and collected them into \(k\)-dictionaries. Let \(Z_n(k)\) be the subset of the \(k\)-dictionary composed by those \(k\)-motifs which are expressed in, \textit{at least}, \(n\) different proteomes. \(Z_1(k)\) is thus the entire set of \(k\)-motifs, referring to the considered group of proteomes. \(Z_2(k)\) is the set of \(k\)-motifs common to \textit{at least} two proteomes; \(Z_1(k) - Z_2(k)\) is, therefore, the subset of the \(k\)-motifs specific to one proteome. Fig. 2 reports, as a function of \(k\), the number of entries in different \(k\)-dictionaries, normalized over the total number of expressed \(k\)-peptides (\(Z_1(k)\)). It is worth noting that, as \(k\) increases, the proteome-specific \(k\)-motifs (\(Z_1(k) - Z_2(k)\)) rapidly overwhelm the shared \(k\)-motifs (i.e. \(Z_2(k)\)). The \(Z_2(k)\) dictionary (open circles in fig.2) has a significant number of entries for low and intermediate \(k\) values, as it contains almost 10\% of the expressed peptides, for \(k = 6\).

We define now the \textit{co-expression} matrix of the set \(Z_n(k)\) in a proteome \(P\) the matrix \(A^{(P)}[Z_n(k)]\); its element \(ij\) counts the number of times \(i\)-th and \(j\)-th \(k\)-motifs, from \(Z_n(k)\), occur together in one of the proteins of \(P\). This matrix resembles the adjacency matrix of the network formed by linking words when they occur in the same phrase in texts written in natural languages, as done in recent linguistic studies\[16\]. In a subsequent more extended paper we shall present the statistical properties of the linguistic co-expression networks built on sets of \(k\)-motifs\[17\].

The pattern of co-expression matrices based on \(Z_n(k)\), for a given value of \(k\), is far from trivial in all the considered proteomes, with many groups of \(k\)-motifs co-expressed in one or more proteins up to several tens of times. On the other hand we have noticed that co-expression matrices generated by equally-populated sets of expected \(k\)-peptides are sparse, with just very few and tiny elements different from zero.

The different co-expression patterns of \(k\)-motifs in different proteomes are the basis of the method we propose in this letter.

The observations reported above might be resumed as follows: (1) there is a consistent set of \(k\)-motifs which are common among the considered organisms of a given kingdom; this set might constitute a sort of basic dictionary collecting robust pieces of information, stable
across the taxa; (2) there is a larger set of proteome-specific $k$-motifs $[Z_1(k) - Z_2(k)]$, whose evolution occurred within a specific taxon and might be considered as the manifestation of a linguistic specificity of that species; (3) there is a consistent set of $k$-motifs, $Z_2(k)$, containing the common $k$-motifs together with a number of $k$-motifs which are quite specific but, nevertheless, common to a few species.

It is reasonable to assume that common and proteome-specific $k$-motifs somehow interact: the usage of proteome-specific terms might influence the usage of the common $k$-motifs, in the sense that the co-expression of the latter might be modulated by usage of the former, giving origin to a specific co-expression pattern.

Let us propose now a definition of the phylogenetic distance among proteomes. From $A^{(P)}[Z_n(k)]$, the symmetric co-expression matrix of a given proteome $P$, we can extract a co-expression vector $V^{(P)}[Z_n(k)]$, whose components are the $n_k(n_k+1)/2$ distinct entries of the matrix ($n_k$ is the number of $k$-motifs in $Z_n(k)$), ordered in an arbitrary but fixed way, e.g. by rows:

$$V^{(P)}[Z_n(k)] = A^{(P)}_{ij} \left[(Z_n(k))\right]$$ (4)

with $j \geq i$ and $s$ ranging from one to $n_k(n_k+1)/2$. We consider the co-expression vector as a linguistic fingerprint of a proteome expressing its peculiar use of both common and proteome-specific motifs. We define a phylogenetic distance $d_{P,P''}(k)$ between two proteomes $P'$ and $P''$ through the scalar product of their co-expression vectors based on a $Z_j(k)$ dictionary:

$$d_{P,P''}(j,k) = 1 - \frac{\sum_s \{V^{(P')}_s[Z_n(k)] \cdot V^{(P'')}_s[Z_n(k)]\}}{\{|V^{(P')}| \cdot |V^{(P'')}|\}}$$ (5)

