Abstract

Synonymous codons, i.e., DNA nucleotide triplets coding for the same amino acid, are used differently across the variety of living organisms. The biological meaning of this phenomenon, known as codon usage bias, is still controversial. In order to shed light on this point, we propose a new codon bias index, CompAI, that is based on the competition between cognate and near-cognate tRNAs during translation, without being tuned to the usage bias of highly expressed genes. We perform a genome-wide evaluation of codon bias for E. coli, comparing CompAI with other widely used indices: tAI, CAI, and Nc. We show that CompAI and tAI capture similar information by being positively correlated with gene conservation, measured by the Evolutionary Retention Index (ERI), and essentiality, whereas, CAI and Nc appear to be less sensitive to evolutionary-functional parameters. Notably, the rate of variation of tAI and CompAI with ERI allows to obtain sets of genes that consistently belong to specific clusters of orthologous genes (COGs). We also investigate the correlation of codon bias at the genomic level with the network features of protein-protein interactions in E. coli. We find that the most densely connected communities of the network share a similar level of codon bias (as measured by CompAI and tAI). Conversely, a small difference in codon bias between two genes is, statistically, a prerequisite for the corresponding proteins to interact. Importantly, among all codon bias indices, CompAI turns out to have the most coherent distribution over the communities of the interactome, pointing to the significance of competition among cognate and near-cognate tRNAs for explaining codon usage adaptation. Notably, CompAI may potentially correlate with translation speed measurements, by accounting for the specific delay induced by wobble-pairing between codons and anticodons.

Introduction

The genetic information carried by mRNA and then translated into proteins is encoded into nucleotide triplets called codons. Four alternate nucleotidic bases (A,U,C,G) compose mRNA, so that there are $4^3 = 64$ possible codons that have to code for only 20 naturally occurring amino acids. The genetic code is therefore redundant: while a few amino acids correspond to a
single codon, most amino acids can be encoded by different codons. Different codons coding for the same amino acid are known as synonymous codons, and in a wide variety of organisms synonymous codons are used with different frequencies—a phenomenon known as codon bias. With the advent of whole-genome sequencing of numerous species, genome-wide patterns of codon bias are emerging in the different organisms. Various factors such as expression level, GC content, recombination rates, RNA stability, codon position, gene length, environmental stress and population size, can influence codon usage bias within and among species [1]. While the biological meaning and origin of codon bias is not yet fully understood, there is a large consensus that the degeneration of the genetic code might provide an additional degree of freedom to modulate accuracy and efficiency of translation [2]. Indeed, population genetic studies [3] have shown that synonymous sites are under weak selection, and that codon bias is maintained by a balance between mutation-selection (random variability in genetic sequences followed by fixation of the optimal codons) and genetic drift (allowing for the occurrence of non-optimal codons). In fact, highly expressed genes feature an extreme bias by using a small subset of codons, optimized by translational selection [4–6]. On the other hand, the persistence of non-optimal codons in less-expressed sequences causes long breaks during protein synthesis; this could be the result of genetic drift and have a key role in the protein folding process [7, 8]. In addition, codon usage appears to be structured along the genome, with neighboring genes having similar codon compositions [9], and codon bias seems positively correlated to gene length (as a result of selection for accuracy in the costly production of long proteins) [10]. In the last years there has been a wide effort in developing effective ways to measure codon bias [11]. The most widely used indices include the Codon Adaptation Index (CAI) [12], the tRNA Adaptation Index (tAI) [13], and the Effective Number of Codons (Nc) [14], each of them having specific advantages and drawbacks. For instance, CAI and tAI correlate well with gene expression levels, however such correlation is a natural consequence of their definition: they are tuned on a reference set of highly expressed genes. Nc is instead basically a measure of the entropy of the codon usage distribution, and thus shows a lower correlation with expression levels.

In this work we propose a novel codon bias index named Competition Adaptation Index (CompAI), which does not rely on information about gene expression levels, but instead has a self-consistent biological meaning—based on tRNA availability and competition between cognate and near-cognate tRNAs. In other words, CompAI is a parameter-free index that does not require a set of reference genes for its calibration, a fact that constitutes its main advantage with respect to CAI and tAI. Moreover, CompAI is designed to extract genetic signals that could be directly correlated to experimental measures for translation speeds, an emerging and challenging issue still to be explored. In order to show the advantage of the novel codon bias index, we perform a genome-wide comparison of CAI, tAI, Nc and CompAI for Escherichia Coli (E.coli). Our analysis reveals that the information on gene conservation across species and gene essentiality is better captured by codon bias metrics that build on tRNA availability (tAI and CompAI). We also study codon bias in relation to the connectivity patterns of the protein-protein interaction network (PIN) [15] of E.coli. We thus show that translational selection systematically favors proteins with the highest number of interactions and belonging to the most densely connected community of the network, at least when the bias is measured by CompAI and, to a smaller extent, by tAI. Additionally, we address the issue of how much a similarity in the codon usage bias of a set of genes is reflected on the interactions among the corresponding proteins. A principal component analysis for the variability of codon bias indices indeed reveals that closeness of a set of genes in the space of the two principal components likely results in the corresponding proteins to interact—in comparison with an appropriate null model.

