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# Peptide Vocabulary Analysis Reveals Ultra-Conservation and Homonymity in Protein Sequences 

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#### Abstract

A new algorithm is presented for vocabulary analysis (word detection) in texts of human origin. It performs at $60 \%-70 \%$ overall accuracy and greater than $80 \%$ accuracy for longer words, and approximately $85 \%$ sensitivity on Alice in Wonderland, a considerable improvement on previous methods. When applied to protein sequences, it detects short sequences analogous to words in human texts, i.e. intolerant to changes in spelling (mutation), and relatively contextindependent in their meaning (function). Some of these are homonyms of up to 7 amino acids, which can assume different structures in different proteins. Others are ultra-conserved stretches of up to 18 amino acids within proteins of less than $40 \%$ overall identity, reflecting extreme constraint or convergent evolution. Different species are found to have qualitatively different major peptide vocabularies, e.g. some are dominated by large gene families, while others are rich in simple repeats or dominated by internally repetitive proteins. This suggests the possibility of a peptide vocabulary signature, analogous to genome signatures in DNA. Homonyms may be useful in detecting convergent evolution and positive selection in protein evolution. Ultra-conserved words may be useful in identifying structures intolerant to substitution over long periods of evolutionary time.


Keywords: peptide vocabulary, vocabulary analysis, word detection, motif, protein structure, bioinformatics, gene families, genome signature, peptide conservation, peptide homonymity

## Introduction

First used at least as early as the beginning of the 1970s, the concept of "the language of the genes" has become a recurring explanatory tool in popular presentations of molecular genetics (Chargaff, 1971; Jones, 1993). Genomes may be compared to libraries of genetic information, with each chromosome as a book, genes as chapters, and DNA bases as the letters in which the text is written (Ridley, 1999). In principle, the linguistic analogy may be applied equally to protein sequences as to DNA, simply by increasing the alphabet from 4 to 20 letters. The prevalence, and utility, of this metaphor in undergraduate teaching and the popular science media, obscures a deeper controversy concerning its genuine applicability in research (Searls, 1993; Ji, 1999; Searls, 2002; Sakakibara, 2005). Attempts have been made to apply generative grammar structures to gene organization in bacteria (Collado-Vides, 1991, 1992, 1996), DNA-protein interaction (Bentolila, 1996; Wang et al. 2005), the problem of gene prediction (Dong and Searls, 1994; Muggleton et al. 2001), protein folding (Gimona, 2006) and RNA structure prediction (Matsui et al. 2004). These efforts in molecular biology are in the tradition of wider attempts to create formal grammars, or to use the grammatical metaphor, for other kinds of biological data (Gutfreund, 1976; Jerne, 1985; Hamilton, 1993; Wang, 2004). A related metaphor is that of genome sequence as a code to be deciphered by the molecular biologist, who thus becomes a "biomolecular cryptologist" (Konopka, 1994; Bodnar et al. 1997). Conversely, techniques developed in molecular biology are now being recycled back into cryptography (Spencer et al. 2004).

Under the terms of these general analogies, short sequences of DNA may be regarded as words. Often, any $k$-mer is referred to as a word (Mantegna et al. 1994; Chatzidimitriou-Dreismann et al. 1996) but here these will be designated strings. Where a string has some local functional significance in a sequence and consequently has been conserved throughout the evolutionary process, it may be referred to as a motif (Waterman, 1989; Hu et al. 2000). Identification of motifs is usually based on large-scale comparative analysis and alignment of related sequences.

Counts of DNA string frequency have been used as a means of differentiating classes of DNA sequence, such as exons, introns and promoters (Beckmann et al. 1986; Solovyev and Lawrence, 1993;

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Solovyev et al. 1994b, 1994a; Bains, 1997; Frontali and Pizzi, 1999; Bultrini et al. 2003), although the meaning of such differences in terms of the linguistic metaphor of the genome has been disputed (Konopka and Martindale, 1995; Chatzidimitriou-Dreismann et al. 1996; Martindale and Konopka, 1996; Tsonis et al. 1997). String counts, after correction for underlying base composition, have been assembled into vectors known as genome signatures, reflecting their apparent distinctiveness between genomes (Karlin and Mrázek, 1997; Karlin et al. 1997; Karlin, 1998; Karlin et al. 1998; Campbell et al. 1999). Such composition-corrected string frequency vectors have proved useful in detecting horizontal gene transfer events between species of bacteria (Karlin, 2001). A further development based on genome signatures is that of compositional spectra, designed to reduce vector size and increase technical tractability (Bolshoy, 2003; Kirzhner et al. 2003).

This paper investigates the meaning of the linguistic metaphor in more detail in protein sequences, with particular emphasis on the identification of words. A protein word, rather than a string, is here taken to be more literally comparable to a word within a text of human origin. Therefore, words are only a subset of strings. Likewise, a word differs from a motif, in that motifs are often fuzzy (meaning tolerant to substitution) and are best viewed in the context of an alignment of related sequences. Within a text of human origin, a word has some contextindependence. It has clear boundaries and may appear flanked by very different text in different cases. Fuzziness is also not tolerated; a word has a correct spelling. The total assembly of detected words is referred to as the vocabulary, and the word detection process as vocabulary analysis.

The pioneering vocabulary analysis in DNA sequences was carried out by Brendel et al. (1986). Their metric was based on contrasting frequencies of substrings within the candidate word. For a string, $s$, of length $k$, its expected occurrence, $E$, is the product of the occurrences of its left and right substrings, divided by the occurrence of its internal substring.

$$
\begin{aligned}
E\left(s_{1} \ldots s_{k}\right)= & f\left(s_{1} \ldots s_{k-1}\right) * f\left(s_{2} \ldots s_{k}\right) \\
& 1 f\left(s_{2} \ldots s_{k-1}\right)
\end{aligned}
$$

For each string, $s$, the difference between its expected occurrence, $E(s)$ as calculated above, and actual occurrence, $f(s)$, is quantified by:

$$
\operatorname{std}(s)=(f(s)-E(s)) / \max \{\sqrt{E(s)}, 1\}
$$

This provides a $z$-score for the actual occurrence of the string. Brendel et al. (1986) define a contrast word as any string where $\operatorname{std}(s) \geq 3$. Brendel et al. (1986) were able to identify several contrast words of lengths $k=3$ to 6 in the genomes of $E$. coli and two coliphages. Conversely, avoided words could also be detected, where $\operatorname{std}(s) \leq-3$. An essentially similar metric has been implemented by others (Phillips et al. 1987a, 1987b; Merkl et al. 1992; Colosimo et al. 1993; Castrignanò et al. 1997; Rocha et al. 1998; Apostolico et al. 2003).

In principle, this method could also be applied to detect contrast words in protein sequences, but the combinatorial explosion caused by the presence of a 20 -letter code in proteins as opposed to the 4-letter code in DNA, has restricted work on string frequency in proteins to $k=2$ (i.e. dipeptides) only (Solovyev and Makarova, 1993). Application of the contrast words method to human texts was extended by Schmitt et al. (1996). Analysing Alice in Wonderland, they found that it performed relatively poorly, essentially due to the fact that the 26-letter alphabet of a text in English has a string combinatorial explosion problem even worse than that of 20-letter protein sequences.

This paper proposes improvements on the contrast words method, initially comparing their performance, in the tradition of Schmitt et al. (1996), on Alice in Wonderland. The most accurate method for identifying true words is then applied to several other human texts, to the NRL3D set of proteins of solved structure, and to the proteome sets of several species from all three superkingdoms (NCBI Taxonomy Browser classification) of cellular organisms.

The concept of synonymity is familiar in molecular biology. Within the degenerate genetic code, many amino acids may be encoded by more than one codon. A protein sequence may therefore be potentially coded by a combinatorially vast number of synonymous DNA sequences. Here the term is used in a more general sense. When two protein strings have different sequences, but perform the same function in their respective proteins, they are said to be functionally synonymous. Fuzzy motifs
are an example of functional synonymity within protein families. The converse concept, that of homonymity, has not been explored (although see Lennon and Nussinov, 1984). Where a non-fuzzy word occurs in two different proteins and performs a different function in each, that peptide word is functionally homonymous. At a trivial level, it is immediately possible to see that the longer a peptide, the less likelihood it has of functional homonymity. The questions of the longest existing homonymous word, the prevalence of peptide homonymity, and its origins are all explored in this paper.

## Methods

## Texts and protein sequence sources

Public domain texts were downloaded from Project Gutenberg (http://www.gutenberg.org). Punctuation, non-alphabetic characters, numbers and spaces were removed. Word counts were caseinsensitive.

The NRL3D set of sequences of proteins of solved structure (Pattabiraman et al. 1990) was downloaded from the University of Hong Kong (http://bioinfo.hku.hk/db/nrl_3d/NRL3D/nrl 3d.seq). Non-contiguous sequences (those annotated as "fragments"), sequences containing ambiguities and exact duplicates were removed using a Perl script. This reduces the number of sequences from 23301 to 6168 . Further trimmings were performed using CD-HIT (Li and Godzik, 2006), which can produce datasets with maximum degrees of pairwise identity. Such reduced sets are subsequently referred to as NRL3D $n n$, where $n n$ is the maximum pairwise identity. The justification for this trimming is that most words will occur in closely related sequences, and will consequently be explicable at a trivial level. Trimming with CD-HIT reduces the number of words detected and maximises the likelihood that they will be found in less closely related proteins, and thereby be potentially more interesting from a functional point of view. As a negative control, trimmed NRL3D data sets were shuffled using shuffleseq (http://emboss.sourceforge.net/apps/ release/4.0/emboss/apps/shuffleseq.html) from EMBOSS (Rice et al. 2000).

