Comparisons between small ribosomal RNA and theoretical minimal RNA ring secondary structures confirm phylogenetic and structural accretion histories

Jacques Demongeot1✉ & Hervé Seligmann1,2

Ribosomal RNAs are complex structures that presumably evolved by tRNA accretions. Statistical properties of tRNA secondary structures correlate with genetic code integration orders of their cognate amino acids. Ribosomal RNA secondary structures resemble those of tRNAs with recent cognates. Hence, rRNAs presumably evolved from ancestral tRNAs. Here, analyses compare secondary structure subcomponents of small ribosomal RNA subunits with secondary structures of theoretical minimal RNA rings, presumed proto-tRNAs. Two independent methods determined different accretion orders of rRNA structural subelements: (a) classical comparative homology and phylogenetic reconstruction, and (b) a structural hypothesis assuming an inverted onion ring growth where the three-dimensional ribosome’s core is most ancient and peripheral elements most recent. Comparisons between (a) and (b) accretions orders with RNA ring secondary structure scales show that recent rRNA subelements are: 1. more like RNA rings with recent cognates, indicating ongoing coevolution between tRNA and rRNA secondary structures; 2. less similar to theoretical minimal RNA rings with ancient cognates. Our method fits (a) and (b) in all examined organisms, more with (a) than (b). Results stress the need to integrate independent methods. Theoretical minimal RNA rings are potential evolutionary references for any sequence-based evolutionary analyses, independent of the focal data from that study.

Ribosomes presumably evolved through serial accretions of tRNAs and tRNA-like RNAs1–11. The ribosomal dimeric RNA core surrounding the peptide synthesis site12–14 also resembles tRNA dimers linked by complementary anticodons, according to the self-referential hypothesis on the origin of translation15–19. Evidence for this process exists also in modern vertebrate mitochondrial ribosomes: regular mitochondrial tRNAs constitutively fulfill SS rRNA functions20,21. In the latter case, extreme mitogenome reduction perhaps reversed evolution to a tRNA-insertion stage, enabling further mitogenome reduction. These evidences suggest that rRNAs derived from tRNAs.

tRNA accretion

Several hypotheses suggest different historical scenarios for tRNA evolution, all assuming accretions of smaller sequences22–41. A similar hypothesis exists for SS rRNAs42. Some evidence suggests that tRNAs originate from stem-loop hairpins initiating replication43–50. Other analyses show striking similarities in nucleotide triplet biases of tRNAs and protein coding genes51,52. Theoretical RNA rings, sequences artificially designed according to coding constraints53,54 seem homologous to tRNAs55–57.

1Université Grenoble Alpes, Faculty of Medicine, Laboratory AGEIS EA 7407, Team Tools for e-Gnosis Medical & Labcom CNRS/UGA/OrangeLabs Telecoms4Health, F-38700, La Tronche, France. 2The National Natural History Collections, The Hebrew University of Jerusalem, 91404, Jerusalem, Israel. ✉e-mail: Jacques.Demongeot@univ-grenoble-alpes.fr
Two main approaches have been developed and used to recover accretion histories of ribosomal RNAs. Both consider secondary structure subcomponents of rRNAs as units undergoing this process. One approach is based on homology, character polarity and cladistic comparisons to infer accretion history from comparisons among numerous sequences. This classical comparative biology method uses parsimony as its main conceptual tool and was also used to recover evolution of molecular functions and protein accretion. Various empirical tests show that this method recovers actual histories better than chance.

A second approach is structure-based, and assumes that the ribosome grew from its spatial core towards its periphery, with the most ancient structural subcomponents located at the physical center of the ribosome, and the more recent ones at its periphery. The method corresponds to that of spatial comparisons in disciplines such as plant community ecology. Structures encompass large amounts of information: in ribosomes, contact biases between amino acids and nucleotide triplets recover the very ancient evolution of genetic code codon-amino acid assignments. Though reasonable, the structural method lacks to our knowledge further empirical tests in contexts of reconstructing biomolecular histories, but one of its merits is that for each taxon for which accurate structural data are available, it produces (slightly) different histories, enabling to search for consensuses.

The theoretical premises of the structural approach are in observations that ontogenies of different structures recover their phylogenies: chemical prebiotic evolution; genetic code evolution; embryology; and ecological communities. Spatial variation in vegetation can reconstruct the ontogeny of forests (forest succession), but plant colonization at forest periphery and clearings differ from de novo colonization of areas where no forest is adjacent and no humus exists: primary and secondary successions differ. In addition, the structural model unrealistically assumes equal ribosomal growth in all directions from the core to the periphery. Its name, the onion peeling model, is formally incorrect (in onions, peripheral rings are most ancient), reflecting emphasis on structure rather than historical process.

Comparing accretion histories: cladistic vs structure. Overall, one can assume that both approaches complete each other, one recovering history using phylogenetic methods, and the other using principles from ecology and embryology for historical reconstruction. Accretion ranks of the 16S rRNA secondary structure subcomponents according to cladistic- and structure-based methods differ (Fig. 1). This analysis shows some
congruence between accretion ranks obtained by the two independent methods, for 26 among 44 secondary structure elements (59%), which is not significantly more than 50% according to a one tailed sign test. The highest percentage of secondary structure subelements with reasonable match between accretion ranks from the two methods is for 16S rRNA domain 3, the lowest percentage is for domain 2. Notably, domain 4, presumed most ancient and consisting of two secondary structure subelements, has one element where both methods are highly congruent, and have very different ranks for the other subelement.