Overall, our study reveals that CompAI captures more information than the other indices about the connection between codon bias and the topology of the interactome. Besides, we
recall that CompAI does not require calibration on gene expression levels and has a consistent biological meaning based on the competition between cognate and near-cognate tRNAs. These observations stress the potential of the new index to both measure and explain codon usage bias, particularly as related to speed and accuracy of gene translation and protein synthesis.

**Materials and Methods**

**Sequences**

In this work we investigate the genome of E.coli K-12 substr. MG1655, whose 4005 coding mRNA sequences have been collected from GenBank [16]. The gene copy numbers coding for each tRNA (tGCN) were derived from the Genomic tRNA database [17].

**Conservation and Essentiality of E.coli genes**

In order to have an index for gene “conservation”, we use the normalized Evolutionary Retention Index (ERI) [18]: for each gene in E.coli’s genome, its ERI measures how much that gene is shared among other 32 bacterial species (having at least an ortholog of the given gene). A low ERI value thus denotes that a gene is specific to E.coli, whereas, high ERI is characteristic of highly shared (and therefore conserved) genes. Concerning gene “essentiality”, we use the classification of Gerdes et al. [18] for the E.coli genome into 606 essential and 2940 non-essential genes, based on experimental measures of gene resistance against transposon insertion.

**Codon Bias Indices**

Codon usage bias can be assessed, for each gene in a given genome, by various indices that can be classified into broad groups based on: (i) codon frequencies; (ii) reference gene sets; (iii) deviation from a postulated distribution; (iv) information theory; (v) interactions among tRNAs (see [11] for an overview). We focus here on the most widely used indices: tAI [13], that belongs to groups (ii) and (v) by requiring calibration on a set of highly expressed genes; CAI [12], a group (i) and (ii) index built on local statistics of codon usage and on a reference list of optimally expressed genes; Nc [14], a group (i) index based on the number of different codons used in a coding sequence. The novel codon bias index we propose in this work, CompAI, is instead based on the competition of cognate and near-cognate tRNAs to bind to the A-site on the ribosome during translation, and is thus a group (v) index that does not need tuning on a reference set of highly expressed genes. While the formal definition for CompAI and the rationale behind are given below, we refer to the S1 File for the definition of CAI, tAI and Nc.

**Competition Adaptation Index (CompAI).** It is generally accepted that translation speed depends on the efficiency of the codon/anticodon pairing in the A site of the ribosome [19]. Hence, for a given codon, the rate of amino acid synthesis is essentially influenced by two dominant processes: the number of collisions of the corresponding tRNA with the ribosomal A site (which strongly depends on tRNA concentration in the cell) and the specificity of the codon-anticodon pairing. Such a pairing process satisfies the Watson-Crick (WC) base-pairing rules (G-C and A-U, and vice versa) for the first two bases, whereas, the rule on the third (or wobble) base is more relaxed and non-standard pairing is allowed in some cases (wobble complementarity) [20]. Hence, there are cases in which several tRNAs pair with the same codon (provided that these are identical in the first two bases) and are called isoacceptor or cognate tRNAs. Codon-anticodon interactions are thus characterized by competition between cognate isoacceptor tRNAs (with WC or wobble complementarity between mRNA codon and tRNA anticodon) on one side and non isoacceptor tRNAs on the other side: near-cognate (with a mismatch in only one of the first two bases) and non-cognate (with at least two mismatches). Discrimination
between correct and wrong tRNA according to base pairing features very high fidelity (error rate $f \sim 10^{-3} \div 10^{-4}$). Rejection of the wrong tRNA can occur in two distinct phases [19, 21]: initial selection of the ternary complex EF-Tu-GTP-aa-tRNA and subsequent proofreading of aa-tRNA after GTP hydrolyzation. The first interaction is fast and does not depend on the choice of codons, in order to allow the ribosome to quickly screen for the available tRNAs. The second step is instead sensitive to base complementarity, featuring the first selection between cognate and near-cognate tRNAs: non-cognate are excluded almost immediately with $f \sim 10^{-1}$, and then a more strict and efficient proofreading takes place, excluding near-cognate with $f \sim 10^{-2}$. This means that near-cognate tRNA (unlike non-cognate tRNA) can enter into the interaction process between the ternary complex and the site A of the ribosome, and (when not accepted, in very few cases) can be rejected at the stage of initial recognition or during proof-reading. In any event, this process results in a time delay of translation, because near-cognate rejection brings the ribosome back to the initial state of waiting for the correct tRNA.

The rationale behind the definition of $\text{CompAI}$ is precisely that of building an index which is based both on tRNA availability and on competition between cognate and near-cognate tRNAs that could modulate the speed of translation of mRNAs into proteins. Note that, since in vivo experimental determinations of tRNA concentrations are available only for few organisms, we will implement $\text{CompAI}$ using the number of tRNA gene copies (tGCN) which, at least in simple organisms, has a high and positive correlation with tRNA abundance [22–25] (a similar approach is adopted in the definition of $\text{tAI}$ [13]). For each codon $i$ we define its absolute adaptiveness value ($W_i$) as:

$$W_i = \left( \sum_{j=1}^{m_i} \text{tGCN}_{ij} \right) \left( \sum_{j=1}^{m_i} \text{tGCN}_{ij} + \sum_{j=1}^{m^*} \text{tGCN}_{ij} \right) \right)$$