Proteomes (meaning predicted protein sets derived from genome projects) were downloaded from the EBI Integr8 database (http://www.ebi. ac.uk/integr8). They were similarly reduced by CD-HIT.

## Vocabulary analysis algorithms

For each text or proteome, and for NRL3D, overlapping strings of all lengths from $k=1$ to 20 were counted using a Perl script running the BioPerl (Stajich et al. 2002) SeqWords module (http://doc.bioperl.org/releases/bioperl-current/ bioperl-live/Bio/Tools/SeqWords.html). The SeqWords output was then analysed in the following ways. Each metric is given an acronym for easier reference.

1) CW: Contrast words method (see Introduction)
This is the method of Brendel et al. (1986). The difference is that the $\operatorname{std}(s)$ threshold was set at 0.1 to maximise the number of candidate words.

## 2) RS: Raw strings

The simplest possible method: all strings of length $k \geq 5$, occurring at $n \geq 20$, were assessed as candidate words.

## 3) ES: Equal substrings

The raw strings extracted as above were trimmed to include only those having equal occurrences of left and right substrings.

$$
f\left(s_{1} \ldots s_{k-1}\right)=f\left(s_{2} \ldots s_{k}\right)
$$

The rationale for this approach is that many true words tend to satisfy this criterion. For instance, in Alice in Wonderland, the true word ALICE is revealed by:

$$
f(A L I C)=f(L I C E)
$$

following to the fact that Alice in Wonderland, despite referring to several species, does not mention lice.
4) CW-ESM: Equal substrings of middle substring of contrast words
Combining methods 1 and 3 , middle substrings were extracted from contrast words with $\operatorname{std}(s)>=0.1$. These were then examined for equal substrings:

$$
f\left(s_{2} \ldots s_{k-2}\right)=f\left(s_{3} \ldots s_{k-1}\right)
$$

The rationale for this approach is the ad hoc empirical observation that false positive contrast words, of which there are many (Schmitt et al. 1996), frequently have true words embedded within them as middle substrings.
5) RS-ESM: Equal substrings of middle substring of raw strings
Combining methods 2 and 3 , since equality of substrings within the middle strings of contrast words was frequently found to be an indicator of a true word, the same was applied to raw strings. The additional proviso was that the left and right substrings of the raw string were not of equal occurrence to each other or the middle substring.

$$
f\left(s_{2} \ldots s_{k-2}\right)=f\left(s_{3} \ldots s_{k-1}\right)
$$

and

$$
f\left(s_{1} \ldots s_{k-1}\right) \neq f\left(s_{2} \ldots s_{k}\right)
$$

and

$$
f\left(s_{1} \ldots s_{k-1}\right) \neq f\left(s_{2} \ldots s_{k-1}\right)
$$

and

$$
f\left(s_{1} \ldots s_{k}\right) \neq f\left(s_{2} \ldots s_{k-1}\right)
$$

The rationale for this was that, for instance, within the raw string DALICET, the true word ALICE is revealed by:

$$
f(A L I C)=f(L I C E)
$$

and

$$
f(D A L I C E) \neq f(\text { ALICET })
$$

and

$$
f(D A L I C E) \neq f(\text { ALICE })
$$

and

$$
f(A L I C E T) \neq f(A L I C E)
$$

CW-ESM and RS-ESM are equivalent, excepting that CW-ESM takes contrast words as its starting point, and RS-ESM uses raw strings. In both cases the candidate word is the middle substring, should it satisfy the criteria given.

## Measurement of accuracy

In human texts it is possible to score true words among the detected candidate words. Accuracy is measured using the Sen2 statistic (Milanesi and Rogozin, 1998):

$$
\operatorname{Sen} 2=T P /(T P+F P)
$$

where $T P$ are those candidate words identified as true positives, and $F P$ are those identified as false positives.

Perl scripts are available on request from the author.

## Assessment of hits

Protein domains were determined by reference to Pfam (http://www.sanger.ac.uk/Software/PfamFinn et al. 2006) and Prosite motifs detected using ScanProsite (http://www.expasy.ch/tools/scan-prosite-de Castro et al. 2006). Alignments were performed using ClustalW (Chenna et al. 2003) or bl2seq (http://www.ncbi.nlm.nih.gov/bl2seq/ wblast2.cgi-Tatusova and Madden, 1999).

## Structural visualization

Solved proteins structures were downloaded from PDB (http://www.pdb.org) and visualization was carried out in MOE (http://www.chemcomp. com).

## Results

## Vocabulary analysis in human texts

Alice in Wonderland is a short novel of 26587 words. The total vocabulary is 2593 different words,

Table 1. Commonest 10 words in Alice in Wonderland, sorted by their occurrence, $n$.

| Word | $n$ |
| :--- | ---: |
| THE | 1631 |
| AND | 865 |
| TO | 728 |
| A | 628 |
| SHE | 541 |
| IT | 530 |
| OF | 512 |
| SAID | 462 |
| I | 410 |
| ALICE | 386 |

of which 1475 are used more than once and 1072 more than twice. For illustrative purposes, the 10 commonest words are shown in Table 1. As might be expected, these are all small prepositions and pronouns, except for the name "Alice" which has 386 occurrences and is the 10th commonest word, and the verb past tense "said" at 462 occurrences.

The words in Table 1 are derived from a spaced text, with only punctuation and other extraneous characters removed. Spaces were then removed for testing of the various metrics. Again for illustrative purposes the top 10 hits using each method are shown (Tables 2 to 6), but the final comparison was made using all the hits for each method (Table 7).

1) $R S$ metric

The commonest raw strings in Alice in Wonderland of length $k=5$ to 20 are tabulated in Table 2.
Only 3 of the commonest raw strings in Table 2 are true discrete words or phrases (DWoPsshaded grey). "Alice" as a raw string has a slightly higher occurrence than the word "Alice" in a spaced text ( 397 vs. 386 - see Table 1) as it also occurs as part of the possessive "Alice's". As Table 2 suggests, RS is a relatively poor metric for identifying true words. Almost all of the raw strings in Table 2 are components of the single DWoP "said the".
2) CW metric

CW (Brendel et al. 1986) performs equally poorly, as previously demonstrated by Schmitt et al. (1996). Table 3 shows the top 10 contrast words of length $k=7$ to 20 , sorted by descending $s t d(s)$. There are only two DWoPs detected.

Table 2. 10 commonest raw strings of $k=5$ to 20 in unspaced Alice in Wonderland. True discrete words or phrases (DWoPs) are shaded.

| Word | $n$ |
| :--- | ---: |
| ALICE | 397 |
| SAIDT | 266 |
| AIDTH | 224 |
| SAIDTH | 222 |
| IDTHE | 221 |
| SAIDTHE | 212 |
| AIDTHE | 212 |
| ANDTH | 169 |
| THING | 169 |
| DALICE | 162 |

Table 3. Top 10 contrast words of $k=7$ to 20 in unspaced Alice in Wonderland, sorted by std(s), their $z$-score. True DWoPs are shaded. $k$ : length of contrast word, $n$ : occurrence, $n-L$ : occurrence of left substring, $n-R$ : occurrence of right substring, $n-M$ : occurrence of middle substring, std(s): z-score (see Introduction).

| Word | $\boldsymbol{k}$ | $\boldsymbol{n}$ | $\boldsymbol{n}$ - $\boldsymbol{L}$ | $\boldsymbol{n}$ - $\boldsymbol{R}$ | $\boldsymbol{n}$ - $\boldsymbol{M}$ | $\boldsymbol{s t d}(\boldsymbol{s} \boldsymbol{)}$ |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: |
| ROUGHTH | 7 | 11 | 14 | 13 | 114 | 7.44 |
| TOTHINK | 7 | 7 | 7 | 7 | 43 | 5.49 |
| TOFTHEW | 7 | 10 | 14 | 25 | 156 | 5.18 |
| OINTHED | 7 | 9 | 21 | 10 | 109 | 5.10 |
| AIDNOTH | 7 | 6 | 6 | 6 | 34 | 4.80 |
| POFTHEE | 7 | 5 | 7 | 6 | 156 | 4.73 |
| DIDTHEY | 7 | 5 | 9 | 8 | 221 | 4.67 |
| THECOUR | 7 | 16 | 16 | 18 | 52 | 4.45 |
| ESAIDTO | 7 | 26 | 40 | 77 | 266 | 4.24 |
| RLITTLET | 8 | 7 | 15 | 14 | 128 | 4.18 |

It was noted that the some of the false positive contrast words in Table 3 contained the true DWoPs "of the" (twice), "in the" and "little" as their middle substrings. This stimulated the further investigation of the middle subwords.

## 3) ES metric

Table 4 tabulates the 10 highest hits with ES, sorted by their occurrence, $n$. This contains 6 true DWoPs (shaded).