Secondary structure classification
The overall impression resulting from Fig. 1 is that both structural and phylogenetic methods have some level of congruence, for a bit more than half of the secondary structure subelements, across all four 16S rRNA structural domains. Hence, for almost half of the secondary structure subelements, we do not know the accretion rank. A third independent method for estimating RNA history could improve the resolution of rRNA accretion ranks.

A method clustering RNA secondary structures found two main RNA secondary structure groups, one characterized by small, presumably ancient tRNA-like secondary structures, and a presumed more derived group, characterized by larger, rRNA-like secondary structures, including viruses91,92. The tRNA-like cluster was...
designated as tRNA-like because it included tRNAs. The decision to assume it is most ancient was not only based on the inclusion of tRNAs in that cluster. This cluster includes a high diversity of RNA types (viroids, ribozymes, tRNAs, replication origins, 5S rRNAs). Ancient groups tend to be more diverse because more time is available for "evolutionary radiation" (this term from species evolution might not be adequate in context of RNA species). The same rationale was applied to functional tRNA species, ranking as most ancient those with the highest diversity of isoacceptor tRNAs.

The decision to consider the other RNA cluster as rRNA-like was because this cluster included all subdomains of small and large rRNAs. Note that this clustering is phenotypic, based on secondary structure similarities, not phylogenetic. The assumption that tRNA-like structures are primitive, and that rRNA-like structures are more derived is in line with the tRNA-accretion hypothesis for rRNA formation. Results show that tRNA-like RNAs have few unpaired nucleotides within stems (bulges); for rRNA-like secondary structures, the proportion of bulges among all unpaired nucleotides is greater. Bulges are targets for regulation and enzymatic degradation, properties of advanced metabolism. In prebiotic conditions, these might be disadvantageous, increasing degradation risks.

**Polarity of the tRNA-rRNA axis of RNA secondary structure evolution**

This assumption about the evolutionary direction of secondary structures was tested explicitly on tRNAs from diverse organisms (organelles, Archaea, Bacteria, Eukaryota and Megavirales). First, similarities of all tRNAs from specific organisms with tRNA-like vs rRNA-like groups were estimated, projecting each tRNA secondary structure on a presumed tRNA-rRNA axis of RNA secondary structure evolution. Then correlations were calculated between the genetic code inclusion rank of the tRNA cognate amino acids and this tRNA-rRNA similarity score, expecting that tRNAs with relatively recent cognates have more rRNA-like secondary structures, and those with ancient cognates, are more typically tRNA-like. Results were overall positive (weakest in Eukaryota), confirming tRNA-rRNA polarity: two independent scales of evolutionary ranks, one for amino acids, and one for RNA secondary structures, converge.

Here again, polarity is not deduced from phylogenetic reconstructions, but from presumed orders of integration of the tRNAs cognate amino acid.

Note that the phylogenetic and the structural methods also make polarity assumptions. In the former, these are deduced from cladistic parsimony principles, in the latter, from structure: the more peripheral a structural element in the ribosome, the more recent, including information on stacking interactions among subdomains. These results strengthen the hypothesis that tRNAs are ancestral and rRNAs derived.

**Independent references for RNA evolution**

The tRNA-rRNA evolutionary axis score is based on a sample of known RNA secondary structures. Hence, it suffers from sampling biases, and from some level of circularity: biological data are used to infer on biological phenomena, a caveat it shares with the phylogenetic method. A possible solution to this is to use as reference theoretical minimal RNA rings, a set of short sequences designed *in silico* according to few basic constraints: the shortest possible sequence coding for a start and a stop codon, and once for each of the 20 biogenic amino acids.

These constraints define at most 25 circular RNA sequences of 22 nucleotides, which code according to partially overlapping codons, along three consecutive translation rounds, for a start codon, 20 different amino acids, and a stop codon. The stop codon is physically next to the start codon, closing the RNA ring. These RNA rings, mainly defined by coding sequences, resemble ancestral tRNAs, with a predicted anticodon and its corresponding cognate amino acid for each RNA ring.

The theoretical minimal RNA rings realistically mimic primitive RNAs and their evolution, along several coding properties and primary and secondary structure properties. These properties coevolve with the genetic code integration order of the cognate amino acid matching the anticodon defined by homology of the RNA rings with ancestral tRNAs. Considering that the design of RNA rings is purely rational and mainly based on the structure of the genetic code, this means that the genetic code's structure intrinsically embeds information on the evolution of these various properties. However, we do not yet understand what determines these complex evolutionary trajectories.

Notably, the tRNA-rRNA scores obtained for secondary structures of these RNA rings, correlate, as observed for real tRNAs, with the evolutionary ranks of integration of the cognate amino acids matching their predicted anticodons. This parallels the result described in the previous section for regular tRNAs and the genetic code integration order of their cognate amino acid. Here too, the polarity results from this order, not from phylogenetic reconstruction.

**Working hypothesis and predictions**

Hence, RNA rings are designed as proto-tRNAs but have also properties that are expected for proto-tRNAs. As plausible proto-tRNAs, they are used here as references for ancestral RNAs, in line with results of evolutionary analyses of their different properties. Analyses use similarities between RNA ring secondary structures and those of structural subelements of 16S rRNAs. The method assumes that high similarities with RNA ring secondary structures indicate ancient structural subelements, and low similarities recent 16S rRNA structural subelements. These similarities are then compared with accretion ranks produced by each of the phylogenetic and the structural hypotheses, expecting: 1. negative correlations if the different methods are producing congruent accretion ranks; 2. these correlations should be most negative for RNA rings with ancient cognate amino acids, and gradually be more positive for RNA rings with recent cognate amino acids.