Here $m_i$ is the number of isoacceptor tRNA sequences (anticodons) that recognize codon $i$ (i.e., containing either the anti-codon $i$ or all its cognates that are read by $i$) and tGCN$_{ij}$ is the gene copy number of the $j$-th of such tRNAs, whereas $m^*$ is the number of tRNA sequences that are near-cognate of $i$ and tGCN$n_{ij}$ is the gene copy number of the $j$-th of such tRNAs (see also S1 Fig). The amount in square brackets represents a penalty introduced by the competition with near-cognate tRNAs, assuming unit or zero values in the cases of smaller and higher competition, respectively. This term thus assumes the role of selective constraint on the efficiency of the codon-anticodon coupling. Importantly, and at odds with $\text{tAI}$, these terms do not result from optimization on expression levels, but have a biological justification based on cognate/near-cognate competition. Note that, in the computation of $W_i$ for a given codon, we count as isoacceptor tRNAs those with WC or wobble base pairing that also carry the same amino acid of $i$’s anticodon. Computation of $\text{CompAI}$ continues by defining for each codon $i$ its relative adaptiveness value $w_i = W_i/W_{\text{max}}$, where $W_{\text{max}}$ is the maximum value between all the $W_i$ of codons. $\text{CompAI}$ of gene $g$ is finally defined as the harmonic mean of the relative adaptiveness of its codons:

$$\text{CompAI}_g = \frac{1}{\sum_{j=1}^{L_g} w_j^{-1}}$$

The choice of the harmonic mean (rather than geometric as for $\text{CAI}$ and $\text{tAI}$) is consistent with
the association of CompAI with the speed of protein synthesis. Indeed, the translation speed of codon \( i \) can be defined as the reciprocal of the concentration of the corresponding tRNA isoacceptors \([26]\). Therefore, if codon \( i \) is read at a speed proportional to \( w_i \), then the average translation speed of a gene is given by the harmonic mean of the \( \{w_i\} \) associated to its codons. CompAI takes values between 0 and 1, where values close to 0 (1) indicate highest (lowest) competition, and therefore a low (high) translation rate.

**Protein-Protein Network Analysis**

In this study we use protein interaction data collected in STRING (Known and Predicted Protein-Protein Interactions) \([15]\). In such database, each predicted interaction is assigned with a confidence level or probability \( w \), evaluated by comparing predictions obtained by different techniques \([27–29]\) with a set of reference associations, namely the functional groups of KEGG (Kyoto Encyclopedia of Genes and Genomes) \([30]\). In this way, interactions with high \( w \) are likely to be true positives, whereas, a low \( w \) likely corresponds to a false positive. Since the percentage of false positives can be very high \([31]\), we select a stringent cut-off \( \Theta = 0.9 \) that allows a fair balance between coverage and interaction reliability (see the probability distribution \( P(w) \) in the left panel of S2 Fig). We thus build the protein-protein interaction network (PIN) of E. coli by placing a link between each pair of proteins (nodes) \( i, j \) provided that \( w_{ij} > \Theta \). The resulting number of connections or degree for a given protein \( i \) is denoted as \( k_i \).

To detect communities of PIN we resort to Molecular Complex Detection (MCODE) \([32]\). In a nutshell, MCODE iteratively groups together neighboring nodes with similar values of the core-clustering coefficient, which for each node is defined as the density of the highest \( k \)-core of its immediate neighborhood times \( k \). The density of a graph \( G \) with \( n \) nodes and \( l \) links is the ratio between \( l \) and the maximum number of possible links, namely \( n(n - 1)/2 \), whereas, a \( k \)-core is a graph \( G \) of minimal degree \( k \), meaning that each node belonging to \( G \) has degree greater or equal than \( k \). MCODE detects the densest regions of the network and assigns to each found community a score that is its internal link density times the number of nodes belonging to it. (Note that communities found by MCODE can overlap, and some nodes can be excluded from all communities: there is no strict partition of the network, which would have been at least questionable since the same protein can be involved in different metabolic processes and thus belong to more than one community. For these reasons and also because it considers both density and connectivity, MCODE performs much better for our purposes than other community detection methods like modularity maximization \([33]\) that are based only on intra/inter-community densities and create a strict partition of the network). We also characterize each found community \( c \) with the mean value \( \bar{x}_c \) and standard deviation \( \sigma_c \) of codon bias values within the community, and use them to compute a Z-score as \( Z_c = (\bar{x}_c - \bar{x}_n)/\sqrt{\sigma_c^2 + \sigma_n^2} \) (where \( \bar{x}_n \) and \( \sigma_n \) are, respectively, the mean value and standard deviation of codon bias values computed on the whole network). In this way, a value of \( Z_c > 1 \) (\( Z_c < -1 \)) indicates that community \( c \) features significantly higher (lower) codon bias than the population mean.

Finally note that each node of PIN is philogenically classified according to the Clusters of Orthologous Groups (COGs) of proteins \([34]\). COGs are generated by comparing predicted and known protein sequences in all completely sequenced genomes to infer sets of orthologs. Each COG consists of a group of proteins found to be orthologous across at least three lineages and likely corresponds to an ancient conserved domain \([34]\).

**Principal Component Analysis**

Principal Component Analysis (PCA) \([35]\) is a multivariate statistical method to transform a set of observations of possibly correlated variables into a set of linearly uncorrelated variables
(called principal components) spanning a space of lower dimensionality. The transformation is defined so that the first principal component accounts for the largest possible variance of the data, and each succeeding component in turn has the highest variance possible under the constraint that it is orthogonal to (i.e., uncorrelated with) the preceding components.