Table 4. Top 10 hits for ES of $k=5$ to 20, sorted by occurrence. True DWoPs are shaded. $k$ : length of raw string, $n$ : occurrence, $n-L$ : occurrence of left substring, $n-R$ : occurrence of right substring, $n-M$ : occurrence of middle substring.

| Word | $\boldsymbol{k}$ | $\boldsymbol{n}$ | $\boldsymbol{n}-\boldsymbol{R}$ | $\boldsymbol{n - L}$ | $\boldsymbol{n - M}$ |
| :--- | ---: | ---: | ---: | ---: | ---: |
| ALICE | 5 | 397 | 397 | 397 | 401 |
| LITTLE | 6 | 128 | 128 | 128 | 128 |
| LITTL | 5 | 128 | 128 | 128 | 193 |
| SAIDALICE | 9 | 116 | 116 | 116 | 116 |
| SAIDALIC | 8 | 116 | 116 | 116 | 116 |
| THOUGH | 6 | 91 | 91 | 91 | 91 |
| HERSELF | 7 | 83 | 83 | 83 | 83 |
| THEQUE | 6 | 77 | 77 | 77 | 78 |
| THEKING | 7 | 62 | 62 | 62 | 62 |
| HEKING | 6 | 62 | 62 | 62 | 64 |

Table 5. The top 10 hits with RS-ESM of $k=5$ to 18 , sorted by their occurrence, $n$. True DWoPs are shaded. $k$ : length of raw string, $n$ : occurrence.

| Word | $\boldsymbol{k}$ | $\boldsymbol{n}$ |
| :--- | :--- | ---: |
| ALICE | 5 | 397 |
| OFTHE | 5 | 156 |
| LITTLE | 6 | 128 |
| SAIDALICE | 9 | 116 |
| LIKE | 4 | 97 |
| THOUGH | 6 | 91 |
| HERSELF | 7 | 83 |
| THEQUE | 6 | 77 |
| THEKING | 7 | 62 |
| TURTLE | 6 | 61 |

ES performs rather better than CW or RS, although it can accumulate nested strings. For instance in Table 4, "saidalic" is found to be a substring of "saidalice", "littl" of "little" and "heking" of "theking". This suggested the combination of ES with the other methods.

## 4) RS-ESM metric

RS-ESM shows a further marked improvement. Nested substrings are avoided, and 9 out of the top 10 hits are true positives (Table 5).

Table 6. Top 10 hits using CW-ESM of $k=5$ to 18 , sorted by std(s), their z-score. True DWoPs are shaded. $k$ : length, $n$ : occurrence of the contrast word in which they are embedded, std(s): z-score.

| Word | $\boldsymbol{k}$ | $\boldsymbol{n}$ | $\boldsymbol{s t d}(\boldsymbol{s})$ |
| :--- | ---: | ---: | ---: |
| OFTHE | 5 | 68 | 5.18 |
| LITTLE | 6 | 44 | 4.18 |
| ALICE | 5 | 198 | 3.21 |
| SHOULD | 6 | 14 | 3.20 |
| THEMARCHHARE | 12 | 4 | 3.17 |
| THEDORMOUSE | 11 | 9 | 3.09 |
| BEGIN | 5 | 11 | 2.63 |
| WHICH | 5 | 8 | 2.51 |
| MINUTE | 6 | 21 | 2.50 |
| VENTURE | 7 | 10 | 2.50 |

Table 7. Comparison of the methods described above. TP: True positive DWoPs detected. Sen2: accuracy (see Methods).

| Method | Hits | TP | Sen2 |
| :--- | ---: | ---: | ---: |
| $R S-E S M, k=2-18, n \geq 2$ | 1312 | 895 | 0.682 |
| CW-ESM, $k=4-18, n \geq 2$ | 673 | 388 | 0.577 |
| $E S, k=5-20, n \geq 20$ | 241 | 61 | 0.253 |
| $R S, k=3-20, n \geq 10$ | 2293 | 540 | 0.235 |
| $R S, k=5-20, n \geq 20$ | 1213 | 206 | 0.170 |
| $C W, k=7-20, n \geq 4$ | 1927 | 117 | 0.061 |

## 5) CW-ESM

CW-ESM appears to be the best method on first examination. All of the top 10 hits are true DWoPs (Table 6). However, a decision on the best method to apply to biological sequences requires a fuller assessment of the output beyond the top 10 hits.

## Comparison of methods

Table 7 compares the methods on Alice in Wonderland. Since the initial string count was to $k=20$, the two ESM methods are limited to $k=18$ as their longest identifiable word.

Table 7 demonstrates that the CW metric is the poorest. Although it generates a large number of hits, the true positive rate is barely more than $6 \%$. RS gives greater numbers of candidate words as the thresholds for occurrence and string length are dropped, but Sen 2 does not rise above $24 \%$. Adding a requirement for equal right and left substrings, ES, brings the number of candidates down dramati-cally-from 1213 hits to 241 hits where $k=5-20$, $n \geq 20$. Sen 2 increases from $17 \%$ to just over $25 \%$. However, for the combination methods, Sen 2 increases considerably. For contrast words (CW-ESM) just under $58 \%$ accuracy can be achieved, and just over $68 \%$ accuracy for raw strings (RS-ESM). The latter also has a larger number of hits, generating 895 true positives. Considering only 1042 words are used more than twice in Alice in Wonderland, this is a reasonable figure.

The next question to be investigated is whether or not the quality of hits varies across $k$. Figure 1 plots the true positive rate against the length of the candidate word for RS-ESM in Alice in Wonderland. Sen 2 increases with length $k$. Although the overall Sen2 is 0.682 (Table 7), Sen 2 rises above 0.8 for $k=11-15$. The majority of strings of length $k=4$ and 5 are false positives.


Figure 1. RS-ESM performed on Alice in Wonderland. Number of hits plotted against $k$. TP: true positives. Sen2 is also plotted ( $\times 100$ ) to show its improvement at higher levels of $k$.

Extending the analysis to a range of other human texts, Figure 2 plots the number of candidate DWoPs detected for each against the length of the text in 1000s of characters (kchar). There is a clear correlation ( $r=0.994$ ) between number of different words and size of text. This has also been observed for the number of different raw strings, a phenomenon known as Heaps' Law (Heaps, 1978).

## Vocabulary analysis in sets of real and shuffled protein sequences

Figure 1 suggests that there may be increased artefactual detection of false positive DWoPs for RS-ESM at $k=4$ and 5 , based on the identification of such false positives in Alice in Wonderland. Greater than $60 \%$ true positivity is only obtained at $k \geq 7$ and $80 \%$ at $k \geq 11$. When a text of human origin is being analysed, one can reliably identify the false positives and thus precisely quantify Sen2. However, in a protein sequence set, whether NRL3D or a naturally occurring proteome, scoring of accuracy requires the use of shuffled sequences. In the shuffled sequences, all hits are by definition artefactual. Figure 3 plots the distribution of candidate words in real and shuffled NRL3D_63 protein sequence set (see Methods) for both $\overline{\mathrm{R}} \mathrm{S}$ ESM and CW-ESM methods. It can be seen that
the shuffled sequence sets give false positives at up to $k=6$ for RS-ESM. However, the ratio of hits of $k=6$ in the real as compared to the shuffled genome is much higher than at $k=5$ or less. Therefore, it seems that $k=6$ should be considered an ambiguous category. Although most hits at $k=6$ are likely to be genuine, there is a far greater risk of a false positive than at $k \geq 7$. The observation that $\operatorname{Sen} 2$ is less than 0.5 for $k \leq 5$ (Fig. 1) also justifies concentration on longer candidate words. This supports the earlier finding by Thode et al. (1996), who found that matches of 6 residues within a window of length 10 , could be found far more frequently between pairs of real proteins than random sequences. By contrast, CWESM, although producing fewer hits, has no hits in shuffled sequences above $k=5$. Therefore, it might be preferred for investigating words of length $k=6$.

## Structural meaning of words

The words of $k=12-18$ identified in NRL3D_63 using RS-ESM are shown in Table 8.

The protein family in which the word is located is designated from the NRL3D annotation, or where that is ambiguous, by reference to Pfam
kword

50

40

30


Figure 2. Number of candidate DWoPs ( $\times 1000$, kword) plotted against length of text (kchar) for RS-ESM, $k=6-18$. Alice: Alice in Wonderland, Gulliver: Gulliver's Travels, Oliver: Oliver Twist, Chaucer: Canterbury Tales in 19th century translation, Origin: Origin of Species, Don Quixote: 19th century English translation of same, KJB: King James Bible.


Figure 3. NRL3D_63 (solid lines) and its shuffled equivalent (dotted lines), tested with both RS-ESM (squares) and CW-ESM (triangles). The logarithm of the number of hits, $n$, is plotted against $k$. $\log (0)$ is arbitrarily designated zero. Pseudocounts are therefore added when $n=1$.