**Materials and methods**

The quantification of similarities between secondary structures is identical to previous analyses. Analyses use similarities between RNA ring secondary structures and those of structural subelements of 16S rRNAs. The method assumes that high similarities with RNA ring secondary structures indicate ancient structural subelements, and low similarities recent 16S rRNA structural subelements. These similarities are then compared with accretion ranks produced by each of the phylogenetic and the structural hypotheses, expecting: 1. negative correlations if the different methods are producing congruent accretion ranks; 2. these correlations should be most negative for RNA rings with ancient cognate amino acids, and gradually be more positive for RNA rings with recent cognate amino acids.
Four secondary structure properties are extracted from secondary structures, as shown as example for structural subelement h45 from the archaean Thermus thermophilus 16S rRNA (Fig. 2): 1. the percentage of nucleotides in stems formed by complementary self-hybridization among nucleotides, %stem among all nucleotides in the sequence; 2. the percentage of nucleotides, among those in loops, that are in loops topping stems (external loops), as opposed to unpaired nucleotides forming bulges within stems (internal loops), %eloops; and the 3. stem and 4. loop GC contents, in percentages.

Similarities between two secondary structure pairs are estimated by Pearson correlation coefficients r between these four variables as obtained for each secondary structure (Fig. 3), in this case between values from Fig. 2 and those of secondary structures formed by two alternative splicings of RNA ring 25, also called AB 53. Table 1 presents the four secondary structure variables for AB for all 22 alternative splicings of that RNA ring. Such data were obtained for all 25 RNA rings. Similar secondary structure data for 22 alternative splicings of RNA ring 13, called AL, were presented previously, and for each comparison, Fig. 3 has four datapoints for each secondary structure variable. For each datapoint, the X-axis is defined by the value obtained for the AB secondary structure, and the Y-axis by the value obtained for the corresponding variable for the 16S secondary structure subelement shown in Fig. 2. These pairings are not arbitrary: the x- and y-axis values are for the same secondary structure property, but for a different secondary structure (x-axis, RNA ring 25; y-axis, RNA structural subelement, in this case h45 of Thermus thermophilus). Similarities are estimated by r, the more positive r, the more similar the secondary structures.

The secondary structure variables of all secondary structure subelements of two Archaea, Thermus thermophilus and Sulfolobus solfataricus, two bacteria, Escherichia coli and Streptomyces coelicolor, and the 18S rRNA of two eukaryotes, Homo sapiens and Saccharomyces cerevisiae, are available at http://apollo.chemistry.gatech.edu/RibosomeGallery/.

Step by step description of analyses.

1. There are 25 RNA rings, each 22 nucleotide long. These are considered according to the splicing matching homology with ancestral tRNAs, as shown previously.
2. Each RNA ring can be spliced at 22 positions, and a different optimal secondary structure (predicted by Mfold) exists for RNA ring sequences spliced at each potential splicing position. The 25 RNA rings form 25 × 22 = 550 secondary structures.
3. Four secondary structure variables are extracted from each of these 550 secondary structures. Table 1 presents as an example these four variables for the 22 alternative splicings of a specific RNA ring, RNA ring 25.
4. For each of the (about 45) structural subelements of small rRNA subunits of the 6 examined organisms, the four secondary structure variables are extracted, as was done for the 550 RNA ring structures at step 3. These variables are presented for the 6 × 45 = 270 secondary structure subelements presented in Tables 2–4.
5. The secondary structures of RNA rings are compared to the secondary structures of rRNA structural subelements by analyses as presented in Fig. 3. These analyses plot the values obtained for each of the 4
secondary structure variables of a rRNA structural subelement as a function of the corresponding values obtained for a given RNA ring secondary structure. A Pearson correlation coefficient r, called rS, estimates similarities between rRNA and RNA ring secondary structures. Figure 3 presents comparisons between 16S rRNA subelement h45 of *Thermus thermophilus* and two RNA ring 25 secondary structures, one obtained by splicing that ring at position 7, and one at position 19.

For each of the 550 RNA ring secondary structures, there are as many rS as there are rRNA secondary structure subelements, about 45.

According to our hypothesis, the (about) 45 rSs comparing a given RNA ring secondary structure to all rRNA secondary structure subelements are potential estimates of the accretion order of the rRNA secondary structures.

These rSs are compared to the accretion order of the rRNA secondary structure subelements, as these were determined by other methods and published by other authors (separately for each cladistic and structural accretion ranks). This comparison is done by calculating the Pearson correlation coefficient between the rS and the accretion orders, producing rH, one for the cladistic method, rHphyl, and one for the structural method, rHstru. Note that rS are z-transformed before calculating rH using the formula $z = -\ln((1 + r)/(1 - r))$. The z transformation linearizes the scale of r, which is not linear.

Hence, each of the 550 RNA ring secondary structures produces one rHphyl and one rHstru per organism. For each organism, there are 550 rHphyls and 550 rHstrus. The minimal and maximal rHphyls and rHstrus for each organism are in Table 5. Table 5 includes percentages of negative rHphyls and rHstrus (the working hypothesis expects negative rHs), and numbers of negative and positive rHphyls and rHstrus that have two tailed $P < 0.05$.

