# Hydrophobicity, expressivity and aromaticity are the major trends of amino-acid usage in 999 *Escherichia coli* chromosome-encoded genes

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#### **ABSTRACT**

Multivariate analysis of the amino-acid compositions of 999 chromosome-encoded proteins from Escherichia coli showed that three main factors influence the variability of amino-acid composition. The first factor was correlated with the global hydrophobicity of proteins, and it discriminated integral membrane proteins from the others. The second factor was correlated with gene expressivity, showing a bias in highly expressed genes towards amino-acids having abundant major tRNAs. Just as highly expressed genes have reduced codon diversity in protein coding sequences, so do they have a reduced diversity of amino-acid choice. This showed that translational constraints are important enough to affect the global amino-acid composition of proteins. The third factor was correlated with the aromaticity of proteins, showing that aromatic amino-acid content is highly variable.

#### INTRODUCTION

This paper investigates the amino-acid usage in *Escherichia coli* proteins, to describe general trends and their biological implications. The method used, correspondence analysis, has also been used to analyze codon usage by Grantham and colleagues (1-3), review in 4). The first factor underlying variations in codon usage is the genome of origin. In addition, there is a considerable within-species codon usage variability. Among *E. coli* genes this diversity is linked to gene expressivity: genes with a potentially high expression level are biased towards the subset of codons that are best recognised by the most abundant tRNA species (5). In contrast with codon usage, the interspecific variability in amino-acid usage is low (3). The present study focuses on amino-acid usage of proteins from a single species, *E. coli*, because a large body of sequence data is available for this species.

# **MATERIALS AND METHODS**

#### Data ser

The data set was 999 protein sequences encoded by genes on the *E.coli* chromosome, corresponding to a total of 385,404

amino-acids. As this is about 25% of the estimated total number of chromosome-encoded proteins, the sample is large enough to be representative. The nonoverlapping ECOSEQ6 collection (6) was structured (7) using the entity-relationship model of ACNUC (8-10). The retrieval system, Query, associated with ACNUC, allows elaborate sequence managements. The ECOSEQ6 collection contains the sequences of a single allele per locus, so that there is no overweighting due to sequence redundancy or DNA polymorphism. This is not a negligible problem since, for instance, there are 16 complete sequences of the gnd locus of E. coli in GenBank (11) release 78. The disadvantage is that the allele sequences in Rudd's collection are from different strains, leaving open the possibility of intraspecific variations affecting results. There are not yet enough data to answer this question, but there seems to be very little polymorphism at the amino-acid level, about 1% for the average number of amino-acid differences per site between two alleles (12).

Plasmid-encoded proteins are not included in Rudd's collection. This minimizes the horizontal gene transfer effect, which is more likely for plasmid-encoded protein. The amino-acid usage of proteins encoded by genes recently incorporated in the *E.coli* genome may differ from native *E.coli* proteins.

Partial sequences (7%) were discarded because the amino-acid composition of a fragment could be atypical of the whole protein composition. Poorly documented open reading frames (12%) were discarded to help analysis of results. The Rudd nomenclature, by which most unidentified ORFs are given a name starting with 'y', ensure their easy removal. Information on the remaining sequences is, however, highly variable. Proteins with fewer than 100 amino-acids (5%) were excluded to minimize influence of stochastic variations in the amino-acid compositions of small peptides. The threshold value of 100 amino-acids is roughly the minimum size for a protein to have an enzymatic function (13).

The N-terminal methionine was not removed. This is an arbitrary choice because the rules that govern the removal of N-formylmethionine are not completely understood (14). This choice did not noticeablely alter the results, there were negligible variations only for small proteins with a low methionine frequency. The special case of selenocystein was not handled

because it is too rare; there are only three known selenopolypeptides in *E.coli* (15). Lastly, post-translational modifications were not taken into account.

#### Multivariate analysis

The  $\chi^2$  metric was used as a measure of the distance between the amino-acid composition of two proteins. Correspondence analysis can then extract orthogonal linear combinations of amino-acid frequencies that best summarize the data. These trends are optimal because they take into account most of the initial variability (16, 17). The squared distance between two sequences x and y is defined as:

$$d^{2}(x,y) = n.. \sum_{i=1}^{20} \frac{1}{n_{\bullet i}} \left( \frac{n_{xi}}{n_{x\bullet}} - \frac{n_{yi}}{n_{y\bullet}} \right)^{2}$$

where  $n_{xi}$  and  $n_{yi}$  are the number of amino-acids of kind i in sequence x and y,  $n_x$  and  $n_y$  are the total number of amino-acids in sequences x and y,  $n_i$  the total number of amino-acids of kind i in the dataset and  $n_i$ , the total number of amino-acids in the dataset. The advantage of the  $\chi^2$  metric over the usual Euclidian distance used in principal component analysis of compositional data (18), is that information on rare amino-acids are not masked by frequent amino acids because of the  $1/n_{ij}$  weighting.

The correspondence analysis was computed with the program MacMul (19, 20) on a Macintosh plus. The results were checked by running a different program (21) on a different computer (Sun SPARCStation 10) to ensure that there were no computational errors. Analysis of results was facilitated by the interactive DIGTT software (22). The absence of bias due to the low frequencies of rare amino-acids was checked by removing them and repeating the analysis.