Table 8. Words of length $k=12-18$ in NRL3D_63, using RS-ESM. The protein family is derived from the NRL3D annotation.

| Word | Length | Protein family | No. of proteins |
| :--- | :---: | :--- | ---: |
| TCNVAHPASSTKVDKKI | 17 | immunoglobulin | 3 |
| LLQLTVWGIKQLQAR | 15 | gp41 | 3 |
| DATDRCCFVHDCCY | 14 | phospholipase | 5 |
| EKPYKCPECGKSFS | 14 | zinc finger domain | 1 (internal repeat) |
| LGGTCVNVGCVPKK | 14 | 2 kinds of reductase | 3 |
| LGRSGYTVHVQCNA | 14 | viral coat protein | 3 |
| TLGNSTITTQEAAN | 14 | viral coat protein | 3 |
| AFLGIPFAEPPVG | 13 | lipase/acetylcholinesterase | 3 |
| LGNGGLGRLAACF | 13 | phosphorylase | 3 |
| LLRISLLLIQSWL | 13 | growth hormone | 3 |
| TTPPSVYPLAPGS | 13 | immunoglobulin | 3 |
| AVLPGDGIGPEV | 12 | dehydrogenase | 3 |
| CLNVGCIPSKAL | 12 | dehydrogenase | 3 |
| FDTGSSNLWVPS | 12 | pepsin | 3 |
| HVQCNASKFHQG | 12 | viral coat protein | 3 |
| LRKAMKGLGTDE | 12 | annexin | 3 |
| PKDATDRCCFVH | 12 | phospholipase | 4 |
| QSQIVSFYFKLF | 12 | interferon | 4 |
| SDGIMVARGDLG | 12 | pyruvate kinase | 3 |
| SHVSTGGGASLE | 12 | phosphoglycerate kinase | 3 |
| SNASCTTNCLAP | 12 | phosphatase | 3 |

(Finn et al. 2006). Most of these hits are found within fairly well conserved proteins, often orthologues having the same essential function in different species within the same major phylogenetic class. In some cases, however, the hits are found to be stretches of total conservation within otherwise somewhat divergent proteins, often having slightly variant functions and from rather more distant species. The 14-mer LGGTCVNVGCVPKK
is found in glutathione reductase (EC 1.6.4.2) from humans and E. coli, and in the related enzyme trypanothione reductase (EC 1.6.4.8) from two genera of trypanosome.

Although LGGTCVNVGCVPKK is completely conserved within an alignment having generally poor levels of conservation (Fig. 4), spanning bacteria, trypanosomes and humans, all these proteins possess a pyridine nucleotide-disulphide

| 1 GRT | YLVIGGGSGGLESAWRAAELG-ARAAVV 4 |
| :---: | :---: |
| 1GER B | -MTKHYDYIAIGGGSGGIASINRAAMYG-QKCALI 33 |
| 1BZL A | -MSKIFDLVVIGAGSGGLEAAWNAATLYKKRVAVI 34 |
| 1 1TYP A | -MSRAYDLVVIGAGSGGLEAGWNAASLHKKRVAVI 34 |
|  | :* : .**.****: : .** : *: : |
| 1GRT | ESH---------KLGGTCVNVGCVPKKVMWNTAVHSEFMHDH--ADYGFP 88 |
| 1GER B | EAK---------ELGGTCVNVGCVPKKVMWHAAQIREAIHMYG-PDYGFD 73 |
| 1BZL A | DVQMVHGPPFFSALGGTCVNVGCVPKKLMVTGAQYMEHLRESAGFGWEFD 84 |
| 1TYP A | DLQKHHGPPHYAALGGTCVNVGCVPKKLMVTGANYMDTIRESAGFGWELD 84 |
|  | : : **************:* * : |
| 1GRT | SCEGKFNWRVIKEKRDAYVSRLNAIYQNNLTKSH-IEIIRGHAAFTSDPK 137 |
| 1 GER B | TTINKFNWETLIASRTAYIDRIHTSYENVLGKNN-VDVIKGFARFVD--A 120 |
| 1BZL A | RTTLRAEWKNLIAVKDEAVLNINKSYDEMFRDTEGLEFFLGWGSLESKNV 134 |
| 1 1TYP A | RESVRPNWKALIAAKNKAVSGINDSYEGMFADTEGLTFHQGWGALQDNHT 134 |
|  | : :*. : |

Figure 4. The first 150 residues of the alignment of the four sequences containing the word LGGTCVNVGCVPKK (shaded). The proteins are identified by their PDB designations-1GRT: human glutathione reductase; 1GER: E. coli glutathione reductase; 1BZL: Trypanosoma cruzi trypanothione reductase; 1TYP: Crithidia fasciculata trypanothione reductase.


Figure 5. Superposition of sequence LGGTCVNVGCVPKK in the 4 proteins aligned in Figure 4. The helical backbone is shown in black. Despite the variability of the other parts of these proteins, LGGTCVNVGCVPKK represents a region of extreme structural, and presumably functional, conservation between E. coli, humans and trypanosomes.
oxidoreductase domain (Pfam PF07992). LGGTCVNVGCVPKK is also recognised by ScanProsite (de Castro et al. 2006) as containing a pyridine_redox_1 motif (ProSite PS00076). The word therefore may be taken to have equivalent function within these proteins and is not a homonym as defined in the Introduction. LGGTCVNVGCVPKK in all four cases is found at the beginning of a long helix. Superposition of the structures of the 4 proteins to $1.894 \AA$ in MOE demonstrates excellent conservation even over the poorly conserved regions. LGGTCVNVGCVPKK assumes a highly similar structure in all cases (Fig. 5).

As an additional example, AFLGIPFAEPPVG is found in the N -terminal regions of acetylcholinesterase (EC 3.1.1.7) from mouse and the electric ray and also in triacylglycerol lipase (EC 3.1.1.3) from yeast. As before, the word represents a stretch
of total conservation in an otherwise low identity alignment (Fig. 6). Despite this, all 3 of these proteins contain a carboxylesterase domain (Pfam PF00135), and their solved structures may be superposed over their full length to $3.70 \AA$ in MOE (not shown).

One phenomenon that appears in the output, that has no analogue in texts of human origin, is the detection of homopolymers. The longest homopolymeric word in NRL3D_63 is the heptamer AAAAAAA, detected in antifreeze protein A from the flounder and also in an amine dehydrogenase from Thiobacillus versutus. However, it occurs in the extreme C -terminus and N -terminus respectively of these two proteins. Homopolymers are a consequence of regions of low complexity within coding DNA, and have no analogue within human texts. They formally constitute words, and

| 1EVE | DHSELLVNTKSGKVMGTRVPVLSSHISAFLGIPFAEPPVGNMRFRRPEPKKPWSGVWNAS |
| :---: | :---: |
| 1MAAD | EDPQLLVRVRGGQLRGIRLKAPGGPVSAFLGIPFAEPPVGSRRFMPPEPKRPWSGVLDAT |
| 1LPM |  |
| 1EVE | TYPNNCQQ----------YVDEQFPGFSGSEMWNPNREMSEDCLYLNIWVPS-PRPKS-T |
| 1MAAD | TFQNVCYQ---------YVDTLYPGFEGTEMWNPNRELSEDCLYLNVWTPY-PRPASPT |
| 1LPM | SYGPSCMQQNPEGTYEENLPKAALDLVMQSKVFEAVSPSSEDCLTINVVRPPGTKAGANL |
| 1EVE | TVMVWIYGGGFYSGSS---TLDVYNGKYLAYTEEVVLVSLSYRVGAFGFLALHGSQ-EAP |
| 1MAAD | PVLIWIYGGGFYSGAA---SLDVYDGRFLAQVEGAVLVSMNYRVGTFGFLALPGSR-EAP |
| 1LPM | PVMLWIFGGGFEVGGTSTFPPAQMITKS IAMGKPI IHVSVNYRVSSWGFLAGDEIKAEGS |
|  |  |

Figure 6. The first 150 residues of the alignment of the three sequences containing the word AFLGIPFAEPPVG (shaded). 1EVE: Torpedo californica acetylcholinesterase; 1MAAD: mouse acetylcholinesterase chain D; 1LPM: Candida rugosa lipase.
indeed in the case of AAAAAAA a homonym, within the terms of the algorithms used, but are neglected owing to their lack of likely functional significance.

Leaving aside the homopolymers, there is only one identifiable homonym in NRL3D_63 of $k \geq 7$. SLGDRVT is found in a beta-lactamase from Streptomyces albus and also in two mouse antibody proteins (1JRHL and 1NMCL). The two mouse proteins are $61 \%$ identical as assessed by bl2seq (Tatusova and Madden, 1999), and SLGDRVT is found in both cases in the N -terminus of the solved structure of the protein, where it is part of the V-set domain (Pfam PF07686) The two mouse proteins superpose to $0.821 \AA$ over the entire length of their solved structures (not shown), and their SLGDRVT sequences have good structural alignment of their backbones (Fig. 7).

In the eubacterial lactamase 1BSG, SLGDRVT is found in a different conformation (Fig. 8). In this protein is it part of a helix rich beta-lactamase domain, but does not occur within a helix.

SLGDRVT is the only homonym detectable in NRL3D_63 at $k=7$ using RS-ESM. Although there
are many at $k=6(37$ with CW-ESM and 36 with RS-ESM). As shown in Figure 3, CW-ESM may be preferable to RS-ESM at $k=6$ in that, although less sensitive, it is less inclined to false positives at $k=6$.

In summary, within NRL3D_63, longer words are mostly indicative of conservation. Some of them are islands of ultra-conservation within distinctly divergent proteins. However, annotation or Pfam domain mapping indicates that these are always, at least in the cases examined (both above and data not shown), within proteins of similar general functionality. The longest homomym is a solitary example found at $k=7$ but they appear to be plentiful at $k=6$. The latter however, must be under suspicion of false positivity, owing to the number of hits at $k=6$ in shuffled versions of the NRL3D database. The relative paucity of homonyms of reliable length suggests that future fine-tuning of the algorithm ought to be performed on protein sequence sets where functional annotation of motifs and domains is more complete than in NRL3D.

