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The TYMV tRNA-like structure

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Summary — The genomic RNA from turnip yellow mosaic virus presents a 3'-end functionally and structurally related to tRNAs. This report summarizes our knowledge about the peculiar structure of the tRNA-like domain and its interaction with tRNA specific proteins, like RNase P, tRNA nucleotidyl-transferase, aminoacyl-tRNA synthetases, and elongation factors. It discusses also the biological role of this structure in the viral life cycle. A brief survey of our knowledge of other tRNA mimickers in biological systems, as well as their relevance for understanding canonical tRNA, will also be presented.

turnip yellow mosaic virus RNA / tRNA-like structure / aminoacylation / replication

Introduction

Turnip yellow mosaic virus, the type member of the tymoviruses and causative agent of systemic infections primarily in members of the Cruciferaea, was discovered in the late 40s [1] and since then has been studied extensively (reviewed by Hirth and Givord [2]). This spherical virus has a characteristic icosahedral capsid, formed by 180 copies of the coat protein monomer (20 kDa). The capsid contains one copy of a single stranded (+) sense genomic RNA. This RNA (6318 nucleotides) has been sequenced for two different strains, the type strain originally from Strasbourg [3, 4] and an Australian isolate [5]. It contains two overlapping 5'-proximal genes and a silent 3'-proximal coat protein gene. In infected host cells, a subgenomic RNA of 693 nucleotides is generated and is also encapsidated; its sequence codes for the coat protein. The two 5'-proximal genes are translated from the genomic RNA. A long open reading frame encodes a 206-kDa protein that undergoes specific cleavage resulting in two proteins of about 150 and 70 kDa [6, 7]. A second open reading frame, upstream of the previous one and out of frame, encodes a 69-kDa protein [8]. A cDNA clone from which infectious transcripts of the genomic RNA can be made in vitro has been constructed, and has opened the way to genetic analysis of the viral genome in connection with the viral life cycle [8].

TYMV was the first virus for which a very peculiar feature was discovered, namely the presence of a tRNA-like domain at the 3'-end of its RNA. The first experimental suggestions pointing to a functional tRNA mimicry were reported about 30 years ago when it was suggested that the viral RNA can bind amino acids, predominantly valine, when incubated with bacterial extracts and that this binding is pH dependent and lost at alkaline pH [9]. Firm demonstