Decoding of all codons can be achieved by a subset of tRNAs. In bacteria, certain tRNA species are mandatory, while others are auxiliary and are variably used. It is currently unknown how this variability has evolved and whether it provides an adaptive advantage. Here we shed light on the subset of auxiliary tRNAs, using genomic data from 319 bacteria. By reconstructing the evolution of tRNAs we show that the auxiliary tRNAs are highly dynamic, being frequently gained and lost along the phylogenetic tree, with a clear dominance of loss events for most auxiliary tRNA species. We reveal distinct co-gain and co-loss patterns for subsets of the auxiliary tRNAs, suggesting that they are subjected to the same selection forces. Controlling for phylogenetic dependencies, we find that the usage of these tRNA species is positively correlated with GC content and may derive directly from nucleotide bias or from preference of Watson–Crick codon–anticodon interactions. Our results highlight the highly dynamic nature of these tRNAs and their complicated balance with codon usage.

INTRODUCTION

tRNA is a key molecule in all cells due to its central role in protein translation. Each tRNA is an adaptor molecule that can be charged with an amino acid residue and donate it to an elongating peptide chain based on codon–anticodon recognition. Each tRNA species, distinguished by its anticodon sequence, recognizes a specific set of codons, which encode the amino acid it is loaded with. This specificity reflects the genetic code and dictates the translation of a nucleotide sequence into protein. Codon–tRNA interaction exhibits two types of redundancy, whereby the same codon can be decoded by different tRNA species (having different anticodons, but loaded with the same amino acid), and the same tRNA species can decode different codons of the same amino acid. This is achieved by wobble interactions (non-Watson–Crick base pairing) between the third codon position and the first anticodon position. Such wobble interactions can occur due to Guanine-Uracil (G:U) base pairing, as well as many anticodon modifications that change codon specificity (1). Due to this redundancy, not all possible tRNA species are required in order to decode the 61 sense codons, and various organisms employ different subsets of tRNAs (2).

The variability of the tRNA repertoire has been mostly explored in relation to tRNA abundance, reflected by gene copy number. It has been suggested that selection for efficient translation leads to close correspondence between codon usage and the tRNA pool (3). This led to the development of the tRNA adaptation index (tAI) (4), which was used as a measure of translational selection (5–8). In addition, translational selection was found to be associated with the increased number of tRNA genes (9,10). A binary view of an organism tRNA repertoire is required, namely whether a tRNA species exists or not, has been less investigated. In the work mentioned above, Rocha (9) found that translational selection is associated with the decreased number of tRNA species. A survey of the tRNA repertoire in representative organisms from all kingdoms of life led to the formulation of three common strategies employed to achieve reduction in the number of tRNA species, where by each strategy some of the tRNAs were spared (2). Novoa et al. (11) showed that archaea employ the smallest number of tRNA species, bacteria prefer the use of U34N35N36 tRNAs (subclass represents position numbering convention where anticodon positions 34, 35, and 36 base pair with codon positions 3, 2, and 1, respectively), in correspondence with the presence of the modifying enzyme tRNA-dependent uridine methyltransferase, and eukaryotes prefer the use of A34N35N36 tRNAs, in correspondence with the presence of the modifying enzyme tRNA-dependent adenosine deaminase. These studies mainly involved the common patterns of tRNA repertoire at the kingdom level. However, little attention has been addressed to the variations in tRNA repertoire within a kingdom, and it is currently unknown whether these variations are shaped to provide an adaptive advantage.

Here we used a dataset of 319 fully sequenced genus representative bacteria to find patterns of tRNA usage and or...
ganism traits associated with them. We found that while certain tRNA species are always used, there is a subset of tRNA species that differs substantially between bacteria. Based on the striking absence of \( A_{34}N_{35}N_{36} \) tRNAs and the extended set of wobble rules, this pattern can be explained by the need to decode all sense codons, thus separating the tRNA species of an organism into mandatory and auxiliary tRNAs. The latter are especially intriguing to investigate, as while an organism can theoretically do without them, different organisms show different repertoires of these tRNAs, and it is not clear what underlies this variability and how it evolved. We use bioinformatic methods employing phylogenetic information to explore the changes in the tRNA repertoire during evolution, with a special emphasis on the auxiliary tRNAs, which were found to be highly dynamic, frequently gained and lost along the phylogenetic tree. Through analysis of sequence similarity, we found evidence pointing at multiple pathways of tRNA species gain through horizontal gene transfer (HGT), anticodon mutations, and a combination of both. Finally, the variability in the use of auxiliary tRNAs was found to be mostly connected to the nucleotide content of the genome, but also to some extent to the strength of translational selection. We discuss possible explanations for the observed associations between the tRNA repertoire and both the nucleotide content and translational selection.

**MATERIALS AND METHODS**

**Genome data and organism datasets**

The genomic sequence and genome-related data of 1245 bacteria was retrieved from the NCBI FTP site (ftp://ftp.ncbi.nih.gov/genomes/Bacteria, April 2011). To reduce bias from closely related organisms, one bacterium species per genus was randomly selected as representative. We excluded endosymbiotic bacteria with severely reduced genomes (less than \( 10^6 \) bp) and less than 31 tRNA species. This resulted in a dataset of 319 bacteria.

Species tree in Newick format was downloaded from the MicrobesOnline database (12) and was trimmed to the chosen dataset of 319 bacteria. This tree is based on sequence alignment of 78 sets of protein orthologs, encoded by a single copy in most bacteria. While HGT events are common in prokaryotes, this core of conserved proteins represents a central trend of vertical inheritance (13) and can be used to construct a reliable species tree (14).

**tRNA repertoire**

Sequences of all non-coding RNAs identified in 319 genus representative bacteria (as annotated by RefSeq) were scanned using the program tRNAscan-SE with bacteria-specific parameters (15). This comprised the repertoire of identified tRNAs and their respective anticodons.

**Association with organism traits**

Three genomic traits were considered: genome size, GC content and ENC\(^\text{C}^{\text{diff}}\). Genome size and GC content were calculated from the genomic sequence. ENC\(^{\text{C}^{\text{diff}}}\) is an estimate of translational selection at an organism scale calculated as the difference between the average ENC\(^{\text{C}}\) of all genes and the genes encoding ribosomal proteins, normalized by the average ENC\(^{\text{C}}\) of all genes. ENC\(^{\text{C}}\) is a variant of the effective number of codons (ENC) index (16) that accounts for background nucleotide composition (17). ENC\(^{\text{C}}\) takes the value of 61 when all codons are used at the frequency expected given the nucleotide composition, and its value decreases as codon usage deviates from the expected. Gene ENC\(^{\text{C}}\) was calculated based on the background nucleotide composition of the same gene. The values of the three genomic traits were normalized so that their mean is zero and their standard deviation is one.

We performed regression analysis between the presence/absence profile of each tRNA species and the genomic traits, which takes into account the phylogenetic relationships between the organisms. This was done using the maximum likelihood regression analysis option of the BayesTraitsV2.0 package (18,19) (http://www.evolution.rdg.ac.uk/BayesTraits.html). While designed for purely continuous data, the algorithm can also be used when the dependent trait is binary. For each presence/absence profile of a tRNA species, the algorithm was used to build regression models with each of the three genomic traits and for each pair of genomic traits, a total of six runs. In addition, the likelihood of the data given only the phylogenetic tree was evaluated by calculating the maximum likelihood when the regression coefficient with each of the genomic traits was set to zero. The parameter \( \lambda \), which effectively scales the branch lengths of the phylogenetic tree, was estimated in each run in order to allow variation in the strength of the phylogenetic signal. The likelihood calculated when the regression coefficient was set to zero was compared to the likelihood of the regression with genomic trait X to estimate the statistical significance of the contribution of trait X. The likelihood of the regression with genomic trait Y was compared to the likelihood of the regression with both genomic trait X and genomic trait Y to estimate the statistical significance of the contribution of trait X over trait Y. The statistic used is twice the difference between the maximum likelihood values of the compared models. This statistic is distributed as \( \chi^2 \) with the number of degrees of freedom that is equal to the difference in the number of parameters between the two models, in this case—one. The association of tRNA presence with a genomic trait was considered statistically significant and independent of the other two genomic traits if adding it to the regression of each of the other traits provided statistically significant results. Since the same data was repeatedly analyzed, false discovery rate procedure was used to control for multiple comparisons based on all tests performed (\( p \)-value threshold is \( 8.5 \times 10^{-3} \)).

