

# Coding triplets in the tRNA acceptor-TΨC arm and their role in present and past tRNA recognition

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**The mechanism and evolution of the recognition scheme between key components of the translation system, that is, tRNAs, synthetases, and elongation factors, are fundamental issues in understanding the translation of genetic information into proteins. Statistical analysis of bacterial tRNA sequences reveals that for six amino acids, a string of 10 nucleotides preceding the tRNA 3' end carries cognate coding triplets to nearly full extent. The triplets conserved in positions 63–67 are implicated in the recognition by the elongation factor EF-Tu, and those conserved in positions 68–72, in the identification of cognate tRNAs, and their derived minihelices by class IIa synthetases. These coding triplets are suggested to have primordial origin, being engaged in aminoacylation of prebiotic tRNAs and in the establishment of the canonical codon set.**

**Keywords:** aminoacylation; genetic code; origin of life; synthetase; translation; tRNA

tRNA is a vital component of the translation system due to its role in linking the genetic code with the synthesized protein. The modern tRNA is an L-shaped molecule. One end of the molecule is the universally conserved single-stranded NCCA tail, which carries the cognate amino acid (aa), while at the other end, about 75 Å away, the anticodon (AC) loop that fully characterizes the attached aa is located.

The mature tRNA interacts with three key components of the translation system—the cognate aminoacyl-tRNA synthetase (aaRS) that specifically aminoacylates it; the elongation factor (EF-Tu in bacteria) that accommodates the aa-tRNAs and carries them, in a ternary complex, to the ribosome; and the ribosome where the attached aa is incorporated into the growing polypeptide. The low overall frequency of

aa misincorporation in translation ( $10^{-3}$  to  $10^{-4}$ ) reflects the cumulative fidelity of these three principal reactions [1].

Recognition of aa-tRNA on the ribosome is accomplished *via* the interactions of the mRNA codon accommodated on the small ribosomal subunit, with the AC loop. EF-Tu, which carries a variety of different tRNAs, recognizes each of them specifically. The minimal fragment of tRNA that can interact efficiently with EF-Tu consists of a 10 base-pair helix from the acceptor-TΨC stem, linked to the aa-3' end tail [2], from which nucleotides 63–67 form the contact interface [3].

Aminoacyl-tRNA synthetases are divided into two distinct classes: Class I synthetases possess active sites with the Rossmann fold motif, while class II

## Abbreviations

AC, anticodon; sC/sAC, stem codon and/or stem anticodon; aa, amino acid; aaRS, aminoacyl-tRNA synthetase; pre-3' end, a string of ten nucleotides preceding the tRNA 3' end.

synthetases have active sites composed of antiparallel  $\beta$ -sheets with three highly conserved motifs: 1, 2, and 3 [4]. Usually, there is one synthetase for each aa, 10 of them belonging to class I and 10 to class II, but LysRS has a class I form in addition to the canonical class II synthetase [5]. Synthetases commonly acquire the primary identity of their cognate tRNAs from the anticodon stem loop. Yet, this loop is not always the principal determinant for aminoacylation [6,7] and in some cases no physical contact is made between the aaRS and the anticodon loop [8]. In several tRNAs, major identity elements are found in the core of the molecule, as is the case, for example, with the large variable loop in tRNA<sup>Ser</sup>. An additional set of identity elements classified as ‘determinants’ and ‘antideterminants’ [9], such as the discriminator base (N73) [10] and specific nucleotides in the acceptor stem [11–13], were shown to participate in the tRNA recognition by synthetases.

The recognition determinants embedded in the first base pairs of the tRNA acceptor stems, as well as in RNA minihelices derived from their cognate acceptor-TΨC stems, were demonstrated to carry coding information sufficient for their specific aminoacylation [9,11–16]. This non-AC aminoacylation process was suggested to be controlled by an ‘operational code’ [12] that relates few nucleotides in the acceptor stem to specific aa. Its mode of action was proposed to be a remnant of an early mechanism for specifically aminoacylating proto-tRNAs by proto-synthetases, possibly associated with a second code [12,17], distinct from the nucleotide triplets of the standard genetic code [16].

Here, we analyze large sequence data sets from bacterial tRNA genes and report the occurrence, to a very high extent, of cognate coding triplets in specific locations of the acceptor-TΨC arm, also known as ‘tRNA acceptor branch’, belonging to six aa: Ala, Asp, Gly, His, Pro, and Ser. Coding triplets conserved in positions 63–67 are implicated in the tRNA recognition by EF-Tu, while the triplets conserved in the 68–72 nucleotide range are proposed to be involved in the contemporary tRNA recognition by synthetases of class IIa. In the evolutionary context, these conserved coding triplets are suggested to have served as identifiers of proto-tRNAs in the primordial aminoacylation process and to have had a primal role in the emergence of the contemporary codon set.

## Results

In bacterial tRNAs, the sequence of the string preceding the 3’ end (pre-3’ end), ranging from nucleotides

63 to 72, is diverse [18], even among different strains of the same bacterium (Table 1, File S1). Statistical analysis was performed on large-scale sequence data of the pre-3’ end strings, acquired from the tRNADB database [19,20], <http://trna.bioinf.uni-leipzig.de>, which holds 128 bacterial tRNA sequences obtained from elongator tRNAs involved in protein synthesis, and over 6200 bacterial tRNA genes, belonging to the 20 canonical aa. To avoid the inclusion of pseudo-tRNA genes, that is, those not coding for elongator tRNAs, only tRNA gene sequences having a full 3’ end and a consensus discriminator base [10] were incorporated into the statistics (Materials and methods).

When read from the 5’ to the 3’, these strings, in the tRNAs of nine aa, were found to hold excessive cognate codon and/or anticodon triplets, termed here sC/sACs, that is, stem codons and/or stem anticodons (Tables 1, Table 2 and S1a, File S1). One class of bacteria, the *Mollicutes*, deviated from the statistical behavior exhibited by the other bacteria (Table 1, File S1, Table S2), and its exclusion resulted in a sample of 3324 tRNA gene sequences from over 100 bacterial species. Statistical analysis that incorporated the 350 tRNA genes of *Mollicutes* did not differ significantly from the results obtained with the data sample used in the current study.

### A group of nine conserving amino acids

The occurrence of the cognate codons and anticodons in the pre-3’ end strings of the 20 canonical aa was determined. Two distinct groups with a wide gap between their cognate sC/sAC occurrences were observed: The pre-3’ end strings of the members of the first group, that is, Ala, Arg, Asp, Gly, His, Leu, Pro, Ser, and Val, were found to contain cognate sC/sACs in over 85% of the strings of each aa. These aa were denoted ‘conserving aa’. For the remaining 11 ‘non-conserving aa’, that is, Asn, Cys, Gln, Glu, Ile, Lys, Met, Phe, Thr, Trp, and Tyr, only 0%–40% of their pre-3’ end strings carried a cognate sC/sAC (Table S1b).

For each of the aa of the first group, the set of specific coding triplets that accounts for 98% of the sC/sAC occurrence within the strings carrying any cognate sC/sAC is denoted the ‘conserved sC/sAC set’, and the codons and anticodons belonging to any conserved set are named ‘conserved sC/sACs’ (Table 2; Materials and methods). These conserved sC/sACs are present in 83% or more of the pre-3’ end strings of each conserving aa, and they typically constitute only a subset of the corresponding standard codon and anticodon table (Table 2). Interestingly, the cognate

**Table 1.** Conserved sC/sAC in the tRNA pre-3' end sequences.

Amino acid (codons/anti-codons)	Pre-3' end sequence (nucleotides 63–72) <sup>b</sup>	Organism	Code in tRNAdb <sup>a</sup>
Ala (Conserved codons-GCC, GCU)	<b>GCUAGGCUCC</b>	<i>Bacillus subtilis</i>	tdbR00000007
	<b>GCUUAGCUCC</b>	<i>Escherichia coli</i>	tdbR00000008
	GCAU <b>AGCUCC</b>	<i>Escherichia coli</i>	tdbR00000010
	GUUAGGGUCC	<i>Mycoplasma capricolum</i>	tdbR00000005
Pro (Conserved codon-CCG)	UGUCUUC <b>CCG</b>	<i>Bacillus subtilis</i>	tdbR00000316
	UCUAUC <b>CCG</b>	<i>Escherichia coli</i>	tdbR00000623
	UCUCGUG <b>CCG</b>	<i>Salmonella typhimurium</i>	tdbR00000318
	UGUCUUC <b>CCG</b>	<i>Mycoplasma mycoides</i>	tdbR00000413
Gly (Conserved Anticodons-UCC, CCC)	CAUCA <b>CCCGC</b>	<i>Bacillus subtilis</i>	tdbR00000116
	CUUCG <b>CCCGC</b>	<i>Escherichia coli</i>	tdbR00000117
	CGUU <b>CCCGC</b>	<i>Escherichia coli</i>	tdbR00000118
	CGCUG <b>CCCGC</b>	<i>Salmonella typhimurium</i>	tdbR00000121
	UAUCU <b>CCCGC</b>	<i>Staphylococcus epidermidis</i>	tdbR00000114
	CGUCA <b>CCCGC</b>	<i>Streptomyces coelicolor_A3</i>	tdbR00000113
	CUUCAC <b>CCUGC</b>	<i>Mycoplasma capricolum</i>	tdbR00000111

<sup>a</sup>All the unique sequences of the elongator tRNAs of Ala, Pro, and Gly, as found in the tRNAdb database [19,20].; <sup>b</sup>The conserved codons are bolded, and the anticodons are underlined. In tRNA<sup>Ala</sup>, the location of the conserved coding triplets varies, and in tRNA<sup>Pro</sup> (codon) and in tRNA<sup>Gly</sup> (anticodon), it keeps a fixed position. The *Mycoplasma* (genus of the *Mollicutes* class) sequences, at the bottom of each box, often do not comply with the other bacteria.

**Table 2.** Statistics of the conserved sC/sACs in the pre-3' end strings from bacterial tRNA genes. N—any nucleotide. P—the probability that the observed occurrence of a particular sC/sAC set is random.  $P \leq 0.01$  is considered statistically significant.

Amino acid <sup>a</sup> (no. sequences)	Conserved sC/sAC set <sup>a</sup>		Observed occurrence <sup>b</sup>	Expected occurrence <sup>c</sup> ( $\pm 0.8\%$ )	P
	Codons	Anticodons			
<b>Ala</b> (209)	<b>GCU</b> , GCC		83%	50%	E-22
<b>Asp</b> (110)		<b>GUC</b>	96%	17%	E-106
<b>Gly</b> (243)		<b>CCC, UCC</b>	94%	57%	E-31
<b>His</b> (83)	<b>CAC, CAU</b>		90%	22%	E-51
<b>Pro</b> (180)	<b>CCG</b>		98%	37%	E-64
<b>Ser</b> (290)	<b>UCA, UCC, UCU</b>		99%	46%	E-72
Arg (305)	<b>CGG, AGG</b> CGA, CGC, CGU	CCU	95%	84%	E-7
Leu (366)	CUN	CAG	84%	63%	E-17
Val (201)	GUN	NAC	88%	58%	E-17

<sup>a</sup>Cognate triplets whose conservation is considered as robust, as well as the robustly conserving aa (Materials and methods), are depicted by bold letters.; <sup>b</sup>% of pre-3' end strings holding at least one sC/sAC from their conserved set at any location along the string.; <sup>c</sup>Derived from random oligonucleotides simulation with 'sample' distribution (A, C, G, U) = (10%, 46%, 24%, 20%). The statistics for the other two distributions and the statistics for the three distributions calculated for the remaining 11 aa are given in Table S1a,b, respectively.

sC/sACs conserved in the pre-3' end strings are often not correlated with the anticodon triplet in the AC loop of the same tRNA (File S1).

For each of the nine conserving aa, the observed occurrence of conserved coding triplets in their pre-3' end was compared with its expected occurrence. The expected occurrences were obtained by analyzing a large data set of random 10-mer oligonucleotides. The construction of the randomized sequences and the estimation of the expected occurrence of a specific set of sC/sACs are described in the Materials and methods section. The randomized strings were constructed with

three nucleotide distributions: (a) distribution identical to the average found in all the bacterial pre-3' end strings analyzed ('sample' distribution), that is, A = 10%, C = 46%, G = 24%, and T = 20%; (b) equal distribution of the four nucleotides; and (c) the individual nucleotide distribution found in the pre-3' end string of each aa (Table S1a,b).

Analysis of the observed vs. expected sC/sAC occurrences in the data of the conserving aa confirmed that the observations are clearly nonrandom, that is, that the conserved sC/sAC of the nine conserving aa show observed occurrences that go far beyond the statistical

expectation ( $P \ll 0.01$ ), a result obtained with all three distributions (Tables 2 and S1a). For the non-conserving aa, in the absence of a conserved sC/sAC set, only the combined occurrence of all the cognate sC/sACs was checked. The combined occurrences were found to lie in the range of 0–40% for each aa, which, for Gln, Glu, Lys, Met, Thr, and Tyr, is less than statistically expected. For the remaining five aa, it practically accords with the statistical expectations (Table S1b). It is noteworthy that the coding triplets' statistics in the Thr pre-3' end strings demonstrate idiosyncratic features—it has the highest level of sC/sAC occurrence among the nonconserving aa, 40%, compared with less than 25% in the data of the 10 remaining nonconserving aa. Moreover, 90% of the observed sC/sAC are anticodons, and not even a single sC/sAC is found in positions 70–72 of tRNA<sup>Thr</sup>.

To further benchmark these statistical results, the occurrence of the triplet CCC was examined. This triplet is expected to be the most prevalent for 14 out of 20 aa due to the considerable excess of the nucleotide C in their sequences (Fig. S1). Pre-3' end strings from the tRNAs of the conserving aa display CCC considerably less than inferred from their nucleotide distribution. Surprisingly, the strings of Ala, His, and Asp, which contain over 44% of nucleotide C in their pre-3' end sequences, hardly show any CCC triplets. Conversely, aa whose tRNAs display sC/sACs significantly less than statistically expected (Gln, Tyr, Lys) exhibit the most significant excess of CCC.

### A subgroup of six robustly conserving amino acids

The sC/sACs found in the pre-3' end string of the conserving aa may appear as a single site, which is fully conserved, although the rest of the sequence varies (Pro, Table 1, Fig. 1) or as an accumulation of occurrences from as many as seven different sites (Leu, Val, Fig. 1). The conserved sC/sACs of Ala, Arg, Asp, Gly, His, Pro, and Ser are concentrated in one or two sites, each holding mainly one or two coding triplets. In contrast, the overall high occurrence of the conserved sC/sACs of Leu and Val is the sum of small occurrences scattered throughout their pre-3' end strings.

Components of a conserved set are considered as 'robustly conserved sC/sACs' when they appear, alone or with one additional coding triplet, at a specific location in more than 60% of the pre-3' end strings belonging to a certain aa. This location is termed a 'sC/sAC conservation site'. Overall, the occurrence histogram (Fig. 1) contains nine such sites: one for Pro (positions 70–72, 98% occurrence of the codon CCG); Gly (positions

68–70, 94% occurrence of the anticodon CCC or UCC); Asp (positions 64–66, 91% occurrence of the anticodon GUC); and Ala (positions 68–70, 64% occurrence of the codon GCU. In the remaining sequences, only C69 and U70 are conserved); Arg (positions 66–68, 60% occurrence of the codons CGG or AGG); two sites for Ser (in two partially overlapping positions: 68–70, 63% occurrence of the codon UCU or UCA and positions 70–72, 72% occurrence of UCC, which combine to 94% occurrence of a conserved codon in positions 68–72); and two for His (positions 63–65, 60% occurrence of the codon CAU. In the remaining sequences, only C63 and U65 are conserved. Positions 70–72, 69% occurrence of the codon CAC. In the remaining sequences, only C70 and C72 are conserved, mostly due to the exchange of A71 into G71 in the sequences from *Bacillus*). The corresponding aa and coding sets (Table 2) are termed 'robustly conserving aa' and 'robustly conserved sC/sAC sets' (Materials and methods). Interestingly, in contrast to the accumulation of coding triplets in positions 68–70 and 70–72 (Fig. 1), there are hardly any conserved triplets in positions 69–71.

In a second benchmark test, the occurrences of individual coding triplets per each three-mer location in the pre-3' end strings of every aa were counted, without relating to the nucleotide distribution. Cognate sC/sACs were found at a specific location in more than half of the strings belonging to each of the six robustly conserving aa, but not in the data of the other 14 aa.

Five out of the six robustly conserving aa, that is, Ala, Gly, His, Pro, and Ser, have synthetases that belong to class IIa. The statistical correlation between the list of aa that robustly conserve the sC/sACs, that is, Ala, Asp, Gly, His, Pro, and Ser, and those whose synthetases belong to class IIa, that is, Ala, Gly, His, Pro, Ser, and Thr [21], was computed. The probability  $P$  that such a correlation or higher, could have been obtained randomly, is given by (Materials and methods):

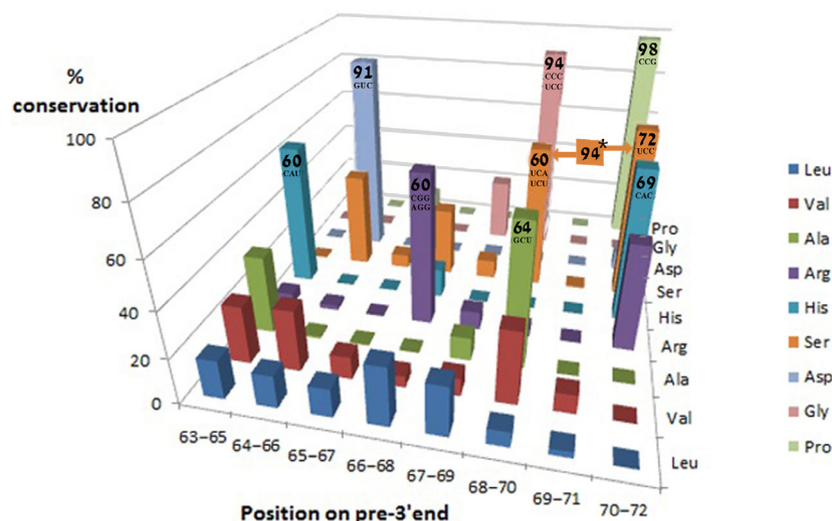
$$P = \frac{\binom{6}{6} \cdot \binom{14}{0} + \binom{6}{5} \cdot \binom{14}{1}}{\binom{20}{6}} \approx 0.002.$$

indicating that the null hypothesis of no correlation between robust conservation and belonging to class IIa is wrong.

### Archaea and eukarya

The genomic tRNA sequences of archaea and eukarya were not subjected to the aforementioned analysis due

**Fig. 1.** Occurrence of the conserved stem codon and/or anticodon triplets in the pre-3' end string of the conserving aa. The column of each 'sC/sAC conservation site' is marked by the identity of its conserved triplets and by the percentages of sequences carrying these triplets in the site. \*The overall occurrence of the conserved sC/sACs of Ser in its two partially overlapping conservation sites (positions 68–70 and 70–72).



to the difficulty to distinguish pseudo-tRNA gene sequences, which are prevalent in the database (Sprinzl M., private communication), from the authentic ones. The 3' end tail, NCCA, used for screening the bacterial data, which excluded 40% of the bacterial sequences, appears only in few archaeal tRNA genes and is absent in eukaryotes. The more reliable elongator tRNA sequences are too scarce to ensure any statistical validity. Part of the 63 unique archaeal and 117 unique eukaryotic sequences, belonging to the elongator tRNAs of the 20 aa, do display access of cognate sC/sACs in their pre-3' end strings (Files S2, S3), mostly for the same aa classified as conserving in bacteria. However, the position and identity of these cognate coding triplets differ from bacteria, leaving open the question of whether the cognate sC/sACs are conserved in the other realms of life besides bacteria.

## Discussion

### Coding triplets in the acceptor-TΨC stem

In total, over 90% of the bacterial pre-3' end sequences from the nine conserving aa contain a conserved codon and/or anticodon triplet (Table 2), although the data probably include unidentified pseudo-tRNA genes randomly carrying the consensus discriminator base, and possibly sequences from additional species besides the *Mollicutes*, whose tRNAs do not comply with the bacterial pattern of sC/sAC conservation. Coding triplets, which are complementary to the conserved sC/sACs presented here, were previously identified in positions 3–5 of prokaryotic tRNAs of four of the conserving aa, that is, Ala, Asp, Gly, and Val,

and were termed 'protocodons' and 'proto-anticodons' [22]. Collectively, these observations point to a highly nonrandom occurrence of conserved sC/sACs, distributed along the pre-3' end strings of the conserving aa.

Nucleotide C is the most abundant (47%) of the nucleotides constituting the 12 robustly conserved sC/sAC triplets (Table 2). The conservation is therefore facilitated by the elevated occurrence of nucleotide C (46%) in the bacterial pre-3' end sequences. Yet, the sC/sAC occurrence in the pre-3' end strings of the conserving aa is of consequence with regard to both the expectation values derived from the randomized oligonucleotides constructed with the C-rich 'sample' distribution (Table 2) and with the individual nucleotide distributions (Table S1a), validating that the conservation of the cognate stem codons and/or anticodons in the pre-3' end string is a genuine phenomenon. Correspondingly, the triplet CCC, which is expected to be highly frequent due to the significant access of nucleotide C in the pre-3' end of 14 aa, is nearly absent in the sequences from the tRNAs of the robustly conserving aa, likely due to a negative correlation with the marked occurrence of the conserved sC/sACs (Fig. S1).

Our statistical results are in line with the ensemble attributes of the 'operational RNA codes', determined from a bacterial classification and regression tree, which was inferred from acceptor stem sequences of a large variety of bacterial organisms [23]. The tree exhibited low degree of degeneracy for the same five aa that display robustly conserved sC/sACs in their acceptor stem, that is, Ala, Gly, His, Pro, and Ser. In contrast, it presented the highest degree of degeneracy for Arg, Leu, and Val, indicating the weakening of the

selective constraints on their tRNA acceptor stem identity elements. The clear accumulation of most species corresponding to each of the five robustly conserving aa, into a single leaf of the tree, corroborates the conservation of coding information in a specific location of their acceptor stems. Likewise, the high degeneracy obtained for Arg, Leu, and Val agrees with the dispersion of the cognate coding triplets throughout their acceptor-TΨC stem (Fig. 1), found in the present study.

### Contemporary significance of the conserved coding triplet

The unexpected retention of cognate coding triplets at specific locations of the acceptor-TΨC stems of Ala, Asp, Gly, His, Pro, and Ser poses the question of the biological significance behind this extreme conservation (Tables 2 and S2).

Five of the robustly conserving aa, that is, Ala, Gly, His, Pro, and Ser, exhibit conserved sC/sACs in the nucleotide range of 68–72 of the acceptor stem (Fig. 1), which forms a contact interface with synthetase [9]. The tRNAs of all these aa are charged by class IIa synthetases, when partition into subclasses is made according to the relevant mode of their binding to the tRNA acceptor stem [21]. The probability that this correlation is random is negligible, 0.2%, suggesting that the high conservation of coding triplets in positions 68–72 relates to their involvement in the aminoacylation mode applied by their cognate class IIa synthetases. This assertion is further supported by the significant decrease in aminoacylation rate of the full-length tRNA<sup>Pro</sup> on mutation of C70 and G72 [24], from the robustly conserved Pro stem codon CCG located in positions 70–72 (Fig. 1).

A plausible aminoacylation mechanism that uses the robustly conserved sC/sACs to discriminate between tRNAs or between their derived minimal substrates requires specific interaction between the catalytic core of the corresponding synthetases and the conserved sC/sACs. The flexible motif 2 loop from the catalytic core of class II synthetases was found to interact, in a base-specific fashion, with nucleotides from the tRNA acceptor stem. In the tRNA<sup>Ser</sup>:SerRS complex, it protrudes into the major groove down to the fifth base pair, interacting with the nucleotide bases [25], that is, with the same nucleotides that are robustly conserved and that constitute the identity determinants for tRNA<sup>Ser</sup> recognition [15]. In the tRNA<sup>His</sup>:HisRS complex (Fig. 2), residues R115 and Q117 from motif 2 loop approach the bases of A71 and C72 to 3.7 and 3.9 Å, respectively, where A71 and C72 make part of

the His robustly conserved codon CAC, located in positions 70–72 of tRNA<sup>His</sup> (Fig. 1). This mode of interaction provides a feasible mechanism for the specific recognition of these tRNAs, either through the involvement of the conserved coding triplets in stabilizing a specific conformation identified by the synthetase, or *via* direct contacts of motif 2 loop with the bases of the conserved coding triplets.

Linking the sC/sAC conservation with their being recognized by motif 2 loop is consistent with the lack of sC/sAC conservation in the pre-3' end strings of the sixth aa charged by a class IIa synthetase, Thr (Table S1b). In the structure of the bacterial tRNA<sup>Thr</sup>:ThrRS complex, the identity determinants held in the stem are recognized by the N-terminal domain of the enzyme [26] and not by motif 2 loop, a functional exception that can account for the lack of sC/sAC conservation. Likewise, the partial loss of sC/sAC occurrence in tRNA<sup>Ala</sup> acceptor stem (Table 2, Fig. 1) can be linked with the reorientation of tRNA<sup>Ala</sup> relative to the core of its cognate synthetase, as seen in the tRNA<sup>Ala</sup>:AlaRS complex from the archaeon *Archaeoglobus fulgidus* (pdb code 3wqy), which is the only currently available X-ray structure of the complex. Although class II consensus catalytic core exists within this AlaRS structure, motif 2 loop does not interact with the acceptor stem. Recognition of the acceptor stem identity determinants is carried out by the helical loops of α11 and α14, from out of the catalytic core [27], a functional exception that is likely to account for the diminishing sC/sAC conservation in the modern tRNA<sup>Ala</sup> acceptor stems. The partial conservation of the CAC codon in positions 70–72 of tRNA<sup>His</sup>, which is manifested mainly in the exchange of A71 into G71 in the sequences from the *Bacilli* class, can be possibly linked with a functional variation as well. A second type of synthetase, histidine tRNA ligase 2, found solely in *Bacillus* [28], may render the sC/sAC conservation in this phylogenetic class surplus.

Involvement of the robustly conserved sC/sACs in the aminoacylation by five class IIa synthetases is further corroborated by the properties of the aminoacylation of minimal RNA substrates. In this process, information contained in the acceptor stem was demonstrated to suffice for charging minihelices derived from the acceptor-TΨC stem, with their cognate aa [9,11–16]. This alternative information source, termed the ‘operational code’ [12], depends on identity determinants held in the first five base pairs of the acceptor stem analogs. Aminoacylation of such minihelices was found to be carried out most efficiently for four out of five robustly conserving aa, that is, Ala, Gly, His, and Ser [29], while no data are currently available for fifth amino acid, Pro. All but one of the

identity determinants recognized by the corresponding synthetases, in the 3' side of the cognate minihelices, are contained in the robustly conserved coding triplets of these aa (Table 3). The coding triplets present in these minihelices can therefore be specifically recognized by motif 2 loop, *via* the same mechanism suggested for the interaction of the loop with the identity determinants located in acceptor stems of full-length tRNAs. This interaction is likely to facilitate the identification of these minihelices and thus grant them with enhanced efficiency in aminoacylation.

The linkage made here between sC/sAC conservation and the contemporary non-AC aminoacylation can be validated or refuted experimentally. Due to the high occurrence of the conserved sC/sACs in the acceptor stems of the robustly conserving aa, the more efficient aminoacylation rates reported in the literature correspond to minihelices carrying a conserved coding triplet. Decrease in aminoacylation rate following the mutation of a nondeterminant nucleotide in a robustly conserved coding triplet can corroborate this linkage.

At the other half of the pre-3' end string, nucleotides 63–67 from the acceptor-TΨC stem establish the contact interface with EF-Tu [3]. The sC/sAC conservation sites in that range, that is, the Asp anticodon GUC in positions 64–66, which is conserved in 91% of the sequences, and the His codon CAU in positions 63–65, which is conserved in 60% of the sequences (Fig. 1), are suggested to contribute to the binding affinity for EF-Tu. In particular, EF-Tu was shown to use the tRNA<sup>Asp</sup> base pairs G51:C63, C50:G64, and G49:U65 for tightly binding it [30], where G64 and U65 are included in the conserved Asp codon GUC. For His, only C63 and U65 are completely conserved, probably reflecting the linkage between the present involvement of specific nucleotides in the recognition of tRNA by EF-Tu and their degree of conservation.

The sC/sACs conserved in pre-3' end strings of the nonrobustly conserving aa Arg, Leu, and Val, which are presently charged by class I synthetases, seem to lack a contemporary role. The conserved sC/sACs of Leu and Val are dispersed throughout the pre-3' end string (Fig. 1) rather than holding specific sites, as would be expected from functional sC/sACs. Consistently, the identity determinants for aminoacylating minimal analogs of tRNA<sup>Leu</sup> are found in the D-arm and in A73, but not in the pre-3' end [31]. 60% of the pre-3' end sequences from tRNA<sup>Arg</sup> actually contain a robustly conserved codon in positions 66–68 (Fig. 1). However, this site is unlikely to serve as an identity determinant, because its location partially overlaps the ranges of tRNA interaction with EF-Tu and with the synthetase. In accord, the base of A20 in the D loop is

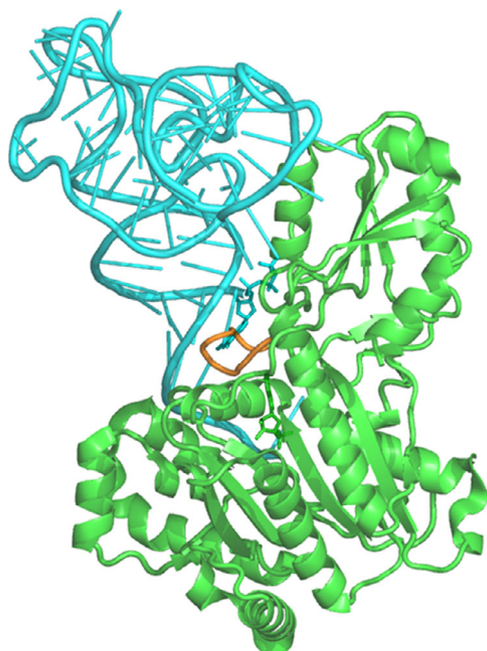
the dominant identity element for non-AC aminoacylation [32].

As expected from the assumed lack of contemporary role, the level of sC/sAC conservation in the pre-3' end strings of these three aa is lower compared with the six robustly conserving aa (Table 2). Also, the number of conserved sC/sACs per aa is larger: five, six, and eight for Leu, Arg, and Val, compared with one to three for the robustly conserving aa. Moreover, the robustly conserved sC/sAC sets, which comprise just codon or just anticodons, are evolutionary more plausible, because they can be the descendants of a single cognate coding triplet that existed in the acceptor-TΨC stem of the ancestral tRNA, while those containing both codons and anticodons, as is the case with Leu, Arg, and Val, cannot. In accord, the 'operational RNA codes' inferred from the bacterial classification and regression tree demonstrated the highest degree of degeneracy for Arg, Leu, and Val, indicating the reduced conservation of their acceptor stem identity elements [23]. These statistical observations seem to point to a gradual process by which, in the absence of a contemporary role, the conserved sC/sACs of Leu, Arg, and Val were shifted throughout the acceptor-TΨC stem, while random mutations supplemented the list of conserved sC/sACs and lowered the overall level of conservation.

Taken together, these five aa, that is, Ala, Gly, His, Pro, and Ser, share three different acceptor stem-related features: robust conservation of the sC/sAC, enhanced efficiency of minimal substrate aminoacylation, and being charged by class IIa synthetases. This correlation suggests the existence of a variant of the aminoacylation mode of action applied by the corresponding class IIa synthetases, which takes advantage of the information embedded in the robustly conserved sC/sACs to discriminate between tRNAs. Such a mechanism can provide reasoning for the enhanced efficiency of charging these minihelices and more importantly—for the extreme conservation (Table 2 and S2) of the coding triplets in the modern acceptor stems of these aa.

### Possible evolutionary implication of the conserved coding triplet

The level of sC/sAC occurrence in the acceptor-TΨC arm is highly correlated with the antiquity of the corresponding aa. Eight out of the nine conserving aa (Table 2), that is, all except His, are listed among the 10 earliest appearing aa, according to a consensus chronology built on 60 criteria [33], which is compatible with the results of the Miller–Urey experiment [34]. In particular,



**Fig. 2.** Synthetase interaction with the acceptor stem. tRNA<sup>His</sup>: HisRS complex (pdb 4rdx). Motif 2 loop (orange) of HisRS (green) bulges from the catalytic core into the major groove of tRNA<sup>His</sup> (cyan) acceptor stem.

the robustly conserving aa Ala, Asp, Gly, Pro, and Ser are among the six most ancient aa. The probability of such a correlation being obtained at random is 0.002, suggesting that the origin of the coding triplets, conserved in the acceptor-TΨC stem, is prebiotic [22].

The evolutionary history of tRNA recognition has gained considerable interest over the years, due to its linkage with the origin of the genetic code, eliciting a plethora of hypotheses. The capability of modern synthetases to utilize information held in the first base pairs of the acceptor stem for specific aminoacylation gave rise to the conjecture that this mode of operation is ancestral. The limited size of the primordial synthetase permitted it to reach only recognition elements residing in the acceptor stem, in the vicinity of the aa attachment site [35,36]. This simple proto-synthetase, which is believed to correspond to the modern catalytic domain [12,14], was suggested to have activated aa and to specifically recognize the tRNA *via* motif 2 loop (Fig. 2), charging in this manner early tRNAs based on their acceptor stem identity determinants [37]. Rodin *et al.* [38] further proposed that the operational RNA code as we see it today, developed from a few ancient anticodon–codon-like pairs located in the first positions of the acceptor stem, and that unrecognized relationship between the acceptor and anticodon points to the historically common root of the two codes.

In line with these hypotheses, we suggest that the primordial aminoacylation of proto-tRNAs by their cognate class IIa synthetases was controlled by the full coding triplets, which are still robustly conserved in the acceptor stem, rather than merely by the discrete identity determinants. The existence of a feasible non-AC aminoacylation mechanism that relies on the interaction of motif 2 loop with the sC/sACs conserved in the acceptor stems of Ala, Gly, His, Pro, and Ser points to ancestry of class IIa synthetases, in accord with previous proposals [39,40]. The robustly conserved coding triplets, suggested here to control the prebiotic aminoacylation, form a subset of the standard genetic code. They incorporate all but one of the cognate identity determinants that constitute the ‘second genetic code’ (Table 3), suggesting that the ‘second code’ [12,17], which was proposed to be distinct from the standard genetic code [16], actually makes part of it. In accord with the conjecture concerned with the common root of the acceptor and anticodon codes [38], the codons and anticodons conserved in the acceptor-TΨC stem are suggested here to have had a primal role in establishing the extant genetic code. A paper focusing on the origin of the conserved coding triplets and their implications on the scenario suggested for the origin of life is now under work (Agmon I. in prep).

The three key components of the protein synthesis machinery, that is, the tRNA, synthetase, and EF-Tu, along with the ribosome, have coevolved into the intricate contemporary translation system, still keeping traces of their prebiotic roots in the form of the conserved coding triplets. The evolutionary path proposed by the coding triplets conserved in the tRNA acceptor stem, commencing from specificity control in a primordial aminoacylation, to the involvement in the specificity recognition by the contemporary synthetase, outlines a continuous route from the prebiotic era to modern biology.

## Materials and methods

### Data set construction

Sequences of the 3' side of the acceptor-TΨC stem from the bacterial tRNAs of each of the 20 canonical aa were obtained from the tRNAdb database [19,20], <http://trna.bioinf.uni-leipzig.de>, 2018. The tRNAdb database holds over 6200 bacterial tRNA genes, belonging to the 20 canonical aa. Only tRNA sequences having a full CCA 3' end and a consensus discriminator base (N73) [10] were incorporated into the statistics, with one exception; the data for Glu contained similar number of sequences with the canonical G73

**Table 3.** Identity determinants vs. robustly conserved sC/sACs in the tRNA acceptor stem.

Amino acid	Ala	Gly	Ser	Ser	His	Pro
sC/sAC conservation sites in the 68–72 range						
Positions	68–70	68–70	68–70	70–72	70–72	70–72
Conserved sC/sACs <sup>a</sup>	GCU	CCC/UCC	UCU/UCA	UCC	CAC	CCG
Identity determinants for minimal substrate aminoacylation <sup>a</sup>	G3: <b>U70</b> [11]	G3: <b>C70</b> , C2:G71 <sup>c</sup> [41]	(C/U) <b>68</b> , (C/U) <b>69</b> , (A/U) <b>70</b> [15]	(A/U) <b>70</b> , <b>C71</b> , <b>C72</b> [15]	<b>C70</b> , <b>A71</b> [41]	<b>C70</b> <sup>b</sup> , <b>G72</b> <sup>b</sup> [24]

<sup>a</sup>In bold—nucleotides from the robustly conserved coding triplets serving as identity determinants.; <sup>b</sup>These identity determinants refer to the recognition of the full-length tRNA.; <sup>c</sup>The identity determinant for aminoacylating minihelices of Gly, C2:G71, is not part of the conserved coding triplets.

and with the noncanonical A73, both groups displaying the same pattern of sC/sAC occurrence. To obtain as much data as possible, tRNAs with noncanonical A73 were incorporated as well. After the exclusion of the partial sequences and of the rest of the noncanonical data, the initial data set used in the current statistical analysis contained 60% of the bacterial tRNA genes from the database.