#### Identification of protein characters

Three scores were computed for each protein to help interpret the results. The GRAVY score (23) is an estimate of the overall hydrophobicity of the protein, the highest scores indicating a hydrophobic character. The GRAVY score is a linear combination of amino-acid relative frequencies:

$$GRAVY = \sum_{i=1}^{20} \alpha_i f_i$$

where  $f_i$  is the relative frequency of amino-acid of kind i in the protein and  $\alpha_i$  the hydropathy index of this amino-acid (23).

The codon adaptation index (CAI) is an empirical measure of synonymous codon usage bias (24), which is positively correlated with the expressivity level of genes.

$$ln(CAI) = \sum_{i=1}^{61} f_i ln w_i$$

where  $f_i$  is the relative frequency of codon of kind i in the coding sequence, and  $w_i$  the ratio of the frequency of codon of kind i to the frequency of the major codon for the same aminoacid, as estimated from examining 25 highly expressed genes (24). Here, CAI has the advantage over other indices, such as the Mean Number of tRNA Discrimination per elongation cycle (5), of being almost independent of amino-acid frequencies.

The AROMATICITY is the relative frequency of aromatic amino-acids,

$$AROMATICITY \ = \ \sum_{i=1}^{20} \! \delta_i f_i \, ,$$

where  $f_i$  is the relative frequency of amino-acid of kind i in the protein and  $\delta_i = 1$  when the amino-acid is aromatic (Phe, Tyr, Trp) and  $\delta_i = 0$  otherwise.

#### **RESULTS**

#### Glbal amino-acid composition

The mean amino-acid composition of the proteins in the dataset (Table 1) was found to be very similar (r = 0.95) to that reported previously (25). The results are also consistent (r = 0.89) with the experimentally determined composition of the total proteins

Table 1. Average amino-acid composition (% ± SD)

<u> AA</u>	Total	IMP	non IMP	N	С
Ala	9.7 ±2.6	10.4 ±2.3	9.6 ±2.7	8.8±3.5	9.6
Arg	$5.8 \pm 2.2$	3.7 ±1.3	$6.0 \pm 2.2$	4.4±2.1	5.5
Asn	3.8 ±1.4	3.0 ±1.2	3.9 ±1.4	}10.5±3.0	9.0
Asp	5.3 ±1.8	$2.5 \pm 0.8$	5.7 ±1.5	)	
Cys	1.2 ±1.0	$1.0 \pm 0.7$	1.2 ±1.0	1.4±1.1	1.7
Gln	4.3 ±1.8	$2.6 \pm 1.2$	4.5 ±1.7	}10.6±3.5	9.8
Glu	$6.1 \pm 2.3$	$2.5 \pm 1.1$	6.6 ±2.0	) .0.0_0.0	,,,,
Gly	7.5 ±2.1	8.7 ±2.0	7.3 ±2.0	8.1±2.8	11.5
				$2.1 \pm 1.2$	
Пе	5.9 ±1.9	$8.1 \pm 2.0$	$5.7 \pm 1.7$	5.0±2.0	5.4
Leu	10.2 ±2.7	13.2 ±2.9	9.9 ±2.5	8.1±2.8	8.4
Lvs	4.7 ±2.3	$2.9 \pm 1.2$	$4.9 \pm 2.0$	$6.5 \pm 3.1$	6.4
Met	2.8 ±1.2	3.9 ±1.3	2.7 ±1.1	1.9±1.1	2.9
Phe	3.8 ±1.6	6.3 ±1.9	3.5 ±1.3	3.8±1.6	3.5
				4.7±1.9	
				6.8±2.7	
Thr	5.3 ±1.5	5.3 ±1.4	5.3 ±1.5	5.9±2.2	4.7
Trp	$1.3 \pm 1.0$	$2.6 \pm 1.2$	$1.2 \pm 0.9$	$1.1 \pm 0.8$	1.1
Tyr	2.7 ±1.3	$3.0 \pm 1.3$	2.7 ±1.3	3.3±1.5	2.6
				$7.0\pm 2.2$	

IMP is the group of 114 integral membrane proteins given in table 3. Column N contains the results previously reported (25) and column C is the experimentally determined total protein composition (26). Since determination of protein composition requires the hydrolysis of all amide bonds, the relative amounts of Asp:Asn and Glu:Gln cannot be estimated, and their values are usually assumed to be 1:1. Here, the ratios were found to be about 3:2, showing that the acidic form was more abundant.

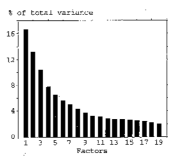


Figure 1. Factors of the correspondence analysis ranked in decreasing order of the fraction of total variance they accounted for.

of *E.coli* (26), although these results are not directly comparable, because of inequal protein concentrations *in vivo*.

On the basis of their average frequencies, amino-acids can be classified as very rare (Trp, Cys), rare (Tyr, Met, His), frequent (Gly, Val), very frequent (Leu, Ala) and intermediary for the remainder. In general, aliphatic amino-acids occur frequently, while aromatic or sulfur containing amino-acids are rare.

The relative frequencies of amino-acids within protein have unimodal, nearly symetric distributions, except for rare amino-acids (Trp and Cys), because they are quite often absent from a protein (about 10% of proteins lacked Trp or Cys in our data set).