We use this technique on the space of codon bias indices, so that each gene of E.coli is represented as a 4-dimensional vector with coordinates (CompAI, CAI, tAI, Nc). Such coordinates are separately normalized to zero mean and unit variance over the whole genome. We then obtain the associated covariance matrix between the four dimensions of codon bias and diagonalize it. The eigenvectors of the covariance matrix, ordered according to the magnitude of the corresponding eigenvalues, are the principal components of the original data.

**Configuration Model**

In order to assess how significant are the codon usage patterns observed for the PIN, we need to compare the E.coli interactome with a suitable null model for it, i.e., an appropriate randomization of the network. Here we follow the most common approach in statistical mechanics of networks of using the **Configuration Model** (CM) [36]. The basic idea is to build the null model as an ensemble $\Omega$ of graphs with maximum entropy, except that the ensemble average of the node degrees are constrained to the values observed for the real network: $\langle k\rangle_{\Omega} \equiv k_i \forall i$. This leads to a probability distribution over $\Omega$ which is defined via a set of Lagrange multipliers $\{x_i\}$ (one for each node) associated to the constraints [37]. Once all $\{x_i\}$ are found, the CM reduces to having a link between nodes $i$ and $j$ with probability $p_{ij} = \frac{x_i x_j}{1 + x_i x_j}$, independently on all other links. Then, the null hypothesis is that any given network property $\chi$ varies in the range $\langle \chi\rangle_{\Omega} \pm \sigma_{\Omega}[\chi]$, where both average and standard deviation of $\chi$ over the ensemble can be obtained either analytically or numerically (by drawing sample networks from $\Omega$) [37]. The number of standard deviations by which the empirical and expected values of $\chi$ differ is given by the Z-score $Z[\chi] = (\chi - \langle \chi\rangle_{\Omega})/\sigma_{\Omega}[\chi]$: large positive (negative) values of $Z[\chi]$ indicate that $\chi$ is substantially larger (smaller) than expected, whereas, small values signal no significant deviation from the null model.

**Results and Discussion**

**Specificity, Essentiality and Codon Bias of E.coli genes**

**Correlations between Codon Bias indices.** As the starting point of our analysis, we first check how the different codon bias indices correlate over E.coli’s genome. Fig 1 shows that, interestingly, CompAI is strongly (and positively) correlated with tAI, whereas it does not show...
any significant correlation with CAI nor with Nc. This result can be easily explained as CompAI and tAI elaborate on the same genetic information, that is the abundance of tRNAs, whereas CAI and Nc are based on codon usage statistics (see the S1 File).

**Codon Bias and ERI.** We move further and analyze the correlation between the various codon bias indices and the evolutionary retention index (ERI) [18] for E.coli genes (we recall that a gene with a low ERI value is peculiar to E.coli, whereas a gene with high ERI is shared among different species). Fig 2 reports the average values and standard deviations of the codon bias indices for every group of genes having the same ERI value. Interestingly, the evolutionary codon adaptation measured by CompAI and tAI tends to increase for genes that are less specific to E.coli. Fig 2 also suggests that it is possible to make a threefold separation of genes by looking at the rate of variation of tAI and CompAI with ERI. We thus identify group A (ERI < 0.2: 1597 low ERI genes that are specific to E.coli), group B (0.2 < ERI < 0.9: 1804 intermediate ERI genes) and group C (ERI > 0.9: 231 high-ERI genes that are highly conserved and shared among several bacterial species). In each group, the correlation between codon bias and ERI is maximized (see the corresponding correlation coefficients in the figure). CAI and Nc are maximized.
instead less structured with respect to ERI, as shown by the very small correlation coefficients (and by the impossibility to identify gene groups).

**Codon Bias and Gene Essentiality.** We now study the patterns of codon usage bias in essential and non-essential genes, according to the classification scheme of Gerdes et al. [18] (see Materials and Methods). As a preliminary result, Fig 3 reports the percentage of essential genes in each set of genes sharing the same ERI. We see that the three groups A, B, C of genes identified as in the previous paragraph feature different percentages of essential genes: approximately, 10% for group A (ERI < 0.2), 15% for group B (0.2 < ERI < 0.9) and more than 30% for group C (ERI > 0.9).

**COGs.** We now perform a kind of gene ontology to check how the three gene groups A, B, C are projected over the clusters of orthologous genes (COGs) and their functional annotations [34]. To this end, for each group we evaluate the Bayesian probability that its genes belong to a
given COG:

\[
Pr(COG|\text{group}) = Pr(\text{group}|COG)Pr(COG)/Pr(\text{group}),
\]

where \( Pr(\text{group}) \) is estimated as the fraction of the genome belonging to the group, \( Pr(COG) \) as the fraction of the genome belonging to the COG and \( Pr(\text{group}|COG) \) is the fraction of genes in the COG that belong to a particular group. Fig 5 shows the histogram of \( Pr(COG|\text{group}) \) over the 17 COGs, for the three groups A, B, C defined above. Assuming an arbitrary discriminating threshold of 10%, we observe that each group is prevalently projected over a limited set of COGs (reported in the legend box of Fig 5). Group A genes (those with low ERI values) mostly insist over COGs K and G (transcription, carbohydrate metabolism); group B (genes with intermediate ERI) is enriched in COGs G and E (again, carbohydrate metabolism, amino acid metabolism and transport); finally, group C (genes with the highest ERI) is dominated by the functional annotations associated with COGs J and L (translation, ribosome structure and biogenesis, replication, recombination and repair). Indeed, group C, composed of the highly adapted, essential, and conserved genes of E. coli, is the set of genes that code for ribosomal proteins.
Codon Bias and the Connectivity Patterns of E. coli’s Protein Interaction Network