The observed occurrence of the cognate codon and anticodon triplets was determined in 10-mer strings consisted of nucleotides 63–72 (termed pre-3' end), by scanning the sequences. Nucleotides 61 and 62 were not included in the study, being conserved and semi-conserved, respectively. The scanning indicated that one class of bacteria, that is, species from six genera of the *Mollicutes* class, which constituted about 9% of the initial data set (350 gene sequences), does not share the statistical behavior of the other bacteria (Table 1, File S1, Table S2), and it was excluded. No other species in the data contained enough entries to be reliably recognized as diverging from the conservation pattern. Excluding the *Mollicutes* resulted in a final data set consisting of 3324 tRNA gene sequences from over 100 bacterial species.

## Definitions

### Conserving amino acid

Amino acids for which more than 85% of their pre-3' end strings contain cognate sC/sACs are denoted 'conserving aa'. Amino acids for which < 40% of their pre-3' end strings contain cognate sC/sACs are denoted 'nonconserving aa'.

### Conserved sC/sAC set

For the conserving aa, it is the minimal set of cognate triplets accounting for 98% of the sequences carrying a cognate coding triplet.

### Robustly conserved sC/sACs

Are conserved sC/sACs that are located at specific sites on the pre-3' end strings, and occur, alone or with a single additional coding triplet (which is also considered as robustly conserved), in > 60% of the sequences.

### Robustly conserved sC/sAC set

For a specific aa, it is the set containing all its robustly conserved sC/sACs, conditional on the requirement that the sC/sAC set is composed of either just codons or just anticodons.

### Robustly conserving amino acids

They are those having a robustly conserved sC/sAC set.

### Calculation of expected occurrences

The expected occurrence of cognate coding triplets in the pre-3' end string was determined by analyzing their occurrence in simulated 10-mer randomized oligonucleotides. A pool of 10<sup>5</sup> randomized strings was generated using MATLAB and Statistics Toolbox (Release 2012b; The MathWorks, Inc., Natick, MA, USA), by producing a string of 10<sup>6</sup> nucleotides with a predetermined nucleotide distribution, randomizing it, and cutting it into 10<sup>5</sup> 10-mer oligonucleotides. For the conserving aa, the percentage of the 10-mer randomized strings carrying at least one cognate triplet from the conserved triplet set is defined as the expected occurrence. For the remaining aa, for which a conserved set of sC/sACs does not exist, the percentage of the 10-mer randomized strings carrying at least one cognate codon or anticodon triplet was set to be the expected occurrence. The error value for the expected percentage, ±0.8%, includes a statistical error of 0.6% and a rounding error of 0.5%. The probability of random occurrence, *P*, was calculated for each aa, from the results of a chi-square test, using a statistical cutoff of *P* ≤ 0.01.

### Benchmark test for robust conservation

The test counts the occurrences of all (coding and noncoding) triplets according to their appearance per 3-mer site within the 10-mer pre-3' end strings. Any cognate coding triplet that appears alone at a specific location, in more than half of the strings of an aa, is assumed to denote a highly conserving aa.

## Calculation of correlation

The probability of obtaining exactly  $m$  common objects in two groups of  $n_1$  and  $n_2$  objects chosen randomly out of a batch of  $n$  objects, is given by

$$\frac{\binom{n_1}{m} \cdot \binom{n-n_1}{n_2-m}}{\binom{n}{n_2}}$$

where

$$\binom{n}{m} = \frac{n!}{m!(n-m)!}$$

## Conclusions

The extreme conservation of cognate coding triplets in the acceptor stems that belong to five amino acids having tRNAs charged by class IIa synthetases points to their involvement in an extant function. A contemporary aminoacylation mode, whereby the coding triplets conserved in these tRNA acceptor stems, or in mini-helices derived from them, assist the recognition by their cognate synthetases, is likely to account for it. The antiquity of the corresponding amino acids suggests that this mode of action is a vestige of a simple ancestral process of specifically aminoacylating proto-tRNAs *via* the acceptor stem coding triplets. This, in turn, suggests the ancestry of class IIa synthetases, the incorporation of the ‘second genetic code’ within the standard code, and the involvement of the stem coding triplets in the establishment of the standard genetic code.

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## Author contributions

IA conceptualized the study and wrote the original manuscript. IA, IF, and TM designed methodology, and analyzed the data. IA and IF provided software. IF and TM revised the manuscript.

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## Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** Benchmark - CCC occurrence for amino acids with large excess of nucleotide C in the sequences of their tRNA pre-3'end.

**File S1.** Bacteria-occurrence of coding triplet in the pre-3'end string.

**File S2.** Archaea- occurrence of coding triplets in the pre-3'end string.

**File S3.** Eukarya- occurrence of coding triplets in the pre-3'end string.

**Table S1.** (a) Occurrence statistics of stem codons and anticodons in the tRNA pre-3'end; random, sample and individual distributions of bacterial data of conserving amino acids<sup>†</sup>. (b) Occurrence statistics of stem codons and anticodons, in the tRNA pre-3'end;

random, sample and individual distributions of bacterial data of the non-conserving amino acids.

**Table S2.** Occurrence of conserved stem codons and anticodons in the tRNA pre-3'end of different bacterial genera.