#### Selection of factors

The relative importance of factors, as juged by the difference with their following factor, was found to vanish with factor 4

**Table 2.** Definition of the first three factors of the correspondence analysis (F1, F2 and F3)

AA	FI	F2	F3	ДF	RF	F1.RF
Ala	-0.319	-0.306	-1.515	35	0.114	-0.036
Arg	+1.395	+1.538	-0.039	5	0.016	+0.023
Asn	+0.239	-0.964	+1.425	14	0.046	+0.011
Asp	+1.428	-0.680	+0.696	14	0.046	+0.065
Cys	+0.102	+1.759	+1.036	1	0.003	+0.000
Gln	+1.204	+1.316	-0.441	16	0.052	+0.063
Glu	+1.902	-0.314	-0.475	9	0.029	+0.056
Gly	-0.740	-0.679	-0.117	20	0.065	-0.048
His	+0.949	+1.496	+0.875	1	0.003	+0.003
Пе	-1.019	-0.853	-0.333	10	0.033	-0.033
Leu	-0.582	+1.171	-0.764	33	0.108	-0.063
Lys	+1.026	-2.302	+0.444	19	0.062	+0.064
Met	-1.107	-0.350	-0.497	6	0.020	-0.022
Phe	-1.888	-0.037	+1.242	8	0.026	-0.049
Pro	+0.123	+0.727	+0.399	25	0.082	+0.010
Ser	-0.527	+0.321	+0.248	29	0.095	-0.050
Thr	-0.097	-0.255	+0.372	27	0.088	-0.009
Trp	-2.233	+2.668	+2.687	l	0.003	-0.007
Tyr	-0.121	-0.347	+3.311	7	0.023	-0.003
Val	-0.520	-0.613	-0.825	26	0.085	-0.044
Σ				306	1.000	-0.069

The computation of F1 score for MalM (accession number = X04477) is explained. AF is the absolute frequency of amino acids in MalM, including initial methionine, and RF the relative frequency. The score of MalM on the first factor (-0.069) is found by summing the products of F1 by RF.

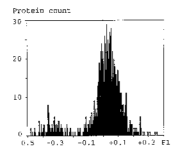


Figure 2. Distribution of scores for correspondence analysis factor 1. The minor peak (11% of total) contains integral membrane proteins.

(Figure 1). The three first factors, which accounted for 40% of the total variability of amino-acid composition of *E. coli* proteins, were then further analysed. These factors are defined in Table 2.