For any given RNA ring secondary structure, there are 6 rHphyls and 6 rHstrus, because analyses were done for 6 organisms. There are in total $6 \times 550 = 3300$ rHphyls and 3300 rHstrus. Further analyses describe general patterns within these data, according to RNA rings, and according to splicing positions.

For each RNA ring, there are 22 secondary structures which produce 22 rHphyls and 22 rHstrus per organism, hence $6 \times 22 = 132$ rHphyls and 132 rHstrus across all 6 organisms. An alternative way to explain this is: for each of the 25 RNA rings, there are $3300/22 = 150$ rHphyls and 150 rHstrus across all 6 organisms. Percentages of negative rHphyls and rHstrus for each RNA ring (calculated among the 132 rHphyls and among the 132 rHstrus, pooling all organisms) are used in the y axis of Fig. 4.

There are 25 RNA rings. Hence, for a given splicing position, there are 25 rHphyls and 25 rHstrus. Pooling these data across 6 organisms, for any given splicing position, there are $6 \times 25 = 150$ rHphyls and 150 rHstrus across all 6 organisms. Percentages of negative rHphyls and rHstrus for each splicing position, calculated from these 150 rHphyls and 150 rHstrus, consist the y axis in Fig. 5.

|    | %stem | %seloop | %GCstem | %GCloop |
|----|-------|---------|---------|---------|
| 1  | 54.5  | 40.0    | 33.3    | 50.0    |
| 2  | 54.5  | 40.0    | 33.3    | 50.0    |
| 3  | 54.5  | 40.0    | 33.3    | 50.0    |
| 4  | 54.5  | 40.0    | 33.3    | 50.0    |
| 5  | 54.5  | 40.0    | 33.3    | 50.0    |
| 6  | 54.5  | 40.0    | 33.3    | 50.0    |
| 7  | 45.5  | 33.3    | 40.0    | 41.7    |
| 8  | 36.4  | 28.6    | 25.0    | 50.0    |
| 9  | 36.4  | 28.6    | 37.5    | 42.9    |
| 10 | 36.4  | 28.6    | 37.5    | 42.9    |
| 11 | 45.5  | 50.0    | 20.0    | 58.3    |
| 12 | 54.5  | 60.0    | 33.3    | 50.0    |
| 13 | 63.6  | 75.0    | 42.9    | 37.5    |
| 14 | 72.7  | 100.0   | 43.8    | 33.3    |
| 15 | 63.6  | 75.0    | 42.9    | 37.5    |
| 16 | 54.5  | 60.0    | 33.3    | 50.0    |
| 17 | 45.5  | 50.0    | 20.0    | 58.3    |
| 18 | 36.4  | 42.9    | 25.0    | 50.0    |
| 19 | 45.5  | 83.3    | 30.0    | 50.0    |
| 20 | 45.5  | 83.3    | 30.0    | 13.3    |
| 21 | 45.5  | 83.3    | 40.0    | 41.7    |
| 22 | 54.5  | 40.0    | 33.3    | 50.0    |

Table 1. Secondary structure variables extracted as in Fig. 2 explanations, for the 22 secondary structures formed by RNA ring 25 (AB, TATGAATGGTGGCATTCAAGACTA)53, according to 22 splicing positions. Splicing at position 1 corresponds to the splicing position producing highest homology with an ancestral tRNA53, each splicing of the RNA ring is shifted by a single nucleotide. These secondary structure data were used previously57,109,110,111.
| #  | Phyl | Str | Thermus thermophilus | Sulfolobus solfataricus |
|----|------|-----|----------------------|------------------------|
|    |      |     | %ste #eloo %GCst %GClo | %ste #eloo %GCst %GClo |
| h1 | 25   | 3.3 | 43.5 30.8 60.0 30.8 | 19.0 0.0 50.0 58.8 |
| h2 | 29   | 4   | 75.0 0.0 133.3 0.0 | 66.7 0.0 50.0 0.0 |
| h3 | 10   | 4   | 62.5 0.0 60.0 50.0 | 71.4 0.0 75.0 25.0 |
| h4 | 7    | 94.1| 0.0 93.8 0.0 100.0 | 30.8 60.0 88.9 |
| h5 | 20   | 9   | 54.5 0.0 58.3 40.0 | 54.5 0.0 83.3 40.0 |
| h6 | 10   | 12  | 72.7 25.0 81.3 58.3 | 62.1 54.5 61.1 45.5 |

Table 2. Variables extracted from secondary structures of 16S rRNA of archaeans *Thermus thermophilus* and *Sulfolobus solfataricus*. Columns are: 1. secondary structure subelement of 16S rRNA; 2 and 3. accretion ranks according to phylogenetic and structural models, respectively59,78 and 4–7, secondary structure variables as in Table 1 (explained in Figs. 2 and 3). Domains range from 1. h1-h18; 2. h19-h27; 3. h28-h43; 4. h44-h45.
Analyses in Table 5, Figs. 4 and 5 each take into consideration all 3300 rHphyls and 3300 rHstrus. Hence, these are not biased representations of the data. They show separately effects of each ‘treatment factor’ (organism, RNA ring, splicing position) on each rHphyl and rHstru.