**Communities.** We now turn our attention to the network of interacting proteins in E. coli. We start by studying codon bias in relation with the connectivity patterns of the network. First, note that the degree distribution of proteins is scale-free (see the right panel of S2 Fig), meaning that the network features a large number of poorly connected proteins and a relatively small number of highly connected hubs. Fig 6 notably shows that these hub proteins are systematically characterized by higher values of codon bias of the corresponding genes—when this is measured by \( tAI \) and \( CompAI \). \( CAI \) and \( Nc \) are instead clearly less sensitive to protein connectivity.

We move further and consider codon bias in relation with the community structure of the PIN. We recall that a community is a group of proteins that are more densely connected within each other than with the rest of the network. Table 1 shows the features of the communities that are assigned by MCODE a score higher than 10, together with their COG composition, average degree and, for ERI and the various codon bias indices, the internal average value \( \bar{r} \), and the Z-scores (comparing the distribution of bias inside the community with that of the whole network). We see that such topologically determined communities, ordered by score, are evolutionarily and functionally characterized by a dominant COG, shared by the majority of the proteins in the community. This suggests that the identified communities can be associated with specific metabolic functions: they correspond to functional modules, essential for the life-cycle of the organism.

Let us focus on the first community, that includes only 60 proteins (4.5% of the whole network) but as much as 32.6% of the total number of links in the network, and that basically
Fig 6. Relation between the various codon bias indices of genes and the degree $k$ of the corresponding proteins in the PIN of E.coli. Solid lines are linear fits. CompAI and tAI of a gene definitely increase with the connectivity of the corresponding protein in the PIN, whereas the other two indices are less sensitive to this parameter.

doi:10.1371/journal.pone.0142127.g006

Table 1. Features of top-scoring communities. Number of nodes ($n$), community score ($n$ times the internal density), mean degree ($k$), predominant COG label and percentage; then, for ERI and the codon bias indices, mean values $x$, internal to the community and $Z$ scores (between square brackets). Values $Z > 1$ are reported in bold.

| ID | n  | score | $k$ | COG  | ERI  | CompAI  | CAI  | tAI  | Nc   |
|----|----|-------|-----|------|------|---------|------|------|------|
| 1  | 60 | 54.9  | 63.15 | J (90.0%) | 0.91 [1.66] | 0.13 [1.40] | 0.75 [0.05] | 0.38 [1.35] | 49.16 [-0.06] |
| 2  | 31 | 28.6  | 35.03 | N (74.2%)  | 0.38 [0.21] | 0.08 [-0.35] | 0.75 [0.07] | 0.32 [0.14] | 49.88 [0.12] |
| 3  | 21 | 19.1  | 25.85 | C (97.6%)  | 0.53 [0.65] | 0.09 [0.38] | 0.74 [-0.13] | 0.34 [0.72] | 50.18 [0.2] |
| 4  | 15 | 13.9  | 18.40 | M (66.7%)  | 0.82 [1.31] | 0.09 [0.07] | 0.75 [0.15] | 0.31 [0.07] | 49.32 [-0.02] |
| 5  | 13 | 11.7  | 10.77 | P (76.9%)  | 0.20 [-0.29] | 0.08 [-0.26] | 0.77 [0.40] | 0.33 [0.54] | 48.57 [-0.22] |
| 6  | 12 | 11.5  | 11.50 | U (48.9%)  | 0.20 [-0.29] | 0.07 [-0.82] | 0.76 [0.26] | 0.28 [-0.63] | 48.92 [-0.12] |
| 7  | 11 | 10.6  | 19.82 | P (63.6%)  | 0.56 [0.70] | 0.09 [0.44] | 0.76 [0.28] | 0.34 [0.72] | 48.74 [-0.14] |
| 8  | 10 | 10.0  | 11.60 | C (75.0%)  | 0.04 [-0.86] | 0.07 [-0.66] | 0.76 [0.28] | 0.29 [-0.45] | 47.78 [-0.30] |

doi:10.1371/journal.pone.0142127.t001
overlaps with the main core of the PIN (i.e., the k-core with the highest possible degree). Notably, proteins belonging to this community have on average a codon bias index (as measured by tAI and, even more, by CompAI) that is significantly higher than the average of the rest of the network (the Z-score is bigger than 1). As noticed above, this core is essentially composed of ribosomal proteins, that are usually highly expressed, have the highest codon usage bias, and are broadly conserved and essential across different taxa [38].

**Principal Component Analysis.** Finally we perform PCA over the space of the four codon bias indices (CompAI, CAI, tAI, Nc) measured for each E.coli gene. The two first principal components (PC1 and PC2) turn out to represent as much as 85% of the total variance of codon bias over the genome (left plot of Fig 7). Projection of the first two principal components on the individual codon bias indices (loadings) shows that none of the four indices predominantly contributes to the data variability (right plot of Fig 7). Thus, the placement of a gene in the PC1-PC2 plane depends on a weighted contribution of all the indices. Interestingly, the genes encoding for the proteins of the eight top MCODE communities are well localized and separated in this reduced space (Fig 8). In particular, the first community (i.e., the core of ribosomal proteins characterized by high values of both CompAI and tAI) is located in the upper left part of the graph, isolated from the others. This represents an important evidence: proteins that belong to the densest connected cores of the interactome are well-localized in the space of the
two principal components. In other words, if a set of proteins are physically and functionally connected in a module, then their corresponding genes should share common codon bias features. Conversely, we can obtain an estimate for the conditional probability $Pr(link|d)$ of a functional interaction between proteins, provided that their relative genes fall within a distance $d$ in the plane of the two principal components PC1 and PC2. Reasonably, we compare $Pr(link|d)$ estimated on the real interactome with $\langle Pr(link|d) \rangle_{CM}$ estimated on the Configuration Model (CM) which, we recall, is a degree-conserving randomization (re-wiring) of the network. Fig 9 shows the Z-score for $Pr(link|d)$ as a function of $d$, and reveals a peculiar behavior: for small distances ($d \leq 2$) the probability of finding a connection between two proteins is much higher than what could have been expected from a (degree-conserving) random link placement. Conversely, for medium distances ($3 \leq d \leq 9$), the linking probability is lower than that of the CM, whereas, the real network and the CM become compatible for large distances, where, however, connections are rather few. This analysis shows that sets of genes sharing similar codon usage
patterns encode for proteins that are much more likely to interact than in situations where chance alone is responsible for the structure of the interactome.

Conclusions

In this work we have introduced CompAI, a novel codon bias index that is inspired by tAI, though conceptually distinct. In fact, CompAI does not make reference to lists of highly expressed genes, and is thus unsupervised and based on intrinsic information about co-evolution of genes that code for proteins and tRNAs. Conceptually, the definition of CompAI is based on a model that postulates a competition between cognate and near-cognate tRNAs for the same codon, exposed on the ribosome at each step of protein synthesis. Competitive mechanisms in the machinery of ribosomal translation of genes into proteins have been repeatedly suggested and studied in the literature [39] and deserve further attention in order to understand their role for translation efficiency.

Our genome-wide analysis of codon bias in E.coli using CompAI as well as other commonly used indices revealed that codon usage metrics resting on counting tRNA genes (CompAI and tAI) are strongly and positively correlated among themselves—in spite of their conceptually different definition. It would then be quite interesting to check in the future whether this correlation is specific to E.coli or it is universally observed in the genomes of bacterial species that are either ecologically and evolutionarily close or, by contrast, very far from E.coli. We also found that both CompAI and tAI correlate with ERI, the degree of conservation for a gene among similar species, and gene essentiality, whereas, CAI and Nc are less sensitive to these
quantities. *CompAI* and *tAI* values also allow to distinguish three groups of genes, that are differently characterized by codon choice adaptation, ERI and degree of essentiality, and that also feature specific predominant COG signatures. In particular the third group (C), composed of the few genes that are highly conserved and with the strongest codon bias adaptation, consists for 30% of essential genes with predominant COGs J and L—that refer to translation, ribosome structure and biogenesis, replication, recombination and repair. These represent house-keeping and control functions that must be continuously executed by the cell, meaning that the genes responsible for them have to be expressed most of the time during the cell cycle. These observations strongly support the idea that an increasing selection of codons and, in parallel, a correlated modulation of tRNA availability co-evolved along the evolutionary history of a species.

Finally, we have addressed a theme as relevant as the connection between codon usage bias and protein functional or physical interactions. Our main result indicates that, in the course of the evolution of a genome, the functional structuring of the complex of interactions between proteins has interfered with the peculiar codon-coding formulation of the corresponding genes. In particular we have shown here, for the first time to our knowledge, that communities of highly connected proteins in the interactome of *E. coli* correspond to encoding genes that share the same degree of evolutionary adaptation, as expressed by codon bias indices that synthetically represent genetic information encoded in the tRNAs sector of the genome. Indeed, *CompAI*, that is based on a simple representation of tRNA competition, seems to detect the codon bias signal behind communities more consistently than the other indices here considered. Conversely, we have provided evidence that if two genes have similar codon usage patterns then the corresponding proteins have a significant probability of being functionally connected and interacting. This result points out that codon bias should be a relevant parameter in the fundamental problem of predicting unknown protein-protein interactions from genomic information. This study is a first exploratory step towards a more complete investigation on how communities within protein-protein interaction networks rest on a consistent but still to be decoded codon bias signal. Indeed, the connection of the topology of a network with an underneath semantics is far from trivial, as recently pointed out in the specialized literature [40]. Biological PINs and codon bias offer an interesting case study worth to be investigated in the wider perspective of multilayer network theory [41].