Table 3. List of proteins (score < -0.2) in the minor peak of F1 scores

Prot.	Fl	FUNCTION
CyoD	-0.483	Component of the cytochrome c utiquinel oxidase
S∄hD Mvx⊆	-0.482 -0.473	Anchor polypeptide of succinate dehydrogenase Methyl viologen resistance
TnaH BicA	-0.460	Transport of Tryptoppan
LacY	-0.456 -0.455	Involved in bicyclomycin resistance Transport of Lactose
DmsC NirC	-0.450	Anchor polypeptide of the anarobic dimethylsulfoxide reductase
CodB	-0.431 -0.427	Transport of mitrito Transport of cytosine
RhaT NupG	-0.427 -0.426	Transport of L-rhamiose
MreD	-D.426	Transport of nucleosides Envolved in the formation of rod shape of the cell
AraJ AppB	-0.422 -0.415	Transport of arabinose polymers (putative) Component of a third cytochrome exidese (putative)
Mtr	-0.413	Transmort of tryptophan
FrdD MglC	-0.412 -0.409	Anchor protein of the funarate reductase complex Transport of beta-methylgalactoside
PstC	-0.404	Transport of phosphate Transport of nitrate Transport of potassium Transport of lysine and cadaverine (putative)
Nark TrkG	-0.402 -0.398	Transport of potassium
CadB	-0.396 -0.396 -0.395	
CADB	-0.395	Component I of the cybochrome o ubiquinel-8 exidase Transport of 1-arabinose
AraH PotE	-0.392 -0.391	Transport of L-arabinose Transport of butrescine
CyoC	-0.390	Transport of putrescine Component III of the cytochrome o ubiquinal-8 axidase
Livn UgpE	-0.382	Transport of branched-chaip amino acids Transport of en-dlycerol-3-phosphate
Rfe	-0.374	Transport of sn-glycerol-3-phosphate Synthesis of lipid I
CynE Put P	-0.372	Synthesis of the cytochrome o ubiquinol-8 oxiduse Transport of proline Synthesis of polar bead of phospholipida
CdsA	-0.370	Synthesis of polar head of phospholipids
CynX UdpA	-0.369	Transport of sn-glycerol-3-phosphate
TekH NarV	-0.367	Transport of sn-glycerol-3-phosphate Transport of potassium Component of the second nitrate reductase
PotC	-0.366 -0.365	Transport of potassium Component of the second nitrake reductase Transport of approvidine and putrescine Transport of phenylalagine Transport of lysite
PheP	-0.365 -0.365	Transport of phenylalanine Transport of lysice
LysP TdcC	-0.364	2
OhpC HycC	-0.363 -0.362	Transport of hexoses phosphates Component of the formate bydrogene lyass
FhuB	-0.362	Component of the formate bydrogene lyass Transport of ferric bydroxenute
TyrP LapA	-0.361 -0.361	Transport of tyrosine Lipoprotein signal peptidase
AroF	-0.361 -0.360	Lipoprotein signal poptidase Transport of aromatic amino-acids
GabP ≥otB	-0.350 -0.358	Transport of 4-aminobutyrate Transport of spermidine and putroscine Transport of glutamate Component II of cytochrome d
GltS	-0.358 -0.357	Cransport of glutamate
CydB MraY	-0.353 -0.353	7
BtuC NhaA	-0.353 -0.348	Transport of vitamin B12 Transport of sedium
KdpA	-0.347 -0.346	Transport of potassium Transport of maltone
MalG	wO 739	Transport of maltose Anchor polypeptide of succinate dehydrogenase
SdhC RbsC	-0.339 -0.339 -0.337	Anchor polypeptide of softinate denyaloguals Transport of ribose Involved in multidrug resistance
EmrB FucP	-0.337 -0.332	Tavolved in multidrug resistance Transport of L-fucose
Mel3	-0.337 -0.330 -0.326	Transport of L-fucose Transport of meliblose Transport of meliblose Transport of feric enturobactin
FepQ ManY	-0.328 -0.328	Transport of ferric enturblactin Transport of manage
GlpT	-0.327 -0.326	Transport of manose Transport of glycerol-3-phosphate Component of hydrogeness 1
HyaC MrdB	+0.325	Penicillin binding procein o
CelD	-0.324 -0.324	Transport of bata-glucoside sugars Transport of glycerol
GlpF CysU	-0.324 -0.324	Transmort of sulfate
Narī	-0.324 -0.322 -0.321	Incher palvosatide for cytochrome ENR
PecD XylE	-0.321 -0.321	Transport of ferric dicitrate Transport of Xylose
PstA	-0.321 -0.318 -0.337	Transport of phosphate Transport of branched-chain amino-acids
LivM Ohg/T	-0.317 -0.313 -0.313	Transport of bexosa 6-phosphate Component of bydrogenase 3
RycD FrdC	-0.313	Component of hydrogenase 3 Anchor protein of fumarate reductase
ObiA	-0.310 -0.309 -0.307	Synthogis of ubiquinone Transport of alpha-ketogluterate
KgtP MalX	-0.307 -0.305	Transport of alpha-ketoglutarate
FanF	-0.304	Transport of gantothenate Involved in coll disvision
FtaM GltP	-0,304 -0.300	involved in cell disvision Transport of glutamate
₽epD	-0.298	Transport of ferric enterobactin
ProW PtsG	-0.299 -0.297	Transport of glutamate Transport of erric enterbactin Transport of elycine-betains and proline Transport of glucose Transport of ferric dicitrate.
PecC	-D.295	Transport of ferric dicitrate.
AraE RfcL	-0.294 -0.294	Transport of L-examinose
SimA	-0.294	Sensitivity to microcin B17
PgsA FdnI	-0.291 -0.288	Synthesis of phospholipids Component of pitrate inductible formate debydropomase
DgkA	-0,283	Diacylglycorol kinase Cytochrome b561
Cyba AscF	∸0.282 -0.282	Cytochrome prof Transport of beta-gluposide sugars
DedA SecY	-0.280 -0.277	7 Involved in storein export
PhrE	-0.276	Transport of phosphate Component of the aerobic glycerol-3-phosphate dehydrogenase
GlpG NhaE	-0.273 -0.272	Franchort of sodium
BylF	-0.259	Transport of beta-glucoside sugars
DabB CysW	-0.254 -0.254	? Transport of sulfate and thiosulfate
AppC	-0.253	Component of the third cytochrome oxidate (putative)
BetT Nau£	-0.244 -0.242	Transport of choling Transport of N-acetylglucosamine
PatB	-0.241	Transport of N-acetylglucosamine Component of the pyridine nucleotide transhydrogenase Component I of Sytchinom of
CydA MalP	-0.240 -0.239	Transport of Editose
Srlh	-0.225	Transport of D-glucitol Transport of copper
Cut2 Man2	-0.224 -0.320	Transport of manage
PhnQ	~0.219	Transport of phosphate Regulation of the lenght and number of typel fimbrine
FimH FruA	-0.217 -0.210	Pransport of fructose

These integral membrane proteins are involved in transport, anchoring of dehydrogenases, and synthesis of lipid bilayer components.

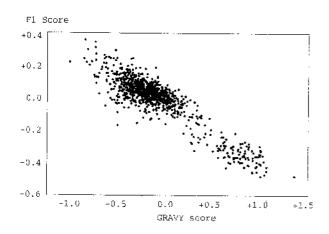
#### Factor 1 (F1)

The first, and thus most important, factor of the correspondence analysis accounted for 17% of the total variability of amino-acid composition of E.coli proteins. The protein F1 scores had a bimodal distribution (Figure 2), indicating that the amino-acid frequencies in the dataset were heterogeneous. The minor peak (11%) contained only integral membrane proteins (Table 3).

Factor 1 was highly correlated (r = 0.90,  $p < 10^{-4}$ ) with the GRAVY score (figure 3). Direct comparison of the GRAVY score and the F1 score coefficients (figure 4) showed a major difference only for Trp. Another difference is that the GRAVY scale assigns the same value to Glu, Gln, Asp and Asn. The coefficients for Glu, Gln, Asp were found to be quite similar in the F1 score, but the coefficient for Asn was different.