In this work we have evaluated phylogenetic distances among a set of prokaryotic proteomes, using the $Z_2(6)$ dictionary. There are several arguments to motivate this choice for the probe-set of motifs. The $Z_2(6)$ dictionary of the set of prokaryotes we are considering has 7712 entries; it contains a balanced mixture of common and proteome-specific tracts. The use of a $Z_j(6)$ dictionary, with $j > 2$ would have produced a distance evaluation on the only basis of strongly conserved motifs, i.e.
those common to a large number of organisms, disregarding the mod-
ulation effect that they could produce on the proteome-specific tracts.
These dictionaries, moreover, are quite small (the \( Z_6(6) \) dictionary, for
instance, contains only 55 entries) and their size markedly decreases
with the increase of both \( j \) and \( k \). On the other side, \( Z_2(k) \) dictionar-
ies with \( k < 6 \) are made by a large number of motifs (e. g. \( Z_2(5) \) has
161903 entries), but the potential increase in sensitivity, putting aside
practical considerations due to the treatment of large matrices, would
be spoiled by some volatility of low \( k \) motifs. Due to the rigidity of the
statistical criterion (3) (same acceptance threshold for all the motifs)
one \( k \)-motif which has passed the test and belongs to the \( k \)-dictionary
could pass the test for the \( k + 1 \) dictionary as part of one of the 40
\((k + 1)\)-peptides which can be obtained by adding one letter at its be-
beginning or at its end. We have observed that this is rarely the case for
low \( k \). So, if \( k \) is too low then many short peptides, which are accepted
as statistically significant and could be the nucleus of biologically rel-
levant tracts of sequence, are lost and not recognized in the \( k + 1 \) test.
When \( k \) is larger than 5, the motifs have been seen to be more stable,
in that they generally appear as part of longer motifs also in the \( k + 1 \)
dictionary. Indeed, some of them are also ”lost” but this can be the
sign of the ”end” of the specific tract. The \( Z_2(6) \) dictionary is thus a
good trade-off between number of entries and balance of common and
proteome-specific tracts. Moreover, \( k = 6 \) seem to be a peculiar length
for peptides: it has been proven that 6-peptides allow a unique recon-
struction of a protein sequence from the collection of its constituent
\( k \)-peptides 19.

By using the definition of eq.(5) we have evaluated all the distances
between the considered set of proteomes. The resulting distance ma-
trix has been processed by the neighbor-joining method 20 using the
PHYLIP package 21. The dendrogram we have obtained through the
procedure outlined above is shown in Fig. 3. In Fig. 4 we show the tree
obtained, for the same set of taxa, from the server of the Ribosomal
Database Project 22. This last phylogeny can be assumed as a refer-
ence, because it is based on the alignment of sequences of RNA from the
Small Ribosomal Subunit. This molecule is ubiquitary and coevolved to
accommodate a well-defined set of ribosomal proteins, hardly subject to lateral gene transfer. The tree from alignment of the SSUrRNA shows the clear separation of the two kingdoms: Archaea and Bacteria, this separation appear less clearly resolved in our tree whose center seems to be an archaeal spot from which emerge *Sulfolobus solfataricus*, a branch of 5 bacteria, a group of Archaea with the bacterium *Deinococcus radiodurans* (D. radi) among them, and a group of Archaea with the bacteria *Chlorobium tepidum* (C.tepi) and *Synechocystis* (Synech) segregated among them. One could be discouraged by this result and think that the method we are proposing is unable to resolve the basic tripartition of the tree of life and that we are mistakenly classifying taxa. One could argue, from a different perspective, that the kind of method we are proposing, based on global statistical properties of the proteomes, is able to reveal phylogenetic associations which are at variance with the fundamental SSUrRNA classification. The stability of the method and its biological foundations have to be further investigated. However it is worth noting that, quite surprisingly, the tree we have reconstructed through a biologically blind criterion mutated from statistical linguistics can be reasonably compared with those obtained through refined and deep whole-genome analyses [23][24]. In particular we believe that whole-genome phylogenies of the kind we are proposing should be confronted with very recent observations suggesting that eukaryotes could originate from the fusion of pre-existing prokaryotic genomes [25]. Moreover, the important distinction between operational and informational genes [26] suggests that we are looking at a possible different statistics of occurrence of the $k$-motifs, which are the probe of our method, over the two kinds of proteins; we also believe that blind approaches based on the statistics of short sequence motifs, as the one we present here, could be less affected by different sources of bias which are however present in statistical phylogenomic studies based on the clustering of entire genes [27].