Since NRL3D is a compendium of proteins of highly diverse origin, but also enriched for sequences of easily solved structure, its vocabulary may be very different in character to that of


Figure 7. Superposition of SLGDRVT in mouse antibody proteins 1JRHL (white) and 1NMCL (black). Backbone traces are rendered as fine lines.


Figure 8. Comparison of SLGDRVT word in Streptomyces albus lactamase 1BSG (white) and mouse antibody protein 1NMCL (black). Backbone traces are rendered as fine lines.
individual proteomes. These were therefore examined for the presence of homonyms and island of extreme conservation.

## Vocabulary analysis on individual proteomes

Figure 9 plots the number of words detected using RS-ESM versus the size of the proteome in terms of number of proteins. All proteomes were previously reduced to no more than $63 \%$ identity by use of CD-HIT, as performed on NLR3D. Figure 9 indicates that Heaps' Law (see Fig. 2 above) also applies to proteomes. This had previously been observed for raw strings in proteins (Mukhopadhyay et al. 2006). The same trend applies when the proteomes are measured in kilo-residues (comparison not shown).

Figure 9 shows the same general relationship for proteomes as is demonstrated in Figure 2 for texts. The correlation is weaker for eukaryotes (not shown in Fig. 9) and archaea ( $r=0.905$ and 0.907 respectively), but comparable for eubacteria ( $r=0.996$ against $r=0.994$ for texts). However, the range of proteome size in eukaryotes is generally not
comparable with the other two superkingdoms, making it difficult to draw any conclusions concerning differences in vocabulary structure between superkingdoms. Supplementary Material Tables 1, 2 and 3 give the full results for the various species.

Table 9 shows that texts of human origin have a far richer vocabulary than proteomes, and that eukaryotes appear to have a richer vocabulary than eubacteria or archaea. However, when only eukaryotic proteomes within the size range of the other two kingdoms are considered, this discrepancy decreases markedly, suggesting that it should be interpreted with caution.

Detailed analysis of all proteome sets would be inappropriate for a single paper. A number of individual proteomes were chosen for further analysis, contrasting the three superkingdoms, and also small and large proteomes where possible.

## Vocabulary analysis in a small eubacterial proteome

Chlamydia muridarum has 916 proteins, of which 914 are no more than $63 \%$ identical, indicating a virtual absence of gene families of closely related


Figure 9. Candidate words of $k=6$ to 18 , detected using RS-ESM, against number of proteins for 35 eubacterial (black circles) and 28 archaeal (grey circles) proteomes.
proteins. Using RS-ESM, C. muridarum contains 34 words of which 17 are $k \geq 7$ (Supplementary Material Table 4). One of these is the homoheptamer DDDDDDD, and 7 others are words that
occur several times within single proteins, indicating repetitive sequences, or occurring in low complexity areas of proteins. Of the remainder, all fall within clearly related proteins, except for two.

Table 9. Summary of RS-ESM results on human texts and phylogenetic kingdoms. "euk. (eub. range)" refers to the eukaryotic proteomes that are within the same size range (in kilo-residues) as the eubacterial proteomes. Likewise, "euk. (arc. range)" refers to those within the same size range as the archaeal proteomes. For the human texts, "number of species" refers to number of texts, and average proteome size to average text size (in kilo-characters).

| Superkingdom | Number <br> of species | Total <br> proteins | av. proteome <br> size (kres) | av. protein <br> len. (res) | Words/kres | Words/prot |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| eukarya | 36 | 382698 | 4902 | 461 | 1.615 | 0.745 |
| euk. (eub. range) | 7 | 15205 | 1025 | 472 | 0.957 | 0.452 |
| euk. (arc. range) | 4 | 3459 | 374 | 432 | 0.424 | 0.183 |
| eubacteria | 35 | 104006 | 947 | 319 | 0.670 | 0.214 |
| archaea | 28 | 65197 | 681 | 292 | 0.665 | 0.194 |
| human texts | 9 | N/A | 1322 | N/A | 13.939 | N/A |

The first of these, GPTGSGK appears at first glance to constitute a homonym, occurring in 4 proteins (SwissProt identifiers Q9PLF7, Q9PLM1, Q9PJG9 and Q9PKD0) with different Pfam domains, although all are ATP-binding proteins. In each case GPTGSGK is found at or near the N-terminus of the main Pfam domain within the protein (being IPPT,AAA_2, GSPII_E and ABC_tran respectively, listed as "various" in Supplementary Material Table 4). In the case of Q9PKD0, GPTGSGK is annotated by ScanProsite as an NP BIND ATP binding motif. Therefore, GPTGSGK is probably not a homonym but rather an ATP-binding cassette conserved or convergently evolved across divergent proteins within C. muridarum.

A second word of interest is an ultra-conserved region within a set of rather divergent transporter proteins, where it constitutes, again like GPTGSGK, the NP_BIND motif for ATP-binding (Fig. 10). GPNGAGGKSTL and GPTGSGK can be represented by the profile GP(T/N)G(A/S)GK.

Of the 17 words in $C$. muridarum of $k=6$, all but 4 appear to be homonyms. However, these must be regarded with suspicion, as they can occur artefactually at $k=6$ in shuffled sequences (Fig. 3). When $C$. muridarum is examined with the less sensitive CWESM algorithm, which is also less liable to artefactual hits at $k=6$, there are only 6 hits at that length, of which only one is a homonym.

## Vocabulary analysis in a large archaeal proteome

Methanosarcina acetivorans has 4467 proteins, of which 4080 are no more than $63 \%$ identical, indicating that around $10 \%$ of the total is comprised
of members of moderately or closely related protein families, in contrast to the virtual absence of such families in C. muridarum. In order to make the analysis more tractable, the M. acetivorans proteome is first trimmed to $40 \%$ maximum identity, reducing it to 3655 proteins. Using RS-ESM, M. acetivorans contains 946 words or which 659 are $k \geq 7$ and 300 are $k \geq 10$. Those satisfying both $k \geq 10$ and $n \geq 9$ are shown in Supplementary Material Table 5.

The words mostly represent islands of extreme conservation within what are fairly divergent families. For instance the 17 -mer HHRIKNNLQVISSLLDL is found in histidine kinases (Fig. 11), where its location corresponds to the start of the HisKA_2 domain (Pfam PF07568). There do not appear to be any homonymous words of $k \geq 10$ in the M. acetivorans proteome. M. acetivorans words are dominated by the preponderance of components of histidine kinase domains and PKD domains (Pfam PF00801).

## Vocabulary analysis in a medium-sized eukaryotic proteome

The fungus Yarrowia lipolytica has 6524 proteins of which 5864 are $<40 \%$ identical. It therefore has almost exactly the same overall proportion (just under $90 \%$ ) of proteins in gene families as $M$. acetivorans. Using RS-ESM, Y. lipolytica contains 1954 words of which 940 are $k \geq 7$ and 401 are $k$ $\geq 10$. All words satisfying both $k \geq 10$ and $n \geq 9$ are shown in Supplementary Material Table 6. By contrast with M. acetivorans, the prominent $Y$. lipolytica words are composed entirely of simple sequence repeats.


Figure 10. N-terminal region of 3 C . muridarum transporter proteins showing the GPNGAGKSTL word (shaded).

Q8TKQ3
Q8TK73 Q8TMC7 Q8TRB3 Q8TSN7 Q8TSN6 Q8TR62 Q8TQC1 Q8TT85 Q8TIW4 Q8TRA6 Q8TT86 Q8TP40 Q8TPM8 Q8TKN1 Q8TQA4 Q8TU70 Q8TMA7 Q8TS36 Q8TM82 Q8TQA5

Q8TKQ3
Q8TK73 Q8TMC7 Q8TRB3 Q8TSN7 Q8TSN6 Q8TR62 Q8TQC1 Q8TT85 Q8TIW4 Q8TRA6 Q8TT86 Q8TP40 Q8TPM8 Q8TKN1 Q8TQA4 Q8TU70 Q8TMA7 Q8TS36 Q8TM82 Q8TQA5
$---V D A A R K K E I H H R I K N N L Q V I S S L L D L Q A E N F S K H E V C K T P K V V E A F K$
$---I E A A R K K E I H H R I K N N L Q V I S S L L D L Q A E K F N K R E G I K D S E V M E A F R$
$---A E K L R K K E I H H R I K N N L Q V I S S L L D L E C D S L L S G-T P D H K K I A E A F R$
$---M K E S R I K E I H H R I K N N L Q V I S S L L D L Q A E K F E D P------T I R Q A F R ~$
-AKIEDARKKEIHHRIKNNLQVISSLLDLQAEKFRDK------DVLEAFR
LAEIDKTRIKEIHHRIKNNLQVISSLLDLQAEKFRDK------EVLEAFR
LATIEIARKKEIHHRIKNNLQVISSLLDLQAEKFGNKKYIMNSEVMDAFR
-ANIEIARKKEIHHRIKNNLQVISSLLDLQADKFDNP-------KVIEAFR
LENIEIARKQEIHHRIKNNLQVISSLLDLQAEKFNNREDIKDSEILEAFR
-VKIEDARKKEIHHRIKNNLQVISSLLDLQAEKFSHREAVPTLEILEAFK
-ANIEIARKKEIHHRIKNNLQVISSLLDLQAEKFRSREHVEDSEVLNAFK
-EKIDKIRIKEIHHRIKNNLQVISSLLDLQAEKFQNK-------EVLEAFR
-EKMERIRIKEIHHRIKNNLQVISSLLDLQAEKFRDK------EVLEAFR
-------IKEIHHRIKNNLQVISSLLDLQIDIFSNREICKTPEVIEAFR
LDKIQIARQKEIHHRIKNNLQVISSLLDLQAGKFNNKEHIRDSEVLEAFK
L---FELASKKELHHRIKNNLQVISSLLDLQADLFKGKKTITDSEVLKAFN
LANLEIARKKEIHHRIKNNLQVISSLLDLQAEKFNSREDIKDSEVLEAFR
---IETVRKQEIHHRIKNNLQVISSLLDLQAEQFKNRECIKNSEVLEAFR
LQEIDIIRKQEIHHRIKNNLQVISSLLDLQAEKFRGKKNIEDSKILEAFK
---IESARKKEIHHRIKNNLQVISSLLDLQAEKFKDREDIKDSEVLEAFR
$---I E I A R K K E I H H R I K N N L Q V I S S L L D L Q A E Q F K N R E N I K D S E V L E A F R ~$