### Factor 2 (F2)

The second factor accounted for 13% of the variability in aminoacid compositions. Protein scores on this second factor had a



**Figure 3.** Correlation of the global hydrophobicity of proteins (GRAVY score) with the correspondence analysis factor 1. Each point represents a protein, the bottom right group is the integral membrane protein group.

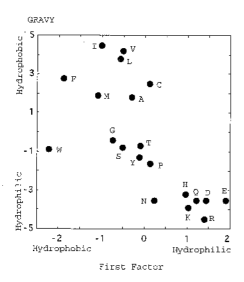
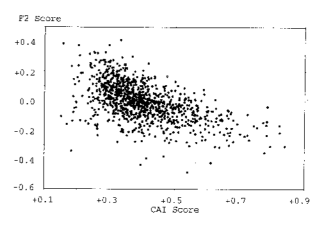
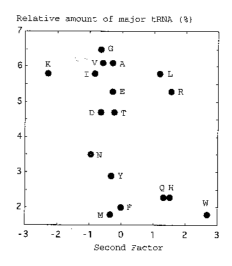


Figure 4. The coefficients for the GRAVY score and for the correspondence analysis factor 1, for the 20 amino-acids.



**Figure 5.** Correlation of the codon adaptation index (CAI) with the correspondence analysis factor 2. Each point represents a protein. Highly expressed genes have a high CAI value.



**Figure 6.** The intracellular concentrations of the major tRNA of amino-acids (36) and the coefficient for correspondence analysis factor 2. The concentrations of the major tRNA for Ser, Pro and Cys were not determined.

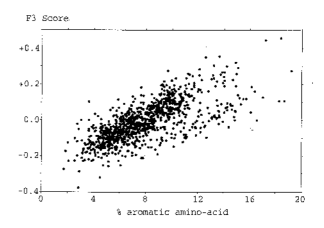


Figure 7. Correlation of the aromaticity with the correspondence analysis factor 3. Each point represents a protein.

unimodal, nearly symmetrical distribution. The F2 scores were correlated (r = 0.55, p <  $10^{-4}$ ) with the CAI scores (Figure 5). The general trend was that proteins with a high CAI value had low F2 scores. This result is highly surprising as CAI score is almost independent of amino-acid composition of the protein: CAI measures the codon usage bias cumulated for each amino acid. Hence, amino acid composition correlates with the choice of codon among synonymous sets.

A comparison of amino-acid F2 coefficients and major tRNAs concentrations (Figure 6) showed three notable exceptions. Lys was more enriched than expected from the relative frequency of its major tRNA, Leu and Arg were avoided despite the relative abundance of their major tRNA.

#### Factor 3 (F3)

The third factor accounted for 10% of the variability in amino-acid compositions. Protein scores on this second factor had a unimodal, nearly symmetrical distribution. The F3 scores were correlated (r=0.70,  $p<10^{-4}$ ) with the aromaticity scores (Figure 7). The general trend was that proteins enriched in aromatic amino-acids had high F3 scores.

## DISCUSSION

The pattern of amino-acid usage was very different from the pattern of codon usage. Analysis of the codons in the coding sequences of *E. coli* emphasises the contrast between lowly and highly expressed genes, with the optimal codons in highly expressed genes. But, as the table of amino-acid frequencies is obtained directly from the table of codon frequencies by summing columns, it seems surprising that the factors reported here have not been described before. One reason is that the column summing which transforms the codon frequency table into the amino-acid frequency table is very special in that frequent codons are summed with rare codons. As the contrast between rare and frequent codons is very important, the amino-acid tendencies are hidden in the least important factors of the codon multivariate analysis.

Integral membrane proteins are known to be enriched in hydrophobic amino-acids. Our correspondence analysis confirmed this and showed that this is the most important factor underlying variations in the global amino-acid composition of *E.coli* proteins.

As factor 1 clearly discriminates integral membrane proteins from the others, computing its value for a new open reading frame could indicate if it codes for an integral membrane protein (a complete example of the computation is given in Table 2). For instance, the protein CutE involved in copper transport in *E. coli* has an F1 score of -0.22. This suggests that it is an integral membrane protein, and not an intracellular protein (27). This prediction of an integral membrane protein is expected to occur for  $1/10^{\text{th}}$  of the *E. coli* coding sequences. Peripheral membrane proteins cannot be identified on the basis of their average aminoacid frequencies because the contribution of membrane-spanning segments to the overall amino-acid composition of the protein is not always sufficient (28).

Factor 2 showed that there was a bias in amino-acid usage for highly expressed genes. There is experimental evidence that the total amount of tRNA for a particular amino-acid parallels the total usage of that amino-acid in proteins for *E.coli* and *Mycoplasma capricolum* (29). Our results also show that proteins encoded by highly expressed genes tend to use amino-acids whose

major tRNA are abundant. This bias is not negligible, since it is the second factor accounting for variability of the amino-acid variability of E.coli proteins. This bias was previously observed in studies on much smaller samples of E.coli proteins (30-32).