To test whether the statistically insignificant association of some auxiliary tRNAs with GC content stems from the low level of variation in their presence/absence profile, for each of these tRNAs we examined the percentage of genomes that possess it or lack it, and chose the highest of those as a measure of variability. We then compared this measure between the set of auxiliary tRNAs that were statistically significantly associated with GC content and all other auxiliary tRNAs by a Mann–Whitney test.
Detecting patterns in tRNA repertoire

A binary matrix representing the presence/absence of each tRNA species in 319 bacteria was constructed. The matrix was clustered based on Hamming distance between the presence/absence profiles of each two tRNA species.

The GC content in which an auxiliary tRNA species tends to appear (usage shift) was defined as follows. Bacteria were divided into two groups based on the presence/absence of the tRNA, and each group was ordered by GC content. The GC content where the usage shift occurs was determined as the average between the GC content of the 75 percentile of bacteria in the ‘absent’ group and the GC content of the 25 percentile of bacteria in the ‘present’ group. This procedure was applied only to tRNA species that showed a statistically significant difference in GC content distribution between the ‘absent’ and ‘present’ groups using a Mann–Whitney test.

Ancestral reconstruction

The GLOOME algorithm (20–22) was used to reconstruct the tRNA repertoire in each internal node of the phylogenetic tree as well as to estimate rates of tRNA gain and loss. The algorithm was applied to 52 tRNA species in all 319 organisms simultaneously (tRNA species never observed in the studied organisms were excluded). The evolutionary model used was ‘variable gain/loss ratio (mixture)’ and gamma rate distribution. Both of these parameters were chosen to allow tRNA species to vary as much as possible in their evolutionary model. We also used high optimization level. The reconstruction assigns a probability for the presence of a tRNA species in each internal node and produces an estimation of gain/loss probability per branch. Nodes were considered as undergoing gain or loss events if the branch leading to them had a gain or loss probability of at least 0.8 (see more details in Supplementary Methods).

Origin of new tRNA species

Clues as to the origin of a tRNA gene are more likely to exist in recently acquired genes. Most recent gain events (MRGEs) were defined as gain events that have no other gain of the same tRNA species below them in the phylogenetic tree. For a MRGE of tRNA-X, all the genes descending from the gain were compared against a database of tRNA genes from all tRNA species and organisms using BLAST. The BLAST score of a tRNA against itself was used to evaluate the quality of the match. Only hits that scored at least 80% of the self-hit score were further considered. Hits were classified as ‘vertical inheritance’ if they had the same anticodon and descended from the same MRGE as the query, or otherwise as ‘non-vertical inheritance’. For each query, the best-scored hits were considered in the vertical inheritance category and in the non-vertical inheritance category. The results of the ancestral reconstruction, MRGE analysis and the mapping of potential tRNA origins were uploaded to the interactive tree of life (iTOL) (23,24) for visualization in tree context.

tRNA co-gain and co-loss and operon structure

Operon predictions were retrieved from the MicrobesOnline database (25). Organisms originating from multiple gain events were suspected to acquire the tRNA genes through horizontal operon transfer. For each two tRNA species that had at least one event of co-gain, the number of organisms descending from these events that contain an operon with genes of both tRNA species was recorded. The same was done for organisms not descending from a co-gain event to provide a background of operon co-existence of the two tRNA species.

Nodes where multiple loss events were predicted were suspected to have lost the tRNA genes through operon deletion. We looked for evidence for gene proximity in organisms evolutionarily closest to the co-loss event. Organisms were considered nearest neighbors if they descended from the parent node of the co-loss event but not from the co-loss event itself. For each two tRNA species that had at least one event of co-loss, the number of nearest neighbors that contain an operon with genes of both tRNA species was recorded. The same was done for organisms not considered nearest neighbors to provide a background of operon co-existence of the two tRNA species.

RESULTS

Patterns of tRNA usage across species

We scanned 319 bacterial genomes to identify their tRNA genes (annotated as tRNA species by the anticodon sequence) (Figure 1A). Pseudo tRNAs were excluded. As has been previously reported (2,11) most A34N35N36 tRNA species (except for A34C35G36 tRNA encoding Arginine) were found to be missing from all or the vast majority of organisms. It is known that Adenosine in the anticodon wobble position (position 34) is very efficiently modified to Inosine (I), which recognizes codons ending with C, U and A. This broad specificity has detrimental effects when a codon quartet is split into two pairs of codons (purine and pyrimidine-ending) encoding different amino acids, as it leads to ambiguities when decoding the A-ending codon. However, when all codons in the quartet encode the same amino acid, this broad specificity is not obviously harmful, as can be seen by the use of such tRNAs in eukaryotes (2). Therefore, while it is not clear yet what underlies the widespread avoidance of A34N35N36 tRNA species in prokaryotes, it seems that there is strong negative selection against them.

The absence of most A34N35N36 tRNAs can explain several observed patterns of tRNA usage. Specifically, U34N35N36 and G34N35N36 tRNA species are present in all, or the vast majority of organisms, since without A34N35N36 tRNAs, both are necessary for decoding the N1N2G3 and N1N2[CU]3 codons, respectively (see the allowed codon–anticodon interactions in Figure 1B). We therefore refer to these tRNA species as mandatory tRNAs. Two other mandatory tRNAs are C34A35U36 and C34C35A36, as they are the sole decoders of the Met and Trp codons. Since the always-present U34N35N36 tRNAs are capable of decoding N1N2G3 codons (Figure 1B), the use of C34N35N36 tRNAs is auxiliary rather than mandatory. Indeed, we found
The tRNA repertoire is highly variable. (A) The number of organisms (out of 319) that have at least one copy of a tRNA species. The numbers above the bars represent the number of codons in the relevant codon quartet that code for the amino acid decoded by the tRNA. (B) A schematic representation of known codon–anticodon interactions. Black solid arrows represent commonly occurring interactions. Gray arrows represent possible interactions when Adenosine is modified to Inosine in the first anticodon position. Black dashed arrows represent modifications-dependent interactions when the first anticodon position contains Uridine. The thickness of the dashed arrows represents interaction efficiency.

that the rest of the C34N35N36 tRNA species, which decode amino acids with several synonymous codons, show usage variability. Notice that the eight G34N35N36 tRNAs decoding 4-fold degenerate amino acids are probably not as essential as those decoding 2- and 3-fold degenerate amino acids since they can be supplemented or even replaced by the corresponding U34N35N36 tRNA under certain conditions (as discussed below, Figure 1B). Indeed these G34N35N36 tRNAs demonstrate greater usage variability and are therefore considered auxiliary tRNAs as well. There are two major exceptions to the pattern described above. Ile tRNA U34A35U36 is not mandatory but rather rarely seen, since in order to prevent misreading of the Met codon A1U2G3 as Ile, a specially modified tRNA (not considered in this work) exists instead of the conventional U34A35U36 (2). Also, the ArgN34C35G36 tRNA species shows different behavior due to the use of A34C35G36 tRNA by most bacteria. In the presence of A34C35G36 (modified to I34C35G36), G34C35G36 tRNAs becomes hardly used and auxiliary (Figure 1A and B). Since I34C35G36 does not decode the C1G2G3 codon, the presence of either U34C35G36 or C34C35G36 is mandatory, but they can be viewed as auxiliary to one another.