Figure 11. HHRIKNNLQVISSLLDL (shaded) in part of a histidine kinase alignment.

## Vocabulary analysis in a large eukaryotic proteome

Brachydanio rerio, the zebrafish, has 14049 proteins, of which 8312 are no more than $40 \%$ identical, indicating that just over $40 \%$ of the total are members of moderately or closely related protein families, a considerably higher proportion than in the smaller eukaryotic proteomes (at $\sim 10 \%$ for Y. lipolytica). Using RS-ESM, B. rerio contains 2938 words or which 1380 are $k \geq 7$ and 452 are $k \geq 10$. All words satisfying both $k \geq 10$ and $n \geq 9$ are shown in Supplementary Material Table 7.

Just as prominent M. acetivorans words are dominated by components of histidine kinase domains and PKD domains, prominent $B$. rerio words are in most cases part of an EGF domain, with a handful of SCRC domains (PF00530). There are also several examples of low complexity words
(Supplementary Material Table 7), similar to $Y$. lipoloytica.

## Discussion

An improved algorithm for vocabulary analysis in texts of human origin, has been applied to proteomes. In its two variants, CW-ESM and RSESM, it achieves an accuracy of $60 \%-70 \%$ (Table 7) and in the case of RS-RSM has approximately $85 \%$ sensitivity. This sensitivity estimate is based on 895 true positive hits as compared to the 1042 words used more than twice in Alice in Wonderland. It remains an approximation as the algorithm detects phrases longer than single words (DWoPs, see Tables 2-6). Although CW-ESM is slightly less accurate than RS-ESM and less than half as sensitive (Table 7), it is less liable to false positives at $k=6$ (Fig. 3). Since many protein
homonyms appear at $k=6$, CW-ESM remains an important accessory algorithm for the study of short words in protein sequences. This paper therefore solves the problem posed by Schmitt et al. (1996) of how to apply the method of Brendel et al. (1986) to longer alphabets. Since the combinatorial explosion problem is greater in human texts than in protein sequences, the adequacy of the algorithm for detecting words in texts implies that it can do the same for proteins, should such words exist.

It is notable that the words detected by the algorithm follow Heaps' Law, a linear increase in word count as text size increases, for both human texts (Fig. 2) and proteins (Fig. 9). A similar result for raw strings in proteins is already known (Mukhopadhyay et al. 2006). Within superkingdoms, Heaps' Law correlations are strongest for human texts and eubacterial proteomes. By contrast, between superkingdoms, eukaryotic proteomes appear to be nearly three times more word-rich on average than the two prokaryotic superkingdoms. However, caution must be exercised in intersuperkingdom comparisons as the average proteome size is almost four times larger for eukaryotes. When only small eukaryote proteomes are used, the proportionately larger number of words decreases to similar levels (Table 9). Heaps' Law therefore appears to deviate from linearity in large eukaryotic proteomes, but eukaryotic proteomes may still be comparable to prokaryotic proteomes at smaller sizes. One possible explanation for this is that larger eukaryotic proteomes are richer in gene families, adding an extra source of words to the general trend implied by Heaps' Law, and supported by the observation that about $40 \%$ of $B$. rerio proteins are $>40 \%$ identical.

It should be noted that vocabulary analysis is not the same as segmentation (Wang, 2001; Cohen et al. 2002), when a text known to be composed of words is split into candidate words. Segmentation is often used in computer analysis of pictographic languages such as Japanese kanji script, where word boundaries are unclear. By contrast, vocabulary analysis algorithms search for the presence of candidate word structures in bodies of symbols that may not necessarily contain them.

The fact that human texts are an order of magnitude more enriched in words than proteomes (Table 9), suggests that the linguistic analogy for biological sequences remains a weak one, and furthermore that segmentation algorithms, relying as they do on complete decomposition of the text
into words, are unlikely to be applicable to protein sequences. Nevertheless, the presence of identifiable word-like structures within proteomes is intriguing. Shuffled proteomes, like shuffled texts, lose their word content. Within a shuffled proteome, false positives are rare and in neither RSESM nor CW-ESM are found at $k \geq 7$. Words of $k=6$ are ambiguous, as they are generated as false positives by RS-ESM (Fig. 3). By analogy with words in human texts, proteome words are suggested to be sequences that are intolerant to mutation but are nevertheless relatively contextindependent in their function.

Analysis of the distribution of words in individual proteomes demonstrates two main categories:

1. conserved stretches within proteins of essentially similar function (see Figs. 4-6).
2. homonyms appearing in proteins of demonstrably different functions (see Fig. 8).
Conserved words can be further split into:
a) relatively uninteresting sequence identities within closely related proteins
b) ultra-conserved words in rather more divergent proteins (see Figs. 4, 5, 6, 10 and 11 for examples).

Homonyms are plentiful but short, rarely $k>6$, whereas ultra-conserved stretches are often much longer, for instance the 17 -mer HHRIKNNLQVISSLLDL (Supplementary Material Table 5 and Fig. 11) which forms a word in a family of histidine kinase proteins in M. acetivorans. Only words of up to $k=18$ were tested in this paper, so no estimate can be made of the longest existing word. M. acetivorans has a low complexity 18 -mer, STDDSTDDSTDDSTDDST (Supplementary Material Table 5), and Y. lipolytica has eight (Supplementary Material Table 6). The longest high complexity words are the 6 EGF domain words and a zinc finger word found in B. rerio (Supplementary Material Table 7).

Not all words can be easily designated as homonyms or conserved. For instance, in C. muridarum GPTGSGK is found in different Pfam domains in different proteins (IPPT, AAA_2, GSPII_E and ABC_tran), initially suggesting homonymity. However, in all these cases GPTGSGK forms part of an ATP-binding cassette. Whether this is best explained by ultra-conservation within highly divergent ATP-binding proteins with a distant

GPTGSGK-containing ancestor or by the multiple evolution of ATP binding capacities by convergence to these words, is debatable. The similar longer word GPNGAGKSTL (Fig. 10) is also an ATP-binding element, but, unlike GPTGSGK, GPNGAGKSTL is always found in the ABC_tran domain (PF00005), and therefore is more likely to be an example of ultra-conservation than convergence.

Homonyms are assumed to have different functions within their respective proteins, especially when they can be shown to have different structures (e.g. see Fig. 8). However, apparent homonyms with similar structures may be a result of convergent molecular evolution on a micro-scale, perhaps the case with C. muridarum GPTGSGK. For a specific structural example, VLVIGA is a 6-mer homonym in NRL3D, occurring in: 1BFD, a benzoylformate decarboxylase (EC 4.1.1.7) from Pseudomonas putida; 1AD3A, an aldehyde dehydrogenase (EC 1.2.1.5) from rat; and in 1D4OA, a bovine NADPH transhydrogenase (EC 1.6.1.1). In each of these cases VLVIGA is found in a different Pfam domain: in TPP_enz_M (PF00295), Aldedh (PF00171) and $\overline{\mathrm{PNT}} \overline{\mathrm{T}}$ (PF02233) respectively. Nevertheless, the conformation of VLVIGA is remarkable similar in each case, always being the point at which a short beta-sheet ends (Fig. 12).

Comparison of different proteomes indicates that the peptide vocabularies can be quite different in character from species to species. For instance, the prominent words in the small eubacterial species $C$. muridarum are dominated by components of adherence factor proteins mixed with a handful of peptides from other domains and some low complexity elements (Supplementary Material Table 4). The large archaeon M. acetivorans has many words from PKD and histidine kinase domains (Supplementary Material Table 5). The fungus Y. lipolytica has only low complexity words within its major vocabulary (Supplementary Material Table 6). Finally, the vertebrate B. rerio is dominated by EGF domain words (Supplementary Material Table 7). A fuller exploration of how typical these vocabulary patterns may be, is beyond the scope of the present paper. However, it indicates that each proteome may potentially be identifiable by the characteristics of its vocabulary; e.g. rich in low complexity, or with certain typical domainlinked vocabularies, raising the prospect of a peptide vocabulary signature analogous to the genome signature found in DNA. This may be useful in metagenomic analysis.