Table 4. Last 10% of CAI distribution

Prot.	CAI	function
		·· ·· ·
GapA	0,840	g:yeeraldehyde-3-phoaphate dehydrogenase (glycolysia)
Omp√ TufA	0.824 0.822	mnjor outer membrane protein (porin) elongation factor EF-TO (translation)
McpA	0.794	elongation tactor EF-TU (translation) refolding of protein under stress condition (translation related) ribosomal protein 59
R⊭sI	0.794 0.785	riboscmal protein 59
Pf1	U.784	DALPAGE IDENTES-TARRE (NOUOXIGGIAA CONADERION OF GIACORE)
RpsA	0.784 0.782	ribosomal protein \$1 ribosomal protein 52
Tst	0.777	elongation factors EF-Ts (translation)
СпрА	0.772	major outor membrane protein (porin) elongation factor EF-G (translation)
Fush TpiA	0.743	elongation Eactor EP-G (CTADSIACION) triosephoaphate isomerase (glycolysis)
RplI	0.729	ribosomel protein L9
SodA	0.724	mandanese superoxide dismutase (radicals destruction)
DnaK Tig	2.723	Major heat shock protein (DNA replication) chaperone (protein export)
PykF	0.715 0.701	removate kinase 7 (mlycolycis)
Pal	0.692	pertidoglycan associated lipoprobuin (structure) polytucleotide phosphorylase: mRNA dogradation (transcription)
Pap PpiB	0.68D 0.679	polyhucleotide phosphorylase: mkWA dogradation (transcription) peptidyl-prolyl dis-brans isomerase (protein folding)
Rp17	0.675	riboscmal protein L20
GlvA	0.674	sarina hydroxymethyltransferase (purings & lipids synthesis)
RplM	0.574 0.573	ribosomal protein L13 pyrevate dehydrogenase (glycolysis)
AceE AtpA	0.571	ATP synthase alpha chain (ATP synthesis)
Rp10	0.571	ribosomal protein L15
DeoD	0.568	puring nucleoside phosphorylase
OutpF ResF	0.667 0.665	major outer membrane protein (porin) ribusomal protein S6
AckA	0.665	acetate kinase (anaerobic growth: acetate production)
Ppa	0.664	inorganic pyrophosphatase ribosomal protein S12
Rpsi. LpdA	0.662	ricosomai protein 512 dilydrolipoamide debydrogenass (glycolysis)
AtpD	0.650	ATP synthese beta chain TATF synthesis)
Adhg	0.659	alcohol dehydrogenase (anacrobic growth in absence of nitate)
Adk FabB	0.652 0.646	adenylate kinasé 3-oxonoy1 ACP synthase I (lipids synthasis)
Ndk	0.646	nucleoside diphosphato Kibase
Rpis	0.643	ribosomal protein L19
Tsx GuaB	0.643 0.640	nucleoside-specific channel-forming protein (transport)
DeoC	0.639	IMP dehydrogenase (GMP synthesis) deoxyribose phosphate aldolase ( deoxy]nucleotide catabolism)
Such	0.637	succinvi-CoA symbhetaso wipha-subunic (TCA Cycle)
PurA	0.636	adenylonuccinate synthetase (AM2 synthesis)
AspA GlnA	0.633 0.629	aspartose glucamine synthetase (amino-acid synthesis)
Vals	0.626	valyl tRNA synthetese (translation) extragenic suppressor (heat shock protein related)
SultB	0.626	extragenic suppressor (heat shock protein related) single-strand DNA-binding protein (DNA replication)
Sab AceF	0.524 0.524	pyruvate dubydrogenase (glycolysis)
RecA	0.621	50S response
H1pA	0,519	histone like protein SLP-1 (structure)
HtpG AspS	0.617 0.613	chaperone (heat shock protein) aspartyl-tRNA synthomese (translation)
PfkA	0.612	<pre>6-phosphe(ructokinase (qlycolysis)</pre>
Cer	0.611	PTS enzyme III glg (transport) ribosomal protein LS
RplE Int9	0.605	initiation factor IF2 (translation)
LysS	0.602	lvevi PNN evnihetasa (franslation)
Eda	0.602	2-ketn-3-decxy-6-phosphogluconate aldoisse
Rp1R LauS	0.599	ribosomal protein 519 leucyl-tRNA synthetase (translation)
GunA	0.599	CMP synthetase (GMP synthesis)
Rpo⊃	0.598	RNA polymerase sigma-subunit (transcription)
Su¢C PatS	D.594 D.594	succinyl-CoA synthetase heta-subunit (TCA cycle) phosphate-specific transport system (transport)
rats RpsK	0.594	ribosomal protein Sil
DebA	0.592	ribosomal protein Sil required for disulphide bond formation
Cadh SecB	0.592	lysine decarboxylass (cadaverine production at tow pu)
Dear	0.588	protein export protein (transport) RNA helicase (ribosome assembly)
RplF	0.585	ribosomal proteit L6
RpsE	0.5\$3 0.581	sibosomal protein 85 glutamyl-tRNA synthetase (translation)
GltX SucB	0.580	dibydrolipoamide succinyltransferase (glycolysis)
LcdE	0.579 0.579	2
Ebs	0.579	Dhistone like protein MLP-2 (structure) ribosme-releasing factor (kranslation)
Frr FldA	0.577	flavodoxin (electron transport)
FrdB	0.571	Eumerate reductase iron-sulfur subunit (TCA cycle)
Glys	0.571	glycyl-ERNA synthetase (translation)
Nira FrdA	0.570	NaDH-dependent mitrito reductase (mitrate assimilation) furarate reductase (TCA cycle)
AccC	0.568	Renards reductase (TCA cycle) biotic carboxylase (falt, acid synthesis) maltone traitclestrine; transport
LamB	0.568 0.568	maltose et maltodextrines transport
MetK NapC	0.568	S-adenceylmathionics synthetase (one carbon metabolism) outer membrane porin of endogenous lambdold bacteriophage
Pgi	0.566	
NarH	0.566	mannitol permease (transport)
MUTA	0.564 0.564	mannitol permease (transport) PTS glucose-specific enzyme IT (transport)
PtsG RpsH	0.563	ribosepal protein 98
RogR	0.562	- PMS notimerace cimma-32 subunit theat shock proboders expression:
Apt	0.562 0.562	adening phosphoribosyl-transference (purine salvage) carbamoyl-phosphate synthetase (Arg 5 pyrimidibe synthesis)
CarB	0.552 0.561	cerbambyr-phosynate synthetase (Arg & pyrintsine synthetis) superoxide dismutane (radical destruction)
GyrB	0.561	superoxide dismutane [redical destruction] DNA gyrase (DNA replication)
Rp10	0.561	riboscmal protein L17
CydB Prs	0.558 0.558	cytochrome D terminal exidase (electron transport) phosphoribosylpyrophosphate synthetase (mucleotide biosynthesis)
	_	

The codon usage of the corresponding genes is good, so that their expressivity level is expected to be high. For instance, the genes for ribosomal proteins, major outer membrane proteins or basic metabolism such as glycolysis belong to this class. Note that genes that are only nurned on under special environmental conditions but are abundantly expressed under those circumstances are also present in this class (e.g. AckA and AdhE in anaerobiosis, SubH and RpoH after heat shock, NarH in presence of nitrate).

The concentrations of the major tRNA for Lys, Leu and Arg did not follow the general trend. The concentration of the major tRNA for Lys was less than expected and the concentrations of the major tRNAs for Arg and Leu were higher than expected. The concentrations of the major tRNA for Leu and Arg may appear high because their intracellular concentrations do not correspond to their effective availability to the ribosome. For instance, two minor leucyl-tRNAs species are the ones most

Table 5. First 10% of CAI distribution

Prot.	CVI	FUNCTION
Rfab	0.151	O Antigeb ligase (LPS core synthesis)
AppY Rfas	0.169	transcriptional regulator LPS core synthesis
RfaK TrkG	0.136 0.188	LPS core synthesis
TdcR	0.189	integral mumbrane protein involved in potassium uptake positive regulatory protein of the the operen
FgpA McrC	0.192	Membrane-Bound phosphatidyl glycerophosphate phosphatase Moddfiles the specificity of McrB restriction resistance against methyl viologen toxicity
MvrC McrA	0.194 0.197	resistance against methyl viologen toxicity methyl cytosine restriction enzyme
PhnQ Lit	0.197	bypothetical protein Elocks backnricphage T4 late gene expression
DacB	0.203	D-alanyl-D-alanine carboxypeptidase in murein metabolism (PBP4)
DsdC FitnB	0.204	Spansoription activator type I fimbriae regulatory protein
ThdF FimZ	0.207	thiophene oxydation regulatory protoin
FecE DicA	0.211	citrate dependant Fe3- transport repressor of division inhibition gene dicB
CynR	0.216 0.217	transcriptional activator for the cyn operon
Rosa Araj	0.226	transcriptional activator of capsular polysaccharida symbhasia
Bg1G RfaZ	0.233 0.235	transport or processing of arabinose polymers positive regulator of bgl operon LPS core synthesis
Pin	0.236	DNA-invortano
Fu∈U	0.237 0.237	? uroporphyrinogen III cosynthetase
PriB RnpA	0.238	primosomal realization protein protein component of ribonuclenses F
ЭдрВ	0.241	phosphatidylglycerophosphate phosphatase B
BarA HipB	0.241	OmpR activator
Sulh REal	0.242	UV-indumible call division inhibitor LPS core synthesis
CysX	0.245	hypothetical protein
Fing UmbC	D.245 0.246	type 1 fimbriae regulatory protein UV repair enzyme
Iap ⊂dh	$0.246 \\ 0.248$	conversion of alkaline phosphatase isozyme CDP-diglyceride hydrolase
AVEA	0.249	Alanino-valino transaminase
RfaP HadS	0.250 0.250	LPS core synthesis EcoE type I restriction-modification enzyme S subunit
BtuC CreB	0.251 0.252	cytoplasmic membrane protein involved in vitamin B12 transport transcriptionnal regulatory protein
LysR	0.253	activetion of lysa trenscription
ProV Rhas	0.253 0.254	transport of glycine betaine/L-proline positive activator of genus required for L-rhammose utilization
FimH DicB	0.254 0.254	regulation of length and mediation of adhesion of type 1 fimbries inhibition of cell division
KfaJ UbiC	0.256	LPS core synthesis choriskate lyase (ubiquinone synthesis)
TdcA	0.257	transcriptional activator for tdc operon
EnvY SrlM	0.258	porin thermoregulatory protein positive regulator for glucitol operon
KgtP AppA	0.260 0.260	alpha-ketoglutarate transport acid phosphatase
MiaA	0.260	(delta)2-isopenteny1 pyrophosphate tRNA transferase
ÇadÇ Mali	0.260 0.261	transcriptional activator repressor protein for maltose regulon
OmpT Bicc	0.261 0.262	outer membrane protease involved in biotin synthesis pathway
LacA	0.263	thiogalactoside transacetylase sequence-specific restriction of cytosine-modified DNA
Morb NlpA	0.263 0.263	zykopplasmic membrane llyrotein
BicB RfaY	0.264	hypothetical protein LPS core synthesis
GlpR BtuD	0.265 0.266	represent of alveerel 3 throughste regular
FepD	0.266	peripheral membrane component of vitamin 312 transport system ferric enterobactin transport protein
RfaB CybB	0.266 0.267	LPS core synthesis protoin
llyaF ArcL	0.267 0.267 0.268	cytochrome b561 protein of hydrogenase-1 operon shikimate kinase II
Flis	0.268	flagellar protein
Pcm PhợQ	0.268 0.268	flagellar protein L-isoaspartyl grotein carboxyl methyltransferase type II regulation of acid phosphatase
Ogt Trg	0.269	O-6-alkylguanine-DWA-alkyltransferase sensory transducer protein
CreD	0.269	?
AraC CreC	0.270	regulatory protein regulation of GreB
Fes Tdk	0.272	enterochelin esterase thymidino kinaso
Nip: SoxR	0.274	lipoprotein regulatory protein for superoxide strength response
Mct3	0.277	centrel of chemotaxis
Dgt TyrP	0.277	dGTPase transport of Tyr
Cŷs2 Rπ⊏	0.278 0.278	AFS klnase ribonuclosso III
MetR	0.279	regulatory protein protein of hydrogenase-1 operon
AyaD MatB	0.279 0.279	DNA mismatch repair
·Cysa EntD	0.280 0.280	regulatory protein enterobactin synthesis
Bola BirA	0.280	control of cell morphology biotim operon-represent and biotim holoenzyme synthetase
Fhr	0.283	deoxyribopyrimidine photolymse
AroE RecF	0.282 0.283	shikimate dehydrogenese control of cocombination