| #  | Escherichia coli | Streptomyces coelicolor |
|----|----------------|-------------------------|
|    | %ste | #eloo | %GCst | %GClo | %ste | #eloo | %GCst | %GClo |
| h1 | 63.6 | 62.5 | 42.9 | 25.0 | 48.9 | 33.3 | 52.2 | 41.7 |
| h2 | 72.7 | 50.0 | 43.8 | 33.3 | 60.0 | 0.0 | 66.7 | 0.0 |
| h3 | 54.5 | 50.0 | 50.0 | 30.0 | 69.0 | 0.0 | 60.0 | 44.4 |
| h4 | 45.5 | 25.0 | 30.0 | 42.9 | 88.9 | 0.0 | 87.5 | 50.0 |
| h5 | 72.7 | 50.0 | 43.8 | 16.7 | 66.7 | 0.0 | 66.7 | 16.7 |
| h6 | 63.6 | 37.5 | 35.7 | 37.5 | 61.1 | 28.6 | 59.1 | 50.0 |
| h6a| 54.5 | 30.0 | 41.7 | 30.0 | 0.0 | 0.0 | 66.7 | 0.0 |
| h7 | 63.6 | 62.5 | 42.9 | 37.5 | 69.6 | 0.0 | 71.9 | 28.6 |
| h8 | 45.5 | 41.7 | 30.0 | 50.0 | 62.5 | 33.3 | 65.0 | 58.3 |
| h9 | 72.7 | 50.0 | 43.8 | 16.7 | 62.9 | 30.8 | 68.2 | 30.8 |
| h10| 72.7 | 50.0 | 56.3 | 0.0 | 44.4 | 80.0 | 100.0 | 80.0 |
| h11| 54.5 | 50.0 | 58.3 | 20.0 | 60.0 | 35.0 | 60.0 | 45.0 |
| h12| 45.5 | 41.7 | 40.0 | 41.7 | 88.9 | 0.0 | 87.5 | 50.0 |
| h13| 63.6 | 37.5 | 35.7 | 37.5 | 61.1 | 28.6 | 59.1 | 50.0 |
| h14| 45.5 | 41.7 | 30.0 | 50.0 | 51.6 | 26.7 | 75.0 | 46.7 |
| h15| 72.7 | 50.0 | 31.3 | 66.7 | 62.9 | 30.8 | 68.2 | 30.8 |
| h16| 72.7 | 50.0 | 56.3 | 0.0 | 44.4 | 80.0 | 100.0 | 80.0 |
| h17| 54.5 | 50.0 | 41.7 | 30.0 | 85.7 | 0.0 | 75.0 | 50.0 |
| h18| 63.6 | 37.5 | 50.0 | 25.0 | 80.0 | 0.0 | 71.9 | 28.6 |
| h19| 63.6 | 30.0 | 35.7 | 37.5 | 57.1 | 33.3 | 68.8 | 33.3 |
| h20| 54.5 | 50.0 | 58.3 | 20.0 | 76.2 | 93.3 | 58.3 | 33.3 |
| h21| 63.6 | 37.5 | 50.0 | 25.0 | 66.7 | 0.0 | 71.4 | 35.0 |
| h22| 45.5 | 41.7 | 30.0 | 50.0 | 66.7 | 0.0 | 71.4 | 35.0 |
| h23| 72.7 | 50.0 | 43.8 | 16.7 | 70.8 | 26.3 | 52.2 | 31.6 |
| h23a| 54.5 | 40.0 | 33.3 | 50.0 | 85.7 | 0.0 | 75.0 | 50.0 |
| h24| 54.5 | 50.0 | 58.3 | 20.0 | 43.9 | 52.2 | 72.2 | 34.8 |
| h25| 63.6 | 37.5 | 50.0 | 25.0 | 52.9 | 25.0 | 66.7 | 37.5 |
| h26a| 63.6 | 62.5 | 42.9 | 37.5 | 26.7 | 54.5 | 66.7 | 37.5 |
| h26| 54.5 | 50.0 | 58.3 | 20.0 | 52.2 | 25.0 | 66.7 | 37.5 |
| h27| 63.6 | 30.0 | 35.7 | 37.5 | 57.1 | 33.3 | 68.8 | 33.3 |
| h28| 54.5 | 50.0 | 31.3 | 66.7 | 82.4 | 0.0 | 64.3 | 0.0 |
| h29| 63.6 | 37.5 | 50.0 | 12.5 | 57.1 | 33.3 | 68.8 | 33.3 |
| h30| 63.6 | 62.5 | 42.9 | 37.5 | 71.4 | 0.0 | 65.0 | 50.0 |
| h31| 54.5 | 50.0 | 41.7 | 40.0 | 28.6 | 40.0 | 50.0 | 35.0 |
| h32| 72.7 | 50.0 | 31.3 | 66.7 | 82.4 | 0.0 | 64.3 | 0.0 |
| h33| 72.7 | 50.0 | 56.3 | 0.0 | 69.6 | 0.0 | 56.3 | 42.9 |
| h33a| 63.6 | 62.5 | 50.0 | 25.0 | 53.3 | 64.3 | 68.8 | 42.9 |
| h34| 54.5 | 50.0 | 50.0 | 30.0 | 75.6 | 0.0 | 67.6 | 27.3 |
| h35| 72.7 | 50.0 | 43.8 | 33.3 | 75.0 | 0.0 | 66.7 | 50.0 |
| h36| 72.7 | 50.0 | 56.3 | 0.0 | 54.5 | 80.0 | 50.0 | 60.0 |
| h37| 72.7 | 50.0 | 37.5 | 50.0 | 50.0 | 75.0 | 87.5 | 25.0 |
| h38| 27.3 | 18.8 | 83.3 | 25.0 | 80.0 | 0.0 | 70.8 | 33.3 |
| h39| 72.7 | 50.0 | 43.8 | 16.7 | 61.5 | 50.0 | 68.8 | 30.0 |
| h40| 81.8 | 75.0 | 38.9 | 25.0 | 42.9 | 25.0 | 75.0 | 56.3 |
| h41| 45.5 | 25.0 | 20.0 | 50.0 | 63.6 | 0.0 | 85.7 | 50.0 |
| h41a| 54.5 | 30.0 | 58.3 | 10.0 | 50.0 | 25.0 | 75.0 | 45.8 |
| h42| 81.8 | 75.0 | 38.9 | 25.0 | 39.0 | 32.0 | 75.0 | 44.0 |
| h43| 45.5 | 25.0 | 40.0 | 41.7 | 37.8 | 34.8 | 64.3 | 43.5 |
| h44| 27.3 | 18.8 | 66.7 | 25.0 | 59.8 | 11.6 | 79.6 | 33.3 |
| h45| 54.5 | 40.0 | 33.3 | 50.0 | 50.0 | 20.0 | 70.8 | 33.3 |