In the last stage of the preparation of this manuscript we became aware of an important study which uses an approach very close to ours [28] and which has been made available on line [29]. In that method a proteome is also sampled for statistically significant 6-peptides; the
background constituted by an uncorrelated random extraction of letters is subtracted. The fingerprint vector of each proteome has $20^6$ components, each one of them expresses the statistical deviation of the occurrence of each peptide from that expected in a random proteome. In our approach the fingerprint is represented instead by the co-expression vector. Following the method proposed in [29] we have derived a distance matrix of the same 18 species here investigated; the phylogenetic tree we have obtained has a more resolved dichotomy between Archaea and Bacteria, and a topology which, though more consistent is still less resolved and definitely not coincident with that of a tree obtained from the distance matrix based on the alignment of the SSU rRNA [22]. It will be interesting to proceed, in the next future, to a careful assessment of the biological information which can be derived from the two approaches. At present we tend to have the following view: the phylogenetic picture based on the tree of life has been put under scrutiny by the large extent of lateral gene transfer between taxa; that challenged phylogenists, using properly selected groups of genes, to reveal evolutionary relations which are not consistent with the universal tree of life. Recently there have been claims for the tree of life to fuse into what has been called the *ring of life* [25].

Phylogenies based on whole genomes are coherent with the view of the three kingdoms Archaea, Bacteria and Eukarya as originating from a world based on gene exchange and fusion of genomes. In particular, testing entire proteomes against patterns of correlated expression of statistically significant sequence motifs seems to be a proper way to cope with the original genome fusion regime and with the *mean field* generated by lateral gene transfers, gene duplication and lost. The method proposed in [28], samples in a more generic way the evolutionary correlations between $k$-motifs and seem to force the trees toward the tree of life shape. Our method, based on patterns of co-expression of $k$-peptides could be more in agreement with the view of a fusion-based ring of life. Of course, quantitative comparison between different methods is now really required, we are planning an extensive quantitative investigation of the relative merits of different approaches in reconstructing the philogeny(ies) of a properly selected set of taxa. In
doing that a clear mathematical setting is of tantamount importance [30][31].

The scientific content of phylogenies that are based on the statistical sampling of entire proteomes and that avoid sequence alignment algorithms has still to be validated. Nevertheless we believe that they can have a practical relevance at least as tools for the rapid molecular classification of the ever increasing number of freshly sequenced genomes.

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>gi|14520241|ref|NP_125715.1| hypothetical MALTOSE/MALTODEXTRIN TRANSPORT ATP-BINDING [Pyrococcus abyssi]

| MVEVRLENLKFGNFTA| VLLGPSGCGKTTTLRMIALEEE | PT | EFL | VLLGPSGCGKTTTLRMIALEEE |
|------------------|--------------------------|----|-----|--------------------------|
| GKIYFDREVTYLPREE | SMVFQSYAVWPHTMYDNIAPFLK | IKKFPDEIDKRV | RWAAGLQIEELD | PAQLSGGQRQRVA | VARAIVVEPD | VLLMDEPLSNDLAKLRVA | MRAEIKLQQLKVTTEYVTHDQVEATMGMDDRIAVMN | RGQLLQVGPTEVYLPNSV | FVATFIGAPEMNIVEVSVDGYLEGKFGIILTLPQDIMELLREDYIGKTVLFGIRPEHM | TVEGVSelahMKTAKLNAVDFVEALGIDTDILHVKGDELKVKLPGHIPIEVGKE | VTVIYLDNMMHVFDKDEKAI

Figure 1: A typical dispersion of 6-motifs (bold) in a protein of the archaeon *P. abyssi*. In the upper part of the figure are reported the 55 6-motifs, belonging to $Z_2(6)$, which are expressed in the protein. Note the clustering and overlap of the motifs.
Figure 2: Relative fraction of $k$-motifs present in the different subsets of the $k$-dictionary: $Z_1(k) - Z_2(k)$ (black circles), $Z_2(k)$ (white circles), $Z_6(k)$ (squares).
Figure 3: Unrooted phylogenetic tree of the considered proteomes: the full name of the species can be easily reconstructed from the abbreviations. After a / the kingdom is indicated: A stands for Archaea, B for Bacteria.
Figure 4: SSUrRNA phylogeny of the 18 species here considered; from [22].