Thode et al. (1996), in pairwise comparisons of proteins, found many matches of 6 residues within windows of 10 , and showed that these occurred far less frequently between pairs of random proteins.


Figure 12. VLVIGA in (from left to right) 1D4O.A, 1BFD and 1AD3.A. The backbone trace is drawn as a black line.

The method of Thode et al. (1996), differs from the one presented here in that they used a criterion of $60 \%$ identity with strings of $k=10$. Unlike the present method, there was no previous algorithmic identification of candidate words by statistical properties. Rather, they commenced with a small group of proteins and extracted all their initial 10 -mer strings from those sequences. These were then compared against the whole protein database, and matches of $6 / 10$ or better recorded. It thus has some similarities to method RS above, but incorporating fuzziness. Regardless of these methodological incongruences, the detection by Thode et al. (1996) of a far greater quantity of short common strings in real protein pairs than in shuffled ones, parallels the results presented in Figure 3. Therefore, it is justifiable to believe that even words of $k=6$ may be mostly due to something other than random coincidence. The nature of this pressure may be conservation, amply demonstrated by the various ultra-conserved words within fairly divergent proteins (Figs. 4, 5, 6 and 11) or it may be convergent evolution. The latter of these raises the possibility than the presence of an apparent homonym within a protein may imply positive selection within the family to which that protein belongs, and which may be detectable using appropriate methods (Yang, 1997; Anisimova and Yang, 2007). For instance, if a candidate homonymic word is found in two proteins of differing function, for instance different Pfam families, and positive selection can be statistically demonstrated in each of those families over the region of the homonym, a selective convergent scenario for the origin of that homonym would be highly suggestive.

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## Peptide Vocabulary Analysis Reveals Ultra-Conservation and Homonymity in Protein Sequences

## Derek Gatherer

## Supplementary Materials

Table S1. Words detected in eukaryotic proteomes using RS-ESM, $k=6-18$.

| Species | Proteins | Kres | Words |
| :---: | :---: | :---: | :---: |
| H. sapiens | 37993 | 16405.3 | 30360 |
| M. musculus | 32971 | 14645.0 | 29463 |
| A. thaliana | 34712 | 14124.8 | 47516 |
| T. nigroviridis | 27836 | 11286.0 | 26742 |
| C. elegans | 22434 | 9699.5 | 14167 |
| D. melanogaster | 14396 | 8055.1 | 7509 |
| D. discoideum | 13017 | 6817.5 | 16628 |
| A. gambiae | 15145 | 6125.0 | 6509 |
| C. briggsae | 13192 | 6038.7 | 6687 |
| G. zeae | 11636 | 5952.6 | 5302 |
| B. rerio | 14049 | 5940.9 | 12267 |
| A. oryzae | 12053 | 5410.2 | 5498 |
| R. norvegicus | 11839 | 5350.2 | 10466 |
| L. major | 8010 | 5137.4 | 4795 |
| D. pseudoobscura | 9877 | 5115.2 | 4412 |
| A. fumigatus | 9906 | 4782.5 | 3891 |
| P. falciparum (3D7) | 5282 | 4001.3 | 10494 |
| C. neoformans | 6569 | 3558.9 | 2787 |
| C. neoformans (JEC21) | 6437 | 3449.1 | 2461 |
| P. yoelii | 7590 | 3385.6 | 9444 |
| Y. lipolytica | 6524 | 3118.5 | 3661 |
| D. hansenii | 6309 | 2902.5 | 2401 |
| S. cerevisiae | 5800 | 2891.7 | 2227 |
| B. taurus | 8292 | 2890.4 | 3652 |
| C. glabrata | 5180 | 2610.3 | 1742 |
| K. lactis | 5326 | 2504.6 | 1249 |
| G. gallus | 5387 | 2443.6 | 3384 |
| S. pombe | 5011 | 2351.3 | 1306 |
| A. gossypii | 4720 | 2314.3 | 1103 |
| T. annulata | 3790 | 2025.0 | 3409 |
| T. parva | 4070 | 1895.3 | 1899 |
| C. hominis | 3886 | 1757.6 | 924 |
| E. cuniculi | 1909 | 693.5 | 308 |
| T. gondii | 489 | 377.9 | 230 |
| G. theta | 598 | 178.5 | 38 |

Table S2. Words detected in eubacterial proteomes using RS-ESM, $k=6-18$.

| Species | Proteins | Kres | Words |
| :---: | :---: | :---: | :---: |
| R. baltica | 7271 | 2290.5 | 1658 |
| Anabaena. sp | 6069 | 1955.5 | 2196 |
| A. tumefaciens (Cereon) | 5305 | 1687.1 | 1195 |
| A. bacterium | 4771 | 1677.5 | 1011 |
| B. fragilis (ATCC 25285) | 4234 | 1537.1 | 841 |
| A. dehalogenans | 4345 | 1516.6 | 1902 |
| B. anthracis (Sterne) | 5288 | 1460.1 | 996 |
| Azoarcus. Sp. | 4490 | 1393.5 | 764 |
| G. violaceus | 4406 | 1377.7 | 1412 |
| E. coli (K12) | 4323 | 1372.0 | 687 |
| C. difficile | 3711 | 1164.1 | 884 |
| L. interrogans (icterohaemorrhagiae) | 3654 | 1150.7 | 705 |
| Acinetobacter. Sp. | 3310 | 1048.4 | 266 |
| A. ehrlichei | 2862 | 984.1 | 524 |
| A. borkumensis | 2752 | 908.2 | 236 |
| D. geothermalis | 2821 | 901.0 | 538 |
| B. abortus | 3023 | 877.9 | 273 |
| T. denticola | 2753 | 863.6 | 573 |
| S. elongatus | 2451 | 770.0 | 378 |
| S. haemolyticus | 2634 | 756.6 | 370 |
| C. chlorochromatii | 1991 | 750.9 | 907 |
| T. thermophilus (HB27) | 2200 | 667.6 | 552 |
| P. amoebophila | 2023 | 658.9 | 1210 |
| F. nucleatum | 2046 | 641.1 | 250 |
| B. longum | 1723 | 638.8 | 181 |
| T. maritima | 1852 | 582.8 | 191 |
| C. jejuni | 1836 | 538.6 | 104 |
| H. pylori (26695) | 1551 | 491.8 | 172 |
| A. aeolicus | 1552 | 488.9 | 121 |
| P. marinus (CCMP 1378) | 1707 | 484.4 | 105 |
| D. ethenogenes | 1502 | 416.5 | 95 |
| B. afzelii | 1257 | 357.8 | 223 |
| C. muridarum | 916 | 324.3 | 40 |
| M. pneumoniae | 687 | 239.7 | 380 |
| A. yellows | 690 | 176.6 | 266 |

Table S3. Words detected in archaeal proteomes, using RS-ESM, $k=6-18$.

| Species | Proteins | kres | Words |
| :--- | :--- | ---: | ---: |
| M. acetivorans | 4467 | 1392.1 | 2317 |
| H. marismortui | 4234 | 1200.1 | 1006 |
| M. barkeri | 3616 | 1126.3 | 1701 |
| M. mazei | 3302 | 9004.3 | 999 |
| M. hungatei | 3095 | 997.2 | 823 |
| S. solfataricus | 2910 | 827.9 | 571 |
| N. pharaonis | 2784 | 815.9 | 387 |
| H. walsbyi | 2644 | 787.4 | 399 |
| S. tokodaii | 2816 | 75.9 | 449 |
| H. salinarium | 2426 | 680.0 | 313 |
| M. burtonii | 2242 | 676.7 | 262 |
| A. fulgidus | 2398 | 660.8 | 473 |
| P. aerophilum | 2589 | 654.9 | 219 |
| P. kodakaraensis | 2301 | 637.7 | 188 |
| S. acidocaldarius | 2221 | 631.7 | 202 |
| P. furiosus | 2045 | 577.7 | 159 |
| P. horikoshii | 2077 | 569.4 | 180 |
| P. abyssi | 1785 | 539.2 | 145 |
| M. thermoautotrophicum | 1869 | 524.7 | 213 |
| M. jannaschii | 182 | 504.9 | 205 |
| M. kandleri | 1687 | 501.0 | 277 |
| M. stadtmanae | 1533 | 493.6 | 113 |
| M. maripaludis | 1722 | 490.7 | 143 |
| A. perrix | 1576 | 482.7 | 79 |
| P. toridus | 1535 | 471.3 | 49 |
| T. acidophilum | 1482 | 453.2 | 56 |
| T. volcanium | 1523 | 452.7 | 17 |
| N. equitans | 536 | 151.5 |  |

Table S4. All words of $k \geq 6$ detected in C. muridarum (proteins $<63 \%$ identical), using RS-ESM.