While the apparent selection against A34N35N36 tRNAs combined with codon–anticodon binding possibilities determines the mandatory and auxiliary tRNAs, an intriguing question regards the pattern of usage variability of the auxiliary tRNAs in the various genomes and the principles underlying it. To try and answer this question, we looked for associations between the use of each auxiliary tRNA species and several genomic traits. The genomic traits selected represent three aspects hypothesized to influence the tRNA repertoire: genome size, nucleotide content and translational selection. Genome size was shown to be correlated with tRNA gene number (4) and may therefore correlate with tRNA presence as well. The nucleotide content of the genome has a tremendous effect on codon usage, mainly through the third codon position (26,27). Co-evolution of codon usage and the tRNA pool may therefore be reflected as association between the nucleotide content measured by the GC content of the organism and the tRNA pool. The third genome property is the strength of selection for efficient translation, shown by Rocha to correlate with the total number of tRNA genes and to inversely correlate with the total number of tRNA species (9). We used ENC_diff as a measure of translational selection, calculated similarly to the measure proposed by Rocha in his work (see detailed description in Materials and Methods and in (28)). Briefly, ENC_diff is the normalized difference between the average ENC of all genes and the average ENC of ribosomal genes, where ENC is a measure of gene codon bias. The stronger the selection towards efficient translation in highly expressed genes, the higher the codon bias in the highly ex-
pressed ribosomal genes compared to all other genes, resulting in higher ENC\textsubscript{diff}.

We evaluated the association between tRNA repertoire and these traits by a regression model. The generally hierarchical evolution of species implies that the phylogenetic signal should influence to a certain extent the observed phenotypes (29). In other words, closely related species are expected to have more similar characteristics than remotely related organisms simply due to differences in divergence time, which may lead to biased estimates of association between genomic traits (29). It is therefore essential, when looking for associations between organism traits, to take into consideration the phylogenetic relatedness between genomes. To this end, we performed regression analysis using a phylogenetic generalized least-square approach implemented in the BayesTraits package (18,19). As suggested by Ives and Garland (30), the values of the continuous genomic traits were normalized so that the calculated regression coefficients will represent effect sizes. Testing the correlation between each two continuous genomic traits while taking into account the phylogenetic tree (downloaded from the MicrobesOnline database (12)) (Supplementary Methods) revealed a statistically significant correlation between genome size and GC content \((R = 0.305, p\text{-value} = 2.2 \times 10^{-5})\) and very weak correlations between ENC\textsubscript{diff} and both genome size and GC content \((R = 0.118, p\text{-value} = 3.8 \times 10^{-2} \) and \(R = -0.176, p\text{-value} = 1.6 \times 10^{-3}\), respectively).

For each tRNA species, the relationship between its binary presence/absence profile and the normalized values of each of the genomic traits was analyzed, taking into account the phylogenetic dependencies. We allowed the algorithm to simultaneously estimate the parameter \(\lambda\), which is a measure of the phylogenetic signal. \(\lambda\) values can range between 0 (no phylogenetic signal) and 1 (phylogeny perfectly explains variation). The values calculated for this parameter were often close to one (Supplementary Table S1), indicating a strong phylogenetic signal. It is therefore remarkable that the variation remaining after controlling for the strong phylogenetic signal is often found to be statistically significantly associated with additional genomic traits, as detailed below. Since the continuous genomic traits are not independent, we considered an association of the dependent variable (tRNA repertoire) with trait X to be independent of trait Y if the likelihood of the regression model with both X and Y traits was statistically significantly higher than the likelihood of the regression model with the Y trait alone (Materials and Methods). As can be seen in Figure 2, genome size is not independent associated with the presence/absence status of any tRNA species except for the rarely used U\textsubscript{34}A\textsubscript{35}U\textsubscript{36}. GC content, on the other hand, is positively (and independently from the genome size and ENC\textsubscript{diff} traits) associated with most of the auxiliary tRNAs, explaining between 3 and 34\% of the usage variability as measured by \(R^2\). Of note, the auxiliary tRNA species that were not statistically significantly associated with GC content were more uniformly present or absent in bacteria (Mann–Whitney test, \(p\text{-value} = 1.3 \times 10^{-3}\), see Materials and Methods). Almost all the auxiliary tRNAs exhibit a negative slope that reflects a negative correlation with ENC\textsubscript{diff}, although a statistically significant and independent negative association with ENC\textsubscript{diff} was observed for only three auxiliary tRNA species (C\textsubscript{34}G\textsubscript{35}A\textsubscript{36}Ser, C\textsubscript{34}G\textsubscript{35}G\textsubscript{36}Pro, C\textsubscript{34}G\textsubscript{35}U\textsubscript{36}Thr). This finding is in agreement with Rocha’s discovery that the total number of tRNA species decreases as translational selection increases (9).

We next turned to examine whether the auxiliary tRNAs usage evolved randomly or in a certain determined order. To address this, we presented the data as a matrix of tRNA species over organisms, where each matrix cell contains information about the presence/absence of tRNA X in organism Y. We clustered this matrix on the tRNA dimension using Hamming distance, which measures the number of differences between the profiles of each pair of tRNA species (Figure 3A). Organisms were arranged by their GC content (Figure 3B) as this property was found to be dominant in its relation to the tRNA profile. As can be seen in Figure 3A, there appears to be an order of auxiliary tRNA gain or loss. G\textsubscript{34}N\textsubscript{35}N\textsubscript{36} and C\textsubscript{34}N\textsubscript{35}N\textsubscript{36} tRNAs form separate clusters, with the exception of C\textsubscript{34}A\textsubscript{35}A\textsubscript{36}Leu and C\textsubscript{34}C\textsubscript{35}U\textsubscript{36}Arg, which are included in the G\textsubscript{34}N\textsubscript{35}N\textsubscript{36} cluster. In general, the usage shift (Materials and Methods) of G\textsubscript{34}N\textsubscript{35}N\textsubscript{36} tRNAs occurs in consistently lower GC-content values than that of their C\textsubscript{34}N\textsubscript{35}N\textsubscript{36} counterparts (Wilcoxon signed rank test \(p\text{-value} = 0.0156\)).

The codon quartet of Arg (tRNA N\textsubscript{34}C\textsubscript{35}G\textsubscript{36}) is an interesting special case as it is the only case where the A\textsubscript{34}N\textsubscript{35}N\textsubscript{36} tRNA is regularly used (Figure 3C and D). Since A\textsubscript{34}C\textsubscript{35}G\textsubscript{36} modified to I\textsubscript{34}C\textsubscript{35}G\textsubscript{36} does not decode the codon C\textsubscript{1}G\textsubscript{2}G\textsubscript{3} (see Figure 1B), an additional tRNA is required. In most bacteria (239) this is achieved by C\textsubscript{34}C\textsubscript{35}G\textsubscript{36}. However in 24 bacteria the additional tRNA is U\textsubscript{34}C\textsubscript{35}G\textsubscript{36}, and in 26 bacteria both tRNAs are employed. Of note, C\textsubscript{34}C\textsubscript{35}G\textsubscript{36} tRNA is used when GC content is high and U\textsubscript{34}C\textsubscript{35}G\textsubscript{36} when it is low. Furthermore, the rare event of supplementing the A\textsubscript{34}C\textsubscript{35}G\textsubscript{36} tRNA with G\textsubscript{34}C\textsubscript{35}G\textsubscript{36} tRNA occurs only when GC content is high. In the infrequent absence of A\textsubscript{34}C\textsubscript{35}G\textsubscript{36} tRNA, both G\textsubscript{34}C\textsubscript{35}G\textsubscript{36} and U\textsubscript{34}C\textsubscript{35}G\textsubscript{36} are required to decode the quartet. In nine out of 19 cases, the C\textsubscript{34}C\textsubscript{35}G\textsubscript{36} tRNA is also found but only when GC content is high. It would appear that even in this more complicated case of tRNA combinatorics, the rule governing the use of auxiliary tRNAs is that G\textsubscript{34}N\textsubscript{35}N\textsubscript{36} and C\textsubscript{34}N\textsubscript{35}N\textsubscript{36} are enlisted when GC content is high and the non-essential U\textsubscript{34}N\textsubscript{35}N\textsubscript{36} is enlisted when GC content is low.

tRNA dynamics is dominated by gene loss

To better understand the dynamic processes that shape tRNA usage, an evolutionary reconstruction is essential. We used the GLOOME software (20–22) to combine tRNA presence/absence profile with the phylogenetic tree and thus reconstructed the probable evolutionary history of each tRNA species. GLOOME implements a stochastic mapping approach to assign gain and loss events onto each branch of a phylogenetic tree based on the topology and branch lengths of the tree. The algorithm was used to infer branch-specific and tRNA-specific gain and loss events using an evolutionary model that permits the gain/loss ratio to vary among tRNAs, thereby allowing for different evolutionary behaviors for the various tRNAs. While the expectation of gain/loss per branch depends on its length, the overall
Figure 2. Auxiliary tRNA presence/absence profile is mostly associated with GC content. Regression analysis of tRNA presence/absence profile with genome size (top), GC content (middle) and ENC’ diff (bottom). RNA species that exhibited little presence/absence variability (when three or less organisms had a different presence/absence status than the majority of organisms) are not presented. In parentheses are the number of codons in the relevant codon quartet that code for the amino acid decoded by the tRNA and the number of organisms in which the tRNA species was found (out of 319). $R^2$ is the amount of variation explained by the regression model (note the different $R^2$ range). Asterisks denote statistically independent genomic traits that when added to regression between the dependent variable (tRNA presence/absence) and each of the additional genomic traits, statistically significantly improved the fit of the data to the calculated regression model.

tRNA gain/loss rates are the sum of gain/loss expectations predicted per branch over all the branches of the tree and are therefore comparable (Figure 4). As expected, mandatory tRNAs have very low loss and gain rates. A notable exception is U$_{34}$A$_{35}$A$_{36}$-Leu where a scenario of multiple loss events that occurred late in evolution and a scenario of few early losses followed by multiple gain events both explain the tRNA pattern and are therefore summed together to produce high gain and loss rates. Compared to mandatory tRNAs, auxiliary tRNAs tend to have much higher gain and loss rates. While the rates are highly varied, the dominant trend is of tRNA loss. Only four auxiliary tRNAs are gained at a higher rate than they are lost. These include the rarely used Arg tRNAs G$_{34}$C$_{35}$G$_{36}$ and U$_{34}$C$_{35}$G$_{36}$ that are gained as a replacement when the commonly used A$_{34}$C$_{35}$G$_{36}$ and C$_{34}$C$_{35}$G$_{36}$ are lost, and the highly dynamic C$_{34}$U$_{35}$C$_{36}$-Glu and C$_{34}$U$_{35}$G$_{36}$-Gln. GLOOME calculates a probabilistic reconstruction of ancestral states (see example in Supplementary Figure S1). While there is a distinct signal of vertical inheritance demonstrated by the clusters of related bacteria, all either possessing or missing a specific tRNA, it is also obvious that multiple gain and loss events have occurred repeatedly and independently throughout evolution for most auxiliary tRNAs. The ancestral reconstruction of all auxiliary tRNAs suggests with high probability that they were present at the bacterial ancestor (Supplementary Methods and Supplementary Table S2), as occurs often in extant GC-rich bacteria. This may point to a GC-rich ancestor or to an underlying environmental influence in which the extensive tRNA set promoted survival. Furthermore, this implies that tRNA species with low gain rate like C$_{34}$A$_{35}$C$_{36}$-Val and C$_{34}$G$_{35}$C$_{36}$-Ala once lost are not gained again, whereas for tRNA species with higher gain rate, the loss is not as final.

Identification of gain and loss events makes it also possible to assess the dependencies between the different tRNA species. We used random simulations (Supplementary Materials and Methods) to generate a background distribution of independently gained and lost tRNA species, to which the actual distribution of gain and loss events was comp-
Figure 3. GC content effect on auxiliary tRNA species is variable. (A) Clustering of the presence/absence (black/white, respectively) profiles of auxiliary tRNA species in 319 bacteria. tRNA species are clustered based on Hamming distance while bacteria are arranged by their GC content presented in (B). The GC content where a tRNA species tends to appear/disappear (usage shift) was defined as the mean between the GC content of the 75 percentile of organisms missing the tRNA and the 25 percentile of organisms that possess the tRNA. Asterisks mark the usage shift location based on B. In parentheses are the number of codons in the relevant codon quartet that code for the amino acid decoded by the tRNA, and the usage shift value, separated by a comma.

(B) Distribution of Arg tRNA species combinations among the genomes. This revealed that the observed distribution is statistically significantly biased towards nodes originating from multiple gain or loss events (Supplementary Figure S2), pointing to strong dependencies between tRNA species. Analysis of the individual dependencies between each pair of tRNA species (Supplementary Figure S3) revealed that C34U35C36-Glu and C34U35G36-Gln are co-gained in 10 nodes out of a maximum of 14 (such a result or higher was never reproduced in $10^4$ random simulations, Supplementary Methods). C34C35C36-Gly, C34G35A36-Ser, C34G35G36-Pro and C34G35U36-Thr also tended to be co-gained with each other more than expected at random. Statistically significant co-loss is found in almost all C34N35N36 auxiliary
Figure 4. Auxiliary tRNA evolution is highly dynamic and dominated by gene loss. Sum of gain (blue) and loss (red) branch probabilities over the phylogenetic tree calculated by GLOOME. tRNA species are listed alphabetically by their anticodon sequence. Light gray dots mark auxiliary tRNA species. Dark grey dots mark Arg auxiliary tRNA species.

tRNAs and to a lesser extent in G34N35N36 auxiliary tRNAs (mainly in G34G35C36-Ala and G34G35G36-Pro).

A detailed examination of sequential gain and loss events (Supplementary Methods and Supplementary Figure S4) revealed that there are also certain tRNA species that appeared in a specific order whenever they are proximal in the phylogenetic tree. Remarkably, these tRNA species (i.e. C34G35G36-Pro, C34G35U36-Thr, C34C35C36-Gly, C34G35A36-Ser and C34U35G36-Gln) are the same tRNA species that tended to be co-gained. While the results were not statistically significant due to the small number of instances of each type, they form a coherent hierarchy of tRNA gain. Loss event order was less clear and loss hierarchy could not be established. It is possible that the selective pressure on tRNA loss is stronger than the selective pressure on tRNA gain or that tRNA loss is more easily achieved than tRNA gain. This can lead to rapid gene eliminations that appear simultaneously.

**tRNA gain through horizontal gene transfer and anticodon mutations**

tRNA dynamics, while dominated by gene loss, also includes gain events. An interesting question regards the source of new tRNA genes in a genome. Possible mechanisms are HGT and anticodon mutations of already present tRNA genes. Tracking gene origin is difficult due to genetic variation that accumulates with time and erases sequence-embedded evidence. tRNA genes are especially problematic since in such short sequences every mutation eliminates a significant evolutionary signal. Furthermore, since tRNA function largely depends on its tertiary structure, structure-preserving mutations are more easily accumulated. It was shown in bacteria that genes of the same tRNA species are conserved in only 60–85% of their sequence (31). However, since sequence similarity rapidly deteriorates, the finding of highly similar sequences probably hints at a common ancestor that was shared not long ago. In order to find such events, we identified the MRGEs of each tRNA species (see schematic example in Figure 5A), and all the tRNA genes that descended vertically from them, and ran a BLAST search of these genes against a database containing all tRNA genes from the 319 bacteria. Since only strong sequence resemblance is likely to suggest a shared origin, the BLAST score cutoff was set at 80% of BLAST score of the query against itself (maximal score), which roughly corresponds to 3–4 mismatches or 1–2 gaps for a typical tRNA gene. Importantly, this procedure is aimed at identifying with high confidence the origin of new tRNA species. It does not provide a comprehensive inventory of HGT of tRNA genes or anticodon mutations as it misses events where gene copy number is changed (not considered as MRGE) or multiple mutations have accumulated (old events that do not pass the strict threshold). Sequence similarity was attributed to vertical inheritance if the query and hit tRNA genes had the same anticodon and descended from the same MRGE (Figure 5B) and to non-vertical inheritance otherwise. For each query, the highest scored hits in each of these two categories were considered. Since a query could show high similarity both to tRNAs belonging to the same clade (explained by vertical inheritance) and to tRNAs of distant organisms (suggesting HGT), this procedure allowed the detection of the source of a horizontally transferred tRNA to an ancestor of a bacterial clade.

Out of 403 BLAST hits that passed the threshold, the vast majority (345) was explained by vertical inheritance. We therefore concentrated on the other 58 results. In 53 out of 58 non-vertical inheritance results the query and hit had the same anticodon, but did not descend from the same MRGE event (Figure 5C), which suggests a HGT event. Twenty-two of them are presented in Figure 6A and appear to represent a single HGT event. In this example, an MRGE of C34G35U36-Thr tRNA was found in a clade belonging to the Enterobacteriaceae family. This gain event appears to be relatively recent since 11 of the organisms in the clade (out of 13 that did not lose the C34G35U36-Thr gene) also show a strong vertical inheritance signal, which
Figure 5. tRNA sequence similarity may reveal the origin of newly acquired tRNA genes. (A) A schematic example of tRNA-X ancestral reconstruction. The shade of the blue dots indicates the probability of tRNA-X presence (zero probability is not presented, low probabilities appear white). Red dots mark nodes of MRGEs, where tRNA-X gain is predicted with probability of at least 0.8, and no additional gain event of tRNA-X is found below it in the tree. Yellow dots mark all the currently extant organisms descending from an ancestor where an MRGE occurred. The grey lines connect two organisms demonstrating high tRNA gene sequence similarity according to BLAST. (B–F) Mechanisms most likely to explain tRNA gene similarity: (B) Vertical gene transfer can explain similarity of two tRNA-X genes (yellow curly lines) descending from the same MRGE. (C) Horizontal gene transfer can explain similarity of two tRNA-X genes descending from different MRGEs. (D) Horizontal gene transfer of tRNA-Y (magenta curly line) followed by mutations transforming tRNA-Y to tRNA-X can explain similarity of tRNA-X and tRNA-Y genes from distantly related organisms. (E) Similar to D but in closely related organisms. Due to vertical gene transfer, closely related organisms have a copy of tRNA-Y. A mutation that transforms tRNA-Y to tRNA-X in one organism creates very similar orthologs of different tRNA species. (F) Gene duplication followed by antidodon mutation creates very similar paralogous genes that encode different tRNA species.

indicates they had little time to diverge. These tRNA genes are very similar to two C34G35U36-Thr genes of the Neisseriaceae family (thus creating the 22 non-vertical BLAST hits), suggesting that C34G35U36-Thr tRNA gene was horizontally transferred from an organism in the Neisseriaceae family to the Enterobacteriaceae ancestor. This explanation is further supported by the fact that organisms in both families are host-associated (most are human pathogens) providing ample opportunities for sharing genetic material.

The second gain mechanism mentioned above is through mutation in the antidodon of an already present tRNA gene that transforms it from one tRNA species to another. There are three possible scenarios involving antidodon mutation: (i) tRNA gene is horizontally transferred and then undergoes antidodon mutation (Figure 5D), resulting in distantly related organisms having very similar tRNA genes that differ in their antidodon. (ii) A mutation occurs in a tRNA gene, transforming it to a different tRNA species (Figure 5E), resulting in closely related organisms having very similar orthologs that differ in their antidodon. (iii) A mutation occurs in a tRNA gene after a duplication event occurred (Figure 5F), resulting in very similar paralogs that differ in their antidodon. We identified two instances of the first scenario and three instances of the third scenario. Figure 6B and C show representative cases of each scenario. In Figure 6B, an MRGE of U34C35G36-Arg tRNA was identified in Natranaerobius thermophilus (taxonomy ID: 457570) and showed the highest similarity to the A34C35G36-Arg tRNA gene of Carboxydothermus hydrogenoformans (taxonomy ID: 246194), which belongs to an entirely different order and is only distantly related. The evolutionary distance points to a HGT and mutation rather than just a mutation. Interestingly, the presumed gene donor is an anaerobic hyperthermophile isolated from a hot swamp whereas the gene acceptor is an anaerobic thermophile isolated from a solar-heated lake. While not sharing similar tastes in pH and salinity, it is possible that at one time these organisms, their ancestors or close relatives shared the same environment and exchanged genes. The MRGE of C34C35C36-Gly tRNA in Capnocytophaga ochracea (taxonomy ID: 521097, Figure 6C) was found to be most similar to the G34C35C36-Gly tRNA gene of the same organism. This suggests that
Figure 6. tRNA genes are acquired by multiple pathways. (A–C) Similarities between tRNA genes can explain how tRNA genes are acquired. Tree visualization is as described for Figure 5A. Arrows connect organisms that have similar tRNA genes. The arrows are black if the similarity is explained by vertical inheritance and green if by non-vertical inheritance. Clades not involved in gene transfer are collapsed (black triangles). (A) C34G35U36-Thr tRNA genes in the Enterobacteriaceae family show similarity within the family and are similar to C34G35U36-Thr in the Neisseriaceae family, suggesting horizontal gene transfer of C34G35U36-Thr from the Neisseriaceae family to the Enterobacteriaceae family, followed by vertical gene spread. (B) The A34C35G36-Arg tRNA gene in Carboxydothermus hydrogenoformans (taxonomy ID: 246194) is the most similar gene to the new U34C35G36-Arg tRNA in Natranaerobius thermophilus (taxonomy ID: 457570), suggesting horizontal gene transfer followed by an A to T mutation in the first anticodon position. (C) G34C35C36-Gly tRNA gene in Capnocytophaga ochracea (taxonomy ID: 521097) is the most similar gene to the new C34C35C36-Gly tRNA of the same organism, suggesting that the G34C35C36-Gly tRNA gene underwent duplication followed by a G to C mutation in the first anticodon position. (D) Alignment of the two tRNA genes discussed in C. The duplication/mutation theory is supported by similarities in the sequences flanking the genes. The anticodons of both genes are marked in red.

the original G34C35C36-Gly gene was duplicated and then mutated from G to C in the first anticodon position. This explanation is supported by sequence similarity mostly upstream to the gene, that was not found with any other tRNA gene, as would be expected from gene duplication (Figure 6D).

These results demonstrate that tRNA genes can be gained through HGT, anticodon mutations or a combination of both. The fact that we did not find any cases of anticodon mutation without gene duplication (Figure 5E) may suggest that these genes are added to, rather than replace, existing tRNA genes, thereby reinforcing the accessory role of the gained tRNAs. Remarkably, in all five cases where an anticodon mutation was thought to explain tRNA gain, the original and the new tRNA genes always decoded the same amino acid (U34C35C36-Gly/C34C35C36-Gly, G34C35C36-Gly/C34C35C36-Gly, U34U35G36-Gln/C34U35G36-Gln and twice A34C35G36-Arg/T34C35G36-Arg). This is not surprising considering that the original genes are built to be recognized by the correct aminoacyl tRNA synthetase, and few if any additional modifications are required to achieve a functional tRNA gene with altered codon specificity.

Operon architecture effect is limited to C34U35C36-Glu and C34U35G36-Gln co-gain

Co-gain or co-loss of tRNA species could occur due to the physical proximity of their encoding genes. It is widely acknowledged that tRNA genes tend to cluster in oper-
ons, often accompanied by tRNA genes (32–36). We set out to examine if multiple events of tRNA gene gain or loss can be explained by operon gain or loss. The operon data, downloaded from MicrobesOnline, covered at least 60% of the tRNA genes of most organisms (average of 84%, Supplementary Figure S5A). The low coverage observed in 20 organisms is due to genome version differences that prevented tRNA gene mapping to operons, but is not expected to bias the results. On average, 23.8% of tRNA genes were found in an operon with other tRNA genes (Supplementary Figure S5B). This is lower than expected based on several well-annotated organisms (Bacillus subtilis 168 - 90.7%, Escherichia coli K12 MG1655 – 69.8%, Listeria monocytogenes EGD-e – 88.1%, Mycobacterium leprae TN – 35.6%, Neisseria meningitidis MC58 – 69.5%, Pseudomonas aeruginosa PA01 – 58.7%, Streptomyces coelicolor A3(2) – 29.2%; data taken from (37,38) and is probably an underestimation due to the high percent of genes that have no operon annotation. In the event of tRNA co-gain driven by a horizontally transferred operon or part of an operon, the operon structure is expected to persist in descending organisms until blurred by gene gains, losses and translocations. We therefore counted for each two tRNA species, the number of organisms descending from a co-gain event in which genes of both tRNA species were found in the same operon (Table 1). The only tRNA pair for which evidence for a co-gain event through operon transfer was found is C34U35G36-Gln/C34U35G36-Gln. Out of 12 organisms descending from C34U35G36-Gln/C34U35G36-Gln co-gain nodes in which both tRNA species were present and had operon information, in seven organisms both tRNA genes were found in the same operon. All the identified tRNAs contain only the two co-gained genes and in all of them the first gene is C34U35G36-Gln followed by C34U35G36-Gln. The operon sequences were aligned and the sequence differences were used to construct an unrooted sequence similarity tree (Supplementary Figure S6). Operons descending from a co-gain event showed higher similarity among themselves as would be expected from vertical inheritance following horizontal operon transfer. Interestingly, in Acidaminococcus fermentans DSM 20731 (Taxonomy ID: 591001) a perfect duplicate of the C34U35G36-Gln gene precedes the operon. It is possible that gaining the operon did not satisfy the need of the organism, and a duplication of C34U35G36-Gln was also required.

A similar reasoning was applied to co-loss events, where evidence for operon structure was expected to be strongest in organisms closest to the co-loss event. The organisms considered closest are those descending from the parent of the co-loss branch but not from the co-loss branch itself, where unless a gain event has occurred, the tRNA species should be missing (Supplementary Table S3). In seven pairs of tRNA species, each representing only one event of co-loss, at least one of the close organisms had an operon with genes of both tRNA species. However, in all these instances, the co-loss event is accompanied by two to 11 additional tRNA losses that have no evidence for sharing the same operon (details can be found in the text accompanying Supplementary Table S3). It therefore appears that strong selection operating in parallel to eliminate tRNA genes explains the multiple losses better than operon loss.

**DISCUSSION**

Analysis of tRNA usage has revealed a subset of tRNA species that we termed auxiliary, since they appear to supplement mandatory tRNA species in some but not all organisms. One of our main aims in this work has been to define and investigate the variability in the use of auxiliary tRNA species in the bacterial world.

**Selection for translational efficiency through tRNA reuse**

When examining the changes in tRNA repertoire in regard to ENC_diff, a measure of translational selection, we found that the presence of auxiliary tRNA species is weakly negatively associated with ENC_diff (Figure 2 and Supplementary Table S1). While the results were statistically significant independently of both GC content and genome size in only three of the tRNA species, the same trend was observed for most auxiliary tRNAs. These findings are in accord with Rocha’s conclusion that translational efficiency is associated with economy in the number of tRNA species accompanied by an increase in their gene copy number (9). We propose that the association of ENC_diff with the reduced number of tRNA species may derive from a more efficient recycling of previously used tRNAs. It was shown for Saccharomyces cerevisiae (39), and multiple bacteria (40) and archaea (41) that successive occurrences of the same amino acid favor codons that use the same tRNA. This suggests that reloading a tRNA that has just donated its amino acid to the elongating protein with a new amino acid is more efficient than waiting for a preloaded tRNA to diffuse to the translation site. Thus, if one tRNA species is the sole decoder of a set of codons, it does not matter which codon is used when the same amino acid is encoded next because all codons will be able to reuse that tRNA species. However, when more than one tRNA species is available, the situation is suboptimal. For example, the U34N35N36 tRNA decoding N1N2A3 or N1N2G3 codons can be reused to decode downstream codons of both types, while the auxiliary C34N35N36 tRNA can only be reused to decode downstream N1N2G3 codons, thus reducing translation efficiency. In the case of 4-fold degenerate amino acid decoding, maximal tRNA reuse is achieved when U34N35N36 tRNA is the sole decoder of the entire codon quartet. This type of superwobbling was shown, however, to be less efficient for the U34C1 conventional wobble interaction (42–44) and is therefore expected to be more prevalent when GC content is low; as was indeed observed (2,45,46). As shown in Figure 3, our analysis revealed as well that C34N35N36 and G34N35N36 tRNA species tend to be both absent only when GC content is low.

The association between the tRNA repertoire and GC content can be explained by selection for Watson–Crick interactions or by nucleotide bias

As GC content decreases, the use of most auxiliary tRNA species decreases. A preference of Watson–Crick over wobble interactions may provide an explanation for these results. While the N1N2U3 codon is always decoded using the conventional wobble interaction (42–44) and is therefore expected to be more prevalent when GC content is low, as was indeed observed (2,45,46). As shown in Figure 3, our analysis revealed as well that C34N35N36 and G34N35N36 tRNA species tend to be both absent only when GC content is low.
species) and the $N_1N_2A_3$ codon is almost always decoded using a Watson–Crick interaction (by the $U_{34}N_{35}N_{36}$ tRNA species), the $N_1N_2C_3$ and $N_1N_2G_3$ codons can either form a Watson–Crick interaction with the $G_{34}N_{35}N_{36}$ and $C_{34}N_{35}N_{36}$ tRNA species, respectively, or a wobble interaction otherwise (Figure 1B). The observed coordinated changes in GC content and the tRNA repertoire maximize the number of Watson–Crick interactions by matching the abundant codons with their fully complementary tRNA species. However, except for superwobbling, which was shown to be less efficient than Watson–Crick and wobble interactions, the literature is inconclusive regarding the efficiency differences of wobble and Watson–Crick interactions. Some works claim that standard wobble interactions are less efficient than Watson–Crick interactions (47–50) whereas others find similar efficiency (51,52), and sometimes even increased efficiency (53,54) of wobble compared to Watson–Crick interactions. It was shown that, at least in the case of $U_{34}N_{35}N_{36}$ tRNA species, efficiency depends on $U_{34}$ modifications (49,54,55), which are unknown for the vast majority of organisms. Furthermore, a clear preference associated with the strength of translational selection was shown for $N_1N_2U_3$ over $N_1N_2C_3$ codons in 4-fold degenerate amino acids (28), indicating selection towards wobble rather than Watson–Crick interaction. It is therefore impossible to firmly conclude that Watson–Crick interactions indeed have an advantage over wobble interactions. However, assuming that Watson–Crick interactions are preferred, the much stronger association observed between the tRNA repertoire and GC content compared to ENC'$_{diff}$ implies that Watson–Crick interactions are maximized in organisms independent of the strength of translational selection, and may suggest a basic translation optimization principle observed by all organisms. If this is true, reducing the number of tRNA species in order to improve translation efficiency should not counter the basic mechanism of maximizing Watson–Crick interactions. The ability to balance these requirements greatly depends on the GC content. When GC content is high, a minimal tRNA set that improves tRNA availability reduces Watson–Crick interactions (the abundant $N_1N_2G_3$ and $N_1N_2C_3$ codons are decoded using wobble rather than Watson–Crick interactions) and should therefore be discouraged. However, when GC content is low, the minimal tRNA set has little effect on Watson–Crick interactions (the abundant $N_1N_2A_3$ and $N_1N_2U_3$ codons continue to be decoded by Watson–Crick and wobble interactions, respectively) and is therefore preferred. Indeed, it is of note that strong translational selection manifested in high ENC'$_{diff}$ tends to occur in organisms with low GC content (data not shown) and may be responsible for the observed weak negative correlation between these two traits (reported above).

An alternative explanation for this phenomenon is that the poorly understood forces shaping nucleotide content (56,57) also influence the nucleotide content of the first anticodon position of tRNA genes. While several studies in both archaea (58) and bacteria (59,60) have shown that the nucleotide content of tRNA genes does not comply with the genomic GC content but is rather restricted by habitat temperature through constraints on folding stability, the wobble position, which does not contribute to structure stability, might be free from these constraints and conform to nucleotide content bias.

Importantly, bacterial GC content does not explain all the variation in auxiliary tRNA usage. It is conceivable that in some cases tRNA genes are gained through mobile elements such as plasmids and prophages and therefore fit the mobile element GC content rather than the GC content of the host bacteria.

### Following tRNA usage through evolution

We explored the changes in tRNA usage throughout evolution by following the tRNA repertoire of a large dataset
of bacteria within their phylogenetic context. This powerful approach revealed that auxiliary tRNAs are highly dynamic genes that tend to be lost at a high rate and also have a substantial gain probability. This analysis also provided evidence of complicated dependencies between tRNA species, hinting towards their ordered acquisition. It is, however, not clear what underlies this order. There is no trivial connection between this order and codon–anticodon interaction, as tRNAs with a very similar interaction potential, such as C34U35G36-Leu and C34A35C36-Val or C34G35G36-Pro and C34G35C36-Ala, are not similarly recruited. While it is possible that the order of tRNA recruitment in the various genomes is linked to the abundance of the codons or amino acids the tRNAs decode, which might affect the strength of selection towards their use, such an association was not identified (data not shown).

Due to specific interests of the scientific community and difficulties in maintaining the organism cultures previously required for sequencing, genome sequencing has been highly biased towards certain clades. This resulted in a few dense clades but also many sparse clades, where the tRNA repertoire has changed considerably, but the lack of bifurcations makes it impossible to understand the order of events. Adding more organisms belonging to the genera already represented is not likely to be helpful since organisms in the same genus usually have similar tRNA repertoires. However, it is possible that the rapidly growing coverage of the prokaryotic world, through culture independent and increasingly cheaper genome sequencing, will provide data to better resolve the order of tRNA gain and loss.

Despite the current data limitation, one pair of tRNA species demonstrated a striking dependency in their pattern of appearance. C34U35C36-Glu and C34U35G36-Gln are either present or absent together in approximately 90% of the organisms. They provide the most extreme example of co-gain since out of 14 and 17 gain events found for C34U35C36-Glu and C34U35G36-Gln, respectively, 10 of them co-occurred (Supplementary Figure S6). Also, while less statistically significant, they are co-lost together more than expected by chance. We also show evidence that these two tRNA species tend to be encoded in the same operon. We originally hypothesized that arranging auxiliary tRNAs in operons may be an evolutionary mechanism to facilitate coordinated changes in their genomic occurrence. However, since such arrangement was only observed for C34U35C36-Glu/C34U35G36-Gln, it does not appear that physical proximity is required in order to induce coordinated change of multiple tRNA species. In most bacteria, the glutamine-specific tRNA Gln is first aminoacylated with glutamate, which is converted to glutamine in an amidotransferase reaction requiring adenosine triphosphate (61,62). The frequent clustering of C34U35C36-Glu and C34U35G36-Gln in operons may therefore promote an efficient regulation by their shared amino acid. Incidentally, the ratio of C34U35C36-Glu and C34U35G36-Gln tRNA molecules effectively participating in translation is likely to be lower than of tRNAs of other amino acids since Glu-tRNA is also a precursor of tetrapyrole pigments (e.g. chlorophylls and hemes) (63–65) and Gln-tRNA amino acid loading is slower due to the additional step of Glu to Gln conversion. It is possible that this reduced yield decreases translational efficiency and is therefore compensated by careful fine-tuning of the tRNA repertoire.

CONCLUSIONS

The tRNA repertoire can have a tremendous impact on organism fitness. This was mostly shown through the association of tRNA gene copy number and codon usage in relation to the strength of translational selection. However, the evolution and possible effects of the types of tRNA species employed have remained mainly unexplored. We were able to show a weak negative association between the repertoire of auxiliary tRNA species and translational selection and a stronger association with nucleotide content. While the former weak association might indicate an adaptive strategy that promotes translational efficiency, it is unclear whether the latter association is adaptive or neutral. Additionally, by incorporating phylogenetic data, changes in the evolution of the auxiliary tRNA repertoire were traced. This revealed the dynamic nature of auxiliary tRNA species and points to the ease in which such genes can be acquired and lost. We believe that the approach we describe for phylogeny-driven analysis of the auxiliary tRNA repertoire coupled with the continuously growing number of sequenced genomes will provide a better resolution of the dynamics of the tRNA repertoire and determine whether tRNA recruitment is ordered, and if so, which properties underlie this order.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENT

We thank Ofir Cohen, Andrew Meade and Mark Pagel for helpful discussions, and Yael Altuvia and Asaf Peer for their useful comments.

FUNDING

Israel Science Foundation administered by the Israeli Academy for Sciences and Humanities (1411/13 to H.M.); Ministry of Science and Technology Eshkol scholarship (to N.W.). Funding for open access charge: The Israel Science Foundation administered by the Israeli Academy for Sciences and Humanities (1411/13).

Conflict of interest statement. None declared.

REFERENCES

1. Agris,P.F. (2004) Decoding the genome: a modified view. Nucleic Acids Res., 32, 223–238.
2. March,C. and Grosjean,H. (2002) tRNomics: analysis of tRNA genes from 50 genomes of Eukarya, Archaea, and Bacteria reveals anticondor-sparing strategies and domain-specific features. RNA, 8, 1189–1232.
3. Ikemura,T. (1981) Correlation between the abundance of Escherichia coli transfer RNAs and the occurrence of the respective codons in its protein genes: a proposal for a synonymous codon choice that is optimal for the E. coli translational system. J. Mol. Biol., 151, 389–409.
4. dos Reis,M., Savva,R. and Wernisch,L. (2004) Solving the riddle of codon usage preferences: a test for translational selection. Nucleic Acids Res., 32, 5036–5044.
13. Koonin, E.V., Wolf, Y.I. and Puigbo, P. (2009) The phylogenetic forest
11. Novoa, E.M., Pavon-Eternod, M., Pan, T. and Ribas de Pouplana, L.
24. Letunic, I. and Bork, P. (2011) Interactive Tree Of Life v2: online tool for phylogenetic tree display and annotation.
19. Pagel, M. (1997) Inferring evolutionary processes from phylogenies.
14. Ciccarelli, F.D., Doerks, T., von Mering, C., Creevey, C.J., Snel, B. and
10. Higgs, P.G. and Ran, W. (2008) Coevolution of codon usage and tRNA genes leads to alternative stable states of biased codon usage.
26. Bibb, M.J., Findlay, P.R. and Johnson, M.W. (1984) The relationship between base composition and codon usage in bacterial genes and its use for the simple and reliable identification of protein-coding sequences.
22. Cohen, O. and Pupko, T. (2011) Inference of gain and loss events from horizontally transferred gene families using stochastic mapping.
9. Rocha, E.P. (2004) Codon usage bias in prokaryotic pyrimidine-ending codons is associated with the degeneracy of the encoded amino acids. 

Nucleic Acids Res., 40, 7074–7083.
29. Felsenstein, J. (1985) Phylogenies and the comparative method. Am. Nat., 125, 1–15.
30. Ives, A.R. and Garland, T. Jr (2010) Phylogenetic logistic regression for binary dependent variables. Syst Biol., 59, 9–26.
31. Saks, M.E. and Conery, J.S. (2007) Anticodon-dependent conservation of bacterial tRNA gene sequences. RNA, 13, 651–660.
32. Nakajima, N., Ozeki, H. and Shimura, Y. (1981) Organization and structure of an E. coli tRNA operon containing seven tRNA genes. Cell, 23, 239–249.
33. Fournier, M.J. and Ozeki, H. (1985) Structure and organization of the transfer ribonucleic acid genes of Escherichia coli K-12. Microbiol. Rev., 39, 379–397.
34. Vold, B.S., Gjørv, C., Narasimhan, N., Strem, M. and Hansen, J.N. (1988) Transcriptional analysis of Bacillus subtilis tRNA-rRNA operons. II. Unique properties of an operon containing a minor 5 S rRNA gene. J. Biol. Chem., 263, 14485–14490.
35. Giroux, S. and Cedergren, R. (1989) Evolution of a tRNA operon in gamma purple bacteria. J. Bacteriol., 171, 6446–6454.
36. Ghatak, A., Majumdar, A. and Ghosh, R.K. (2005) Structural organization of the transfer RNA operon I of Vibrio cholerae: differences between classical and El Tor strains. J. Bacteriol., 187, 469–474.
37. Pettersson, F.B.M. (2009) Ph.D Thesis, uppsala universitet.
38. Ardell, D.H. and Kirsebom, L.A. (2005) The genomic pattern of tDNA operon expression in E. coli. PLoS Comput. Biol., 1, e12.
39. Cannarozzo, G., Schraudolph, N.N., Faty, M., von Rohr, P., Fribert, M.T., Roth, A.C., Gonnet, P., Gonnet, G. and Barial, Y. (2010) A role for codon order in translation dynamics. Cell, 141, 355–367.
40. Shao, Z.Q., Zhang, Y.M., Feng, X.Y., Wang, B. and Chen, J.Q. (2012) Synonymous codon ordering: a subtle but prevalent strategy of bacteria to improve translational efficiency. PLoS One, 7, e33547.
41. Zhang, Y.M., Sun, X.Q., Qiao, Y.F., Chen, J.Q. and Wang, B. (2013) Non-random arrangement of synonymous codons in archaea coding sequences. Genomics, 101, 362–367.
42. Rogalski, M., Karcher, D. and Bock, R. (2008) Superwobbling facilitates translation with reduced tRNA sets. Nat. Struct. Mol. Biol., 15, 192–198.
43. Alkatib, S., Scharff, L.B., Rogalski, M., Fleischmann, T.T., Matthes, A., Seeger, S., Schottler, M.A., Ruf, S. and Bock, R. (2012) The contributions of wobbling and superwobbling to the reading of the genetic code. PLoS Genet., 8, e1003076.
44. Kotche, U. and Podruma, M.V. (2007) Codon reading by tRNAAla with modified uridine in the wobble position. Mol. Cell, 25, 167–174.
45. Andachi, Y., Yamao, M., Muto, A. and Osawa, S. (1989) Codon recognition patterns as deduced from sequences of the complete set of transfer RNA species in Mycoplasma capricolum. Resemblance to mitochondria. J. Biol. Chem., 209, 37–54.
46. Chamband, I., Heilig, R., Ferriss, S., Barbe, V., Samson, D., Galisson, F., Mosser, I., Dyvig, K., Wroblewski, H., Vare, A. et al. (2001) The complete genome sequence of the murine respiratory pathogen Mycoplasma pulmonis. Nucleic Acids Res., 29, 2145–2153.
47. Thomas, L.K., Dix, D.B. and Thompson, R.C. (1988) Modified uridines with C5-methylene substituents at the first position of tRNA. J. Biol. Chem., 263, 6552–6558.
48. Kirino, Y., Hori, N., Yamaizumi, Z., Nishimura, S., Miyazaki, T. and Yokoyama, S. (1994) Recognition of UUN codons by three leucine tRNA species from Escherichia coli. FEMS Lett., 134, 31–34.
49. Ogle, J.M., Murphy, F.V., Tarry, M.J. and Ramakrishnan, V. (2002) Selection of tRNA by the ribosome requires a transition from an open to a closed form. Cell, 111, 721–732.
50. Kurata, S., Weixlbaumer, A., Ohtsuki, T., Shimazaki, T., Wada, T., Kirino, Y., Takai, K., Watanabe, K., Ramakrishnan, V. and Suzuki, T. (2008) Modified uridines with C5-methylene substituents at the first
position of the tRNA anticodon stabilize U.G wobble pairing during decoding. *J. Biol. Chem.*, **283**, 18801–18811.

53. Mitra, S.K., Lustig, F., Akesson, B., Axberg, T., Elias, P. and Lagerkvist, U. (1979) Relative efficiency of anticodons in reading the valine codons during protein synthesis in vitro. *J. Biol. Chem.*, **254**, 6397–6401.

54. Takai, K., Okumura, S., Hosono, K., Yokoyama, S. and Takaku, H. (1999) A single uridine modification at the wobble position of an artificial tRNA enhances wobbling in an Escherichia coli cell-free translation system. *FEBS Lett.*, **447**, 1–4.

55. Murphy, F.V.t., Ramakrishnan, V., Malkiewicz, A. and Agris, P.F. (2004) The role of modifications in codon discrimination by tRNA(Lys)UUU. *Nat. Struct. Mol. Biol.*, **11**, 1186–1191.

56. Lind, P.A. and Andersson, D.I. (2008) Whole-genome mutational biases in bacteria. *Proc. Natl. Acad. Sci. U.S.A.*, **105**, 17878–17883.

57. Hildebrand, F., Meyer, A. and Eyre-Walker, A. (2010) Evidence of selection upon genomic GC-content in bacteria. *PLoS Genet.*, **6**, e1001107.

58. Kawai, Y. and Maeda, Y. (2009) GC-content of tRNA genes classifies archaea into two groups. *J. Gen. Appl. Microbiol.*, **55**, 403–408.

59. Hansen, A.K. and Moran, N.A. (2012) Altered tRNA characteristics and 3’ maturation in bacterial symbionts with reduced genomes. *Nucleic Acids Res.*, **40**, 7870–7884.

60. Dutta, A. and Chaudhuri, K. (2010) Analysis of tRNA composition and folding in psychrophilic, mesophilic and thermophilic genomes: indications for thermal adaptation. *FEMS Microbiol. Lett.*, **305**, 100–108.

61. Sheppard, K., Yuan, J., Hohn, M.J., Jester, B., Devine, K.M. and Soll, D. (2008) From one amino acid to another: TRNA-dependent amino acid biosynthesis. *Nucleic Acids Res.*, **36**, 1813–1825.

62. Guo, L.T., Helgadottir, S., Soll, D. and Ling, J. (2012) Rational design and directed evolution of a bacterial-type glutaminyl-tRNA synthetase precursor. *Nucleic Acids Res.*, **40**, 7967–7974.

63. Stange-Thomann, N., Thomann, H.U., Lloyd, A.J., Lyman, H. and Soll, D. (1994) A point mutation in Euglena gracilis chloroplast tRNA(Glu) uncouples protein and chlorophyll biosynthesis. *Proc. Natl. Acad. Sci. U.S.A.*, **91**, 7947–7951.

64. Luer, C., Schauer, S., Mobius, K., Schulze, J., Schubert, W.D., Heinz, D.W., Jahn, D. and Moser, J. (2005) Complex formation between glutamyl-tRNA reductase and glutamate-1-semialdehyde 2,1-aminomutase in Escherichia coli during the initial reactions of porphyrin biosynthesis. *J. Biol. Chem.*, **280**, 18568–18572.

65. Randau, L., Schauer, S., Ambrogelly, A., Salazar, J.C., Moser, J., Sekine, S., Yokoyama, S., Soll, D. and Jahn, D. (2004) tRNA recognition by glutamyl-tRNA reductase. *J. Biol. Chem.*, **279**, 34931–34937.