Finally we remark that *CompAI* was designed to provide information about the speed of protein synthesis, being based on proofreading delay mechanisms. In this respect, the wobble pairing between codon and anticodon reduces the rate of translation elongation with respect to WC base-pairing [42–44]. This slowing down of translation speed can be taken into account when devising a codon bias index that is intended specifically to measure the speed of protein synthesis. To this end, *CompAI* can be adapted by modifying the expression of the $W_i$ as:

\[
W_i = \left( \sum_{j=1}^{m} tGCN_{ij}^{WC} + \lambda \sum_{j=1}^{m} tGCN_{ij}^{wb} \right) \left[ \frac{\sum_{j=1}^{m} tGCN_{ij}}{\sum_{j=1}^{m} tGCN_{ij} + \sum_{j=1}^{m} tGCN_{ij}^{mc}} \right],
\]

where we have separated the cognate tGCN into WC and wobble (wb) base pairing. The wobble interaction factor $\lambda$ represents a penalty for such a coupling; we have set this value to 1/3, following the argument given in ref. [44]. In this modified version of *CompAI* (that we name *CompAI*$_W$), the first term in parenthesis is related to the different translational speed of codons, whereas, the second term in parenthesis measures the competition between cognate and near-
cognate tRNAs for binding to the ribosome. It is then interesting to investigate the correlation between the original CompAI and CompAI\textsubscript{w}. If such a correlation is computed separately for each of the gene groups A, B and C of E.Coli (that have been defined above, see Fig 2), we observe that, remarkably, CompAI and CompAI\textsubscript{w} are well correlated in group C genes, whereas, in group B and in particular in group A (the one including the lowest ERI gene) the two indices uncouple (see S3 Fig). We can conclude that the contribution due to wobble-pairing is particularly relevant among the set of genes that are quite specific of E.Coli (intermediate and low ERI genes). This observation suggests to further investigate the gene ontology of the set of genes for which the modulation of the translation speed due to wobbling pairing is relevant. As a general conclusion we are inclined to think that in bacterial species there are two groups of genes: i) the more universal, conserved and shared ones (in this paper, group C genes), whose evolvability [45] is small; and ii) specific genes, more evolvable and adaptable to the species’ novel environments. The modulation of the translation rate due to wobble pairing of isoacceptor tRNAs thus seems to be a typical mechanism of adaptable genes, a phenomenon that could be investigated in long-term evolutionary experiments.

Supporting Information

**S1 Fig. Abundance of tGCN cognate and near cognate for each anti-codon in E.coli.** Data taken from [17].

(TIFF)

**S2 Fig.** Left plot: Distribution of confidence levels \( Q(w) \), with the vertical line indicating the cut-off we use to separate true from false positives. Right plot: Distribution of degrees \( P(k) \) when \( \Theta = 0.9 \), with the insets showing the same distribution for the original network (\( \Theta = 0 \)).

(TIFF)

**S3 Fig. Correlation between CompAI and CompAI\textsubscript{w}.** Codon bias values are plotted separately for each of the E.Coli gene groups A, B and C. The black solid line identifies the linear regression over the whole genome (with Pearson’s correlation coefficient \( c = 0.58 \)). The blue solid lines instead represent linear regression on individual groups, with correlation coefficients \( c = 0.43 \) for group A, \( c = 0.62 \) for group B and \( c = 0.83 \) for group C.

(TIFF)

**S1 File.**

(PDF)

Acknowledgments

We thank the anonymous referee for useful suggestions.

Author Contributions

Conceived and designed the experiments: MD GC AS AG. Performed the experiments: MD GC AS AD. Analyzed the data: MD GC AD. Contributed reagents/materials/analysis tools: MD GC AD AS. Wrote the paper: MD GC AG.

References

1. Hershberg R, Petrov DA. Selection on codon bias. Ann Rev Genet. 2008; 42:287–99. doi:10.1146/amnurev.genet.42.110807.091442 PMID: 18983258

2. Shabalina SA, Spiridonov SA, Kashina A. Sounds of silence: synonymous nucleotides as a key to biological regulation and complexity. Nucleic Acids Res. 2013; 41(4):2073–2094. doi:10.1093/nar/gks1205 PMID: 23293005
16. Benson DA, Karsch-Mizrachi I, Clark K, Lipman DJ, Ostell J, Sayers EW. GenBank. Nucleic Acids Res. 1999; 27(1):20–22. PMID: 9950598

17. Eyre-Walker A. Synonymous codon bias is related to gene length in Escherichia coli: selection for translational accuracy? Mol Biol Evol. 1996; 13(6):864–872. PMID: 8754221

18. Alexander Roth MA, Cannarozzi GM. Measuring codon usage bias. In: Cannarozzi GM, Schneider A, editors. Codon Evolution Mechanisms and Models. Oxford University Press; 2012. p. 189–217.

19. Sharp PM, Li WH. The codon Adaptation Index—a measure of directional synonymous codon usage bias, and its potential applications. Nucleic Acids Res. 1987; 15(3):1281–1295. doi: 10.1093/nar/15.3.1281 PMID: 3547335

20. Li GW, Burkhardt D, Gross C, Weissmann JS. Quantifying absolute protein synthesis rates reveals principles underlying allocation of cellular resources. Cell 2014; 157(3):624–635. doi: 10.1016/j.cell.2014.02.033 PMID: 24766808

21. Pop C, Rouskin S, Ingolia NT, Han L, Phizicky EM, Weissman JS. Causal signals between codon bias, mRNA structure, and the efficiency of translation and elongation. Mol Syst Biol. 2014; 10:770. doi: 10.15252/msb.20145524 PMID: 25538139

22. Daubin V, Perrière G. G+C structuring along the genome: a common feature in prokaryotes. Mol Biol Evol. 2003; 20(4):471–483. doi: 10.1093/molbev/msg022 PMID: 12654929

23. Pop C, Rouskin S, Ingolia NT, Han L, Phizicky EM, Weissman JS. Causal signals between codon bias, mRNA structure, and the efficiency of translation and elongation. Mol Syst Biol. 2014; 10:770. doi: 10.15252/msb.20145524 PMID: 25538139