Table 3. Variables extracted from secondary structures of 16S rRNA of bacteria *Escherichia coli* and *Streptomyces coelicolor*. Columns 2–9 correspond to columns 4–11 in Table 2.
Results and discussion

There are 25 theoretical minimal RNA rings. Each has exactly 22 nucleotides, hence each RNA ring has 22 alternative splicing positions. Different splicings produce different sequences forming different secondary structures.

| #  | %ste | #eloo | %GCst | %GClo | %ste | #eloo | %GCst | %GClo |
|----|------|-------|-------|-------|------|-------|-------|-------|
| Homo sapiens | Saccharomyces cerevisiae | Homo sapiens | Saccharomyces cerevisiae |
| h1 | 52.6 | 66.7 | 70.0 | 44.4 | 50.0 | 70.0 | 70.0 | 30.0 |
| h2 | 36.4 | 57.1 | 50.0 | 0.0 | 50.0 | 0.0 | 50.0 | 0.0 |
| h3 | 66.7 | 0.0 | 25.0 | 60.0 | 84.6 | 0.0 | 27.3 | 25.0 |
| h4 | 30.8 | 0.0 | 50.0 | 44.4 | 25.0 | 0.0 | 0.0 | 33.3 |
| h5 | 57.1 | 0.0 | 66.7 | 22.2 | 54.5 | 0.0 | 83.3 | 10.0 |
| h6 | 35.0 | 38.5 | 50.0 | 50.0 | 35.9 | 40.0 | 57.1 | 20.0 |
| h6a | 50.0 | 0.0 | 133.3 | 33.3 | 75.0 | 0.0 | 66.7 | 50.0 |
| h7 | 50.0 | 0.0 | 45.0 | 45.0 | 35.7 | 0.0 | 30.0 | 25.0 |
| h8 | 31.4 | 14.3 | 56.3 | 45.7 | 41.9 | 20.0 | 50.0 | 20.0 |
| h9 | 53.8 | 33.3 | 84.3 | 58.3 | 54.1 | 20.5 | 34.8 | 25.6 |
| h10 | 66.7 | 66.7 | 88.9 | 77.8 | 66.7 | 62.5 | 75.0 | 37.5 |
| h11 | 50.0 | 34.6 | 53.8 | 57.7 | 54.9 | 30.4 | 57.1 | 39.1 |
| h12 | 53.3 | 50.0 | 33.3 | 36.4 | 27.8 | 42.3 | 50.0 | 26.9 |
| h13 | 43.5 | 76.9 | 80.0 | 46.2 | 38.5 | 62.5 | 80.0 | 37.5 |
| h14 | 42.9 | 50.0 | 66.7 | 37.5 | 42.9 | 50.0 | 66.7 | 50.0 |
| h15 | 54.8 | 35.7 | 70.6 | 64.3 | 50.0 | 31.3 | 56.3 | 43.8 |
| h16 | 64.9 | 30.8 | 50.0 | 23.1 | 61.1 | 28.6 | 59.1 | 21.4 |
| h17 | 35.3 | 50.0 | 45.0 | 45.0 | 35.7 | 0.0 | 58.0 | 20.0 |
| h18 | 47.1 | 25.9 | 62.5 | 48.1 | 49.0 | 16.0 | 62.5 | 52.0 |
| h19 | 0.0 | 0.0 | 12.5 | 0.0 | 0.0 | 7.1 |
| h20 | 72.7 | 0.0 | 37.5 | 50.0 | 60.9 | 0.0 | 35.7 | 44.4 |
| h21 | 61.7 | 29.3 | 69.7 | 52.4 | 49.3 | 30.4 | 58.0 | 28.7 |
| h22 | 59.5 | 0.0 | 68.2 | 33.3 | 57.1 | 0.0 | 70.8 | 38.9 |
| h23 | 55.8 | 31.6 | 70.8 | 42.1 | 57.8 | 31.6 | 50.0 | 26.3 |
| h23a | 25.0 | 33.3 | 100.0 | 41.7 | 25.0 | 33.3 | 100.0 | 25.0 |
| h24 | 45.3 | 31.0 | 66.7 | 31.0 | 46.2 | 32.1 | 50.0 | 32.1 |
| h25 | 60.0 | 0.0 | 75.0 | 37.5 | 63.6 | 0.0 | 64.3 | 25.0 |
| h26a | 40.0 | 44.4 | 66.7 | 33.3 | 34.0 | 54.8 | 81.3 | 25.8 |
| h26 | 76.9 | 44.4 | 70.0 | 77.8 | 40.0 | 44.4 | 33.3 | 22.2 |
| h27 | 48.5 | 23.5 | 50.0 | 35.3 | 62.9 | 30.8 | 63.6 | 46.2 |
| h28 | 80.0 | 0.0 | 53.6 | 100.0 | 75.7 | 0.0 | 55.6 | 100.0 |
| h29 | 62.5 | 0.0 | 50.0 | 83.3 | 85.7 | 0.0 | 41.7 | 50.0 |
| h30 | 69.2 | 0.0 | 83.3 | 75.0 | 71.4 | 0.0 | 75.0 | 75.0 |
| h31 | 33.3 | 40.0 | 50.0 | 30.0 | 33.3 | 40.0 | 50.0 | 40.0 |
| h32 | 66.7 | 0.0 | 100.0 | 33.3 | 85.7 | 0.0 | 75.0 | 0.0 |
| h33 | 40.0 | 33.3 | 60.0 | 33.3 | 40.0 | 33.3 | 50.0 | 0.0 |
| h33a | 63.8 | 0.0 | 60.0 | 105.9 | 66.7 | 0.0 | 56.3 | 56.3 |
| h34 | 66.7 | 0.0 | 40.0 | 80.0 | 66.7 | 0.0 | 60.0 | 120.0 |
| h35 | 60.0 | 100.0 | 33.3 | 75.0 | 60.0 | 100.0 | 33.3 | 50.0 |
| h36 | 38.5 | 75.0 | 140.0 | 0.0 | 40.0 | 50.0 | 62.5 | 8.3 |
| h37 | 66.7 | 0.0 | 62.5 | 50.0 | 84.2 | 0.0 | 81.3 | 0.0 |
| h38 | 63.2 | 19.0 | 50.0 | 76.2 | 71.4 | 33.3 | 53.3 | 8.3 |
| h39 | 55.2 | 38.5 | 75.0 | 23.1 | 69.0 | 33.3 | 55.0 | 33.3 |
| h40 | 37.5 | 0.0 | 61.1 | 70.0 | 56.4 | 0.0 | 54.5 | 58.8 |
| h41 | 82.4 | 66.7 | 46.4 | 66.7 | 51.4 | 23.5 | 72.2 | 17.6 |
| h41a | 37.8 | 43.5 | 78.6 | 30.4 | 42.1 | 45.5 | 56.3 | 27.3 |
| h42 | 72.7 | 55.6 | 50.0 | 33.3 | 55.2 | 53.8 | 56.3 | 38.5 |
| h43 | 61.9 | 9.4 | 69.8 | 45.3 | 63.8 | 7.8 | 55.6 | 37.3 |
| h44 | 62.5 | 33.3 | 60.0 | 25.0 | 62.5 | 33.3 | 60.0 | 25.0 |

Table 4. Variables extracted from secondary structures of 16S rRNA of eukaryotes Homo sapiens and Saccharomyces cerevisiae. Columns 2–9 correspond to 4–11 in Table 2.
as shown for RNA ring 25, AB, in Table 1, and previously for RNA rings 9106, (therein Table 1) and 1357, (therein Table 3). Hence, there are 25 × 22 = 550 secondary structures to which secondary structure subelements of the 16S rRNAs can be compared.

The secondary structure variables shown in Table 1 and Figs. 2 and 3 were extracted for each of these 550 RNA ring secondary structures and are compared, as shown in Fig. 3, with the corresponding secondary structure variables of all secondary structure subelements of all six organisms considered here.

Table 5 shows the most negative and the most positive rH correlations between accretion ranks according to phylogenetic (rHphy1) and structural (rHstru) models with secondary structure similarities with RNA rings for 16S rRNAs of six organisms, and percentages of rHs (%neg) that are negative as expected by the working hypothesis among the 550 correlation calculated for each rHphy and rHstru, for each organism. * indicates statistically significant differences (P < 0.05) from 50% (550/2 = 275 negative rHphy and rHstru are expected if the sign of rH has an unbiased distribution between negative and positive trends) according to a chi-square test. “Co” indicates the cognate amino acid corresponding to the anticodon of the RNA ring(s) producing these correlations. Cognate G always corresponds to RNA ring 25 (AB). N indicates numbers of datapoints involved in the calculation of rH correlation coefficients.

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Percentage of negative Pearson correlation coefficients r between accretion ranks (phylogenetic method, filled symbols; structural method, hollow symbols) and similarities between 16S rRNA and RNA ring secondary structures, r’s pooled across organisms and secondary structures formed by the 22 alternative splicing of each RNA ring, as a function of the genetic code integration order of the RNA ring’s predicted cognate amino acid according to Davis’s hypothesis on N-fixing amino acids105. The working hypothesis expects negative r’s in particular for ancient amino acids.

### Table 5. Most negative and most positive Pearson correlations coefficients r (x100) (rH) between accretion ranks according to phylogenetic (rHphy) and structural (rHstru) models with secondary structure similarities with RNA rings for 16S rRNAs of six organisms, and percentages of rHs (%neg) that are negative as expected by the working hypothesis among the 550 correlation calculated for each rHphy and rHstru, for each organism.

| Taxon | Phyl. | rHphy | Stru. | rHstru |
|-------|-------|-------|-------|-------|
| | N | Min | Co | %neg | Max | Co | %neg | Max | Co |
| **Archaea** | | | | | | | | | |
| *Thermus thermophilus* | 38 | −46.1 | G | 62.6* | 44.2 | Sec | 48 | −40.7 | CDEM | 67.9* | 38.7 | G |
| *Sulfolobus solfataricus* | 44 | −54.7 | AKQSW | 86.9* | 37.4 | G | 46 | −50.9 | AGLNPQS | 54* | 51.2 | Sec |
| **Bacteria** | | | | | | | | | |
| *Escherichia coli* | 42 | −37.4 | G | 64.7* | 35.6 | Sec | 48 | −35.6 | ALNPQS | 64.2* | 35.0 | Sec |
| *Streptomyces coelicolor* | 39 | −46.8 | A | 90.9* | 30.7 | R | 48 | −32.9 | GLT | 72* | 32.3 | G |
| **Eukaryota** | | | | | | | | | |
| *Homo sapiens* | 43 | −36.0 | N, R | 53.5 | 32.3 | Sec | 48 | −25.5 | AKQSW | 80.4* | 19.1 | G |
| *Saccharomyces cerevisiae* | 43 | −36.1 | S | 79.1* | 27.5 | Sec | 48 | −36.0 | GT | 53.6 | 34.9 | Sec |
in yeast and in *Sulfolobus*. In addition, percentages of negative rHs are significantly greater for rHphyl than rHstru within three among six species, *Sulfolobus*, *Streptomyces* and yeast. In *Homo*, percentages of negative rHstru were significantly greater than percentages of negative rHphyl. The overall pattern is that results match the working hypothesis, and this more for rHphyl than rHstru. The opposite occurs in *Homo*. This could be interpreted as due to recent evolution of small rRNA structure in that species, but would require additional analyses and data from other species.

A second noteworthy point is that the most positive correlations are in 7 among 12 cases with RNA ring 2, which has a predicted anticodon for a stop codon, coding sometimes for selenocysteine. This is presumably one of the latest amino acids integrated in the genetic code (21st). This result fits the prediction that the most positive correlations between accretion ranks and secondary structure similarities would correspond to RNA rings with recent cognates. In other words, these secondary structures would not be references for initial RNAs starting the accretion process, but for the latest RNAs in the accretion process.

Figure 4 plots percentages of negative r’s between accretion ranks and secondary structure similarities between small rRNA subelements and RNA ring secondary structures, pooling all organisms and alternative splicings of RNA rings. Patterns confirm several points: 1. Most correlations between accretion ranks and secondary structure similarities are negative as expected by the working hypothesis, for most RNA rings; 2. In most cases, there are more negative correlations for the phylogenetic than the structural method for reconstructing accretion ranks; 3. Percentages of negative correlations decrease with the genetic code integration order of the cognate amino acid of RNA rings (see above comments for RNA ring with selenocysteine as predicted cognate).

Figure 5 presents the percentages of negative r’s between accretion ranks and secondary structure similarities between small rRNA subelements and RNA ring secondary structures, pooling all organisms and RNA rings, as a function of RNA ring splicing position. Results show that correlations are most frequently negative, meaning fitting the working hypothesis, when RNA rings are spliced at position “1”. This is the position defined by the highest homology between the RNA ring and an ancestral tRNA. This observation is also in line with the working hypothesis that RNA rings are proto-tRNAs, and that accretion of proto-tRNAs, tRNAs and tRNA-like RNAs formed rRNAs. Note that the assumption that RNA rings are proto-tRNAs is under debate. Nevertheless, and apparently confirming this status of proto-tRNAs, pseudo-phylogenetic analyses of RNA ring sequences reveal two clusters of RNA rings, one coinciding with RNA rings whose presumed cognate amino acid is the cognate of tRNAs for which the tRNA acceptor stem includes a primitive code.

Particularly noteworthy is that results of analyses presented here for the small rRNA subunit are in line with results obtained for the large rRNA subunit. These analyses compared structural subelements of the large rRNA subunit with the same RNA ring secondary structures as those used here. As described here for the small rRNA, for the large rRNA subunit, comparisons with RNA ring secondary structures show that: a. are slightly more congruent with the phylogenetic than the structural method; b. results are strongest for comparisons with RNA rings with predicted ancient cognate amino acids; c. weakest for comparisons with RNA rings with predicted recent cognate amino acids.

Conclusions
Results are strong corroboration of the working hypothesis that tRNA accretions formed rRNAs. They show that RNA rings are likely proto-tRNAs, and that these are good reference points for primitive RNAs in general, and tRNAs in particular. Results confirm that RNA ring cognates are good estimates for RNA ring evolutionary ranks, and that similarities between secondary structures bear information on evolutionary direction of RNA secondary
structures, from tRNA to rRNA-like, also among rRNA structural subelements. This has been suggested by several previous lines of analyses presented in the Introduction,[10,12,15,21,56,57,91,92,95,96,97,98] expanding upon evidences for common origins for tRNAs and rRNAs[1,2]. Analyses presented here for the small rRNA subunit show greater congruence between accretion orders derived from the secondary structure method used here and the phylogenetic method than between the former and the structural method. Similar analyses done for the large rRNA subunit produce qualitatively similar results, independently confirming our method and evolutionary conclusions. Overall, both phylogenetic and structural methods produce accretion orders that are congruent with the secondary structure method applied through the tRNA-rRNA axis of RNA secondary structure evolution. It is probable that the structural methods are more prone to errors due to evolutionary convergences than the phylogenetic method, though convergences remain the main difficulty in reconstructing evolution.

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Additional information
Correspondence and requests for materials should be addressed to J.D.

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