| Word | $\boldsymbol{k}$ | Protein family | $\boldsymbol{n}$ |
| :--- | :--- | :--- | :--- |
| GGKGGLTVQIGG | 12 | adherence factor | 3 |
| FQEEHGHCRVP | 11 | helicase | 4 |
| GPNGAGKSTL | 10 | ABC transporter | 3 |
| EPAPEPAPE | 9 | low complexity | 3 |
| SDTESTNGN | 9 | low complexity | 3 |
| NPQLASWV | 8 | helicase | 4 |
| SGSGKSSL | 8 | ABC transporter | 3 |
| IHDVEQNG | 8 | DUF1547 (PF07577) | 3 |
| GPTGSGK | 7 | various | 4 |
| FRVTDPN | 7 | adherence factor | 3 |
| GIEGLIH | 7 | S1 RNA binding | 3 |
| DDDDDDD | 7 | low complexity | 3 |
| EGRCMGL | 7 | adherence factor | 3 |
| NDVTPAD | 7 | adherence factor | 3 |
| KTAAKKA | 7 | histone-like | 3 |
| LGGGAIL | 7 | Chlam_PMP (PF02415) | adherence factor |
| HGIWIAG | 7 | low complexity | 3 |
| SSSSSS | 6 | homonym | 3 |
| RSLLNK | 6 | homonym | 5 |
| VLLGLG | 6 | helicase | 4 |
| GKLSED | 6 | adherence factor | 4 |
| SFRAIP | 6 | homonym | 4 |
| LPLFSL | 6 | homonym | 3 |
| SSSFAL | 6 | homonym | 3 |
| IAILLS | 6 | homonym | 3 |
| RLKTIL | 6 | homonym | 3 |
| ALGIAA | 6 | homonym | 3 |
| VVLFDE | 6 | homonym | 3 |
| AASLIR | 6 | homonym | 3 |
| SLQEGL | 6 | homonym | 3 |
| ALPGVG | 6 | MMR_HSR1 (PF01926) | homonym |
| PNVGKS | 6 | homonym | 3 |
| EKILSL | 6 |  | 3 |
| VLSYEL |  | 3 |  |

Table S5. All words of [ $k \geq 10$ AND $n \geq 9$ ] in M. acetivorans (proteins $<40 \%$ identical), sorted by occurrence, $n$.

| Word | $\boldsymbol{k}$ | Protein family | $\boldsymbol{n}$ |
| :--- | :--- | :--- | :--- |
| STDDSTDDSTDDSTDDST | 18 | low complexity | 27 |
| EIHHRIKNNLQVISSLL | 17 | histidine kinase | 27 |
| HHRIKNNLQVISSLLDL | 17 | histidine kinase | 21 |
| NMPVEYFDFNGN | 12 | PKD domain | 20 |
| VAYFHNMDWIE | 11 | PKD domain | 20 |
| GDGLYEDLTGNGEFSFVD | 18 | PKD domain | 19 |
| DLDGDGLYEDLTG | 13 | PKD domain | 17 |
| VVLATLTVSGKEKGSAN | 17 | PKD domain | 15 |
| VSGKEKGSANLSIGV | 15 | histidine kinase | 15 |
| ISSLLDLQAEKF | 12 | histidine kinase | 14 |
| PLGIIVNELVSNSLKHAF | 18 | PKD domain | 13 |
| GSANLSIGVKRLE | 13 | PKW complexity | 13 |
| YSFLPVYSFLPVYSFLPV | 18 | PKD domain | 12 |
| EGAADVVLATLTVSGKE | 17 | histidine kinase | 11 |
| TVPEENITVPEEN | 13 | PKD domain | 11 |
| AVPLGIIVNELVSNSLK | 17 | PKD domain | 10 |
| GTAPLTVNFTDQSTGSP | 17 | PKD domain | 9 |
| STGSPTSWFWDFGDG | 15 | PKD domain | 9 |
| VSEASGSTVTLYFDP | 15 | PKD domain | 9 |
| PTSWFWDFGDGANST | 15 | histidine kinase | 9 |
| LSPLPDQEYAPKDL | histidine kinase | 9 |  |
| DITERKKAEEAL | 14 | 9 |  |
| MDTAVPLGII | 12 | 9 |  |

Table S6. All words of [ $k \geq 10$ AND $n \geq 9$ ] in $Y$. lipolytica (proteins $<40 \%$ identical), sorted by length, $k$.

| Word | Protein family | $\boldsymbol{k}$ | $\boldsymbol{n}$ |
| :--- | :--- | :--- | ---: |
| QQQQQQQQQQQQQQQQQQ | low complexity | 18 | 67 |
| ATDTGATATDTGATATDT | low complexity | 22 |  |
| TVTGPTAGTTTITGTDGK | low complexity | 18 | 15 |
| SYSPTSPSYSPTSPSYSP | low complexity | 18 |  |
| IFIFIFIFIFIFIFIFIF | low complexity | 18 |  |
| YDSYDSYDSYDSYDSYDS | low complexity | 14 |  |
| PLAEPMPLPLAEPMPLPL | low complexity | 18 | 11 |
| SGSGSSGSGSSGSGSSGS | low complexity | 10 |  |
| ATDTATDTAATDTATDT | low complexity | 9 |  |
| GSGSGSGSEGSGSGSGS | low complexity | 18 | 31 |
| SQSQSQSQSQSQSQSQS | low complexity | 18 | 19 |
| NGNGSDGSNGNGSDGSN | low complexity | 18 | 17 |
| GSGSGSGSDSGSGSGSG | low complexity | 17 | 16 |
| SSSIPTGDVSSATPTGD | low complexity | 17 | 13 |
| DASSSIPTGDVSSATPT | low complexity | 11 |  |
| PTGDVSSATPTGDASSS | low complexity | 17 | 17 |
| TGGADASSTGGADASST | low complexity | 17 | 17 |
| TATDTGATDTATDTGAT | low complexity | 17 | 10 |
| TEQITVAPTGPVTTKTV | low complexity | 17 | 9 |
| KQKQKQKQKQKQKQKQK | low complexity | 17 | 9 |
| ATQTGGNGNNSGSNTAT | low complexity | 17 | 9 |
| ATDTGATATDTGATDT | low complexity | 17 | 9 |
| SPSYSPTSPSYSPTS | low complexity | 17 | 12 |
| ATDTGATATDTATD | low complexity | 17 | 17 |
| EPVTSEPVTSEPVT | low complexity | 17 |  |
| PGPAPSPGPGPAPS | low complexity | 16 | 13 |
| SDSDSDSDSDSDS | low complexity | 15 | 12 |
| DSDSDSDSDSDSD | low complexity | 14 | 10 |
| PSSTEAPSSTEAP | low complexity | 14 | 14 |
| GSNTATQTGGNGN | low complexity | 13 | 10 |
| TKTVTGPTAGT | low complexity | 13 | 13 |
| ASASASASASA | low complexity | 13 | 13 |

Table S7. All words of [ $k \geq 10$ AND $n \geq 12$ ] in $B$. rerio (proteins $<40 \%$ identical), sorted by occurrence.

| Word | Protein family | $k$ | $n$ |
| :---: | :---: | :---: | :---: |
| DDDDDDDDDDDDDDDDD | low complexity | 18 | 184 |
| QQQQQQQQQQQQQQQQQ | low complexity | 18 | 38 |
| YQCKCEGLFVWPNDTCHA | EGF domain | 18 | 22 |
| GSFNCSCLSAFTVTDRNQ | EGF domain | 18 | 19 |
| AQAQAQAQAQAQAQAQAQ | low complexity | 18 | 16 |
| KKKKKKKKKKKKKKKKKKK | low complexity | 18 | 16 |
| NGTEYECKCEVDHVWPSN | EGF domain | 18 | 14 |
| CGLNGTEYECKCEVDHVW | EGF domain | 18 | 14 |
| CGPNSICNNTIGSYNCSC | EGF domain | 18 | 14 |
| MSDPEPCRIKQEETEELI | zinc finger | 18 | 13 |
| YSNCTNEIGSYNCSCLDG | EGF domain | 18 | 12 |
| CDVITNGSCTCINGLPA | EGF domain | 17 | 22 |
| NGSCTCINGLPADGQFC | EGF domain | 17 | 21 |
| VCSLNETRYQCKCEGLF | EGF domain | 17 | 19 |
| ECLFSPPVCGPYSNCTN | EGF domain | 17 | 17 |
| тнтнтнтнтнтнтнтнT | Iow complexity | 17 | 17 |
| CRELDCGAPVQVLRAA | SCRC (PF00530) | 16 | 19 |
| DINECEDAASVCGQYS | EGF domain | 16 | 17 |
| TDRNQPVSNSNPCNVC | EGF domain | 16 | 17 |
| TCGCIQALPSEGSLCQ | EGF domain | 16 | 15 |
| CDAAFDQQDAEVVCR | SCRC (PF00530) | 15 | 25 |
| NSIGSFNCSCLSAFT | EGF domain | 15 | 19 |
| IGGYMCSCWNGFNVS | EGF domain | 15 | 15 |
| QVCDSIVGSTCGCIQ | EGF domain | 15 | 14 |
| SINNTCEDVNECLKS | EGF domain | 15 | 12 |
| SNSNPCNVCSLNET | EGF domain | 14 | 18 |
| PERPPVSAPAPERP | low complexity | 14 | 16 |
| LTETQVKIWFQNRR | homeobox | 14 | 12 |
| DIDECLFSPPVCG | EGF domain | 13 | 12 |
| PVCGPYSNCTNE | EGF domain | 12 | 15 |
| PGGVGGVPGGVG | low complexity | 12 | 13 |
| NLPINSNNTCTD | EGF domain | 12 | 13 |
| LRAAAFDKGD | SCRC (PF00530) | 10 | 13 |