24. Benson DA, Karsch-Mizrachi I, Clark K, Lipman DJ, Ostell J, Sayers EW. GenBank. Nucleic Acids Res. 1999; 27(1):20–22. PMID: 9950598

25. Kanaya S, Yamada Y, Kudo Y, Ikemura T. Studies of codon usage and tRNA genes of 18 unicellular organisms and quantification of Bacillus subtilis tRNAs: gene expression level and species-specific
diversity of codon usage based on multivariate analysis. Gene. 1999; 238(1):143–155. doi: 10.1016/S0378-1119(99)00225-5 PMID: 10570992

26. Zhang G, Ignatova Z. Generic algorithm to predict the speed of translational elongation: implications for protein biogenesis. PLoS One. 2009; 4(4):e5036. doi: 10.1371/journal.pone.0005036 PMID: 19343777

27. Chien CT, Bartel PL, Sternglanz R, Fields S. The two-hybrid system: a method to identify and clone genes for proteins that interact with a protein of interest. Proc Natl Acad Sci USA. 1991; 88(21):9578–9582. doi: 10.1073/pnas.88.21.9578 PMID: 1946372

28. Phizicky EM, Fields S. Protein-protein interactions: methods for detection and analysis. Microbiol Rev. 1995; 59(1):94–123. PMID: 7708014

29. Puig O, Caspary F, Rigaut G, Rutz B, Bouveret E, Bragado-Nilsson E, et al. The tandem affinity purification (TAP) method: a general procedure of protein complex purification. Methods. 2001; 24(3):218–229. doi: 10.1006/meth.2001.1183 PMID: 11403571

30. Kanehisa M, Goto S. KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acids Res. 2000; 28(1):27–30. doi: 10.1093/nar/28.1.27 PMID: 10592173

31. Huang H, Jedynak BM, Bader JS. Where have all the interactions gone? Estimating the coverage of two-hybrid protein interaction maps. PLoS Comput Biol. 2007; 3(11):e214. doi: 10.1371/journal.pcbi.0030214 PMID: 18039026

32. Bader GD, Hogue CWV. An automated method for finding molecular complexes in large protein interaction networks. BMC Bioinformatics. 2003; 4:2. doi: 10.1186/1471-2105-4-2 PMID: 12525261

33. Newman MEJ. Modularity and community structure in networks. Proc Natl Acad Sci USA. 2006; 103(23):8577–8582. doi: 10.1073/pnas.0601602103 PMID: 16723398

34. Tatusov RL, Natale DA, Garkavtsev IV, Tatusova TA, Shankavaram UT, Rao BS, et al. The COG database: new developments in phylogenetic classification of proteins from complete genomes. Nucleic Acids Res. 2001; 29(1):22–38. doi: 10.1093/nar/29.1.22 PMID: 11125040

35. Jolliffe IT. Principal Component Analysis. Series: Springer Series in Statistics. Springer; 2002. 487: p. 28.

36. Park J, Newman MEJ. Statistical mechanics of networks. Phys Rev E 2004; 70(6):066117. doi: 10.1103/PhysRevE.70.066117

37. Squartini T, Garlaschelli D. Analytical maximum-likelihood method to detect patterns in real networks. New J Phys. 2011; 13:083001. doi: 10.1088/1367-2630/13/8/083001

38. Butland G, Peregrín-Alvarez JM, Li J, Yang W, Yang X, Canadien V, et al. Interaction network containing conserved and essential protein complexes in Escherichia coli. Nature. 2005; 433(7025):531–537. doi: 10.1038/nature03239 PMID: 15690043

39. Fluit A, Pienaar E, Viljoen H. Ribosome kinetics and aa-tRNA competition determine rate and fidelity of peptide synthesis. Comput Biol Chem. 2007; 31(5–6):335–346. doi: 10.1016/j.compbiolchem.2007.07.003 PMID: 17897886

40. Hric D, Darst RK, Fortunato S. Community detection in networks: Structural communities versus ground truth. Phys Rev E. 2014; 90(6):062805. doi: 10.1103/PhysRevE.90.062805

41. Boccaletti S, Bianconi G, Criado R, del Genio CI, Gómez-Gardeñes J, Romance M, et al. The structure and dynamics of multilayer networks. Physics Reports. 2014; 544(1):1–122. doi: 10.1016/j.physrep.2014.07.001

42. Sorensen MA. Absolute in vivo translation rates of individual codons in Escherichia coli: the two glutamic acid codons GAA and GAG are translated with a threefold difference in rate. J Mol Biol. 1991; 222:265–280. doi: 10.1016/0022-2836(91)90211-N PMID: 1960727

43. Stadler M, Fire A. Wobble base-pairing slows in vivo translation elongation in metazoans. RNA. 2011; 17(12):2063–2073. doi: 10.1261/rna.02890211 PMID: 22045228

44. Spencer PS, Siller E, Anderson JF, Barral JM. Silent substitutions predictably alter translation elongation rates and protein folding efficiencies. J Mol Biol. 2012; 422:328–335. doi: 10.1016/j.jmb.2012.06.010 PMID: 22705285

45. Pigliucci M. Is evolvability evolvable? Nature Reviews Genetics. 2008; 9:75–82. doi: 10.1038/nrg2278 PMID: 18059367