The codon usage of the corresponding genes is poor, so that their expressivity level is expected to be low. For instance, many regulatory genes belong to this

bound to ribosomes during exponential growth in minimal medium (33). The difference between the effective and measured tRNA concentration could be attributed to the participation of the major leucyl-tRNA species in a reaction other than translation, such as the addition of leucine directly to the amino termini of certain ribosomal proteins (34). This would explain why the effective concentrations of the major tRNA for Leu and Arg could be overestimated from their intracellular concentrations, but does not explain the case of the major tRNA for Lys. However, The comparison of tRNA concentrations from differents authors (35, 36) introduces a note of caution with respect to the interpretation of quantities of tRNA in cells.

To validate the interpretation of factor 2, the first and last 10% of the CAI distribution were extracted (Table 4 and 5), and the mean major tRNA frequencies for the proteins were computed in these two extreme classes. The distributions for the two classes were different (figure 8), showing that proteins with high CAI values are enriched in amino-acids carried by the most abundant major tRNA.

Further discussion about the bias in the amino-acid composition of proteins encoded by highly expressed genes should be taken with care because they are based on a logical construction and cannot be directly challenged by experiment. At first glance it seems that it is simpler for tRNAs to adapt their concentration to the amino-acid content of proteins than the reverse because the mutation expense is lower; changing the tRNA concentrations

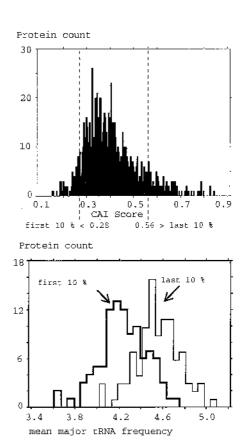


Figure 8. Top: distibution of CAI values for the 999 protein genes of the dataset. The dotted lines indicate the first and last 10% of the distibution. Bottom: distribution of the mean major tRNA frequency for the proteins of the first and last 10% of the CAI distribution.

requires fewer mutation events, such as gene duplication or altered promoter efficiency, than does altering coding sequences, where many sites must be modified. But this cannot explain why the amino-acid compositions of the product of highly expressed genes should be different. This requires that the amino-acid composition of highly expressed genes is particular for some other reason. The simplest explanation is a straightforward adaptation of what is visible at the codon level: highly expressed genes reduce the diversity of codon choices to increase translation efficiency (4). By analogy, proteins encoded by highly expressed genes use a reduced diversity of amino-acid choices to increase translation efficiency.

The fact that proteins encoded by highly expressed genes have a bias of amino-acid usage is an interesting example of the interdependence between translational constraints and overall properties of the protein. The translational constraints seem to be greater than expected since, in addition to selecting the codon corresponding to the most frequent isoacceptor tRNA, they are sufficient to modify the global amino-acid composition. The translational constraints which were known to affect the 'genotype' of proteins, are sufficient to affect their 'phenotype'.

Factor 3 showed that aromatic amino-acids represent a group of amino-acids which frequency is highly variable among proteins. An interpretation is that the biosynthesis of these aminoacids is expensive for the cell, so that there is a selective pressure to reduce the aromaticity of proteins. The fact that these aminoacids are rare (Table 1) is consistent with this hypothesis. However, these amino-acids do not completly disappear, so that there should be an inverse tendency to maintain them in proteins. This inverse tendency could be attributed either to a simple mutationnal drift or more likely to a selective advantage due to a contribution to the stabilization of the three-dimensional structure of the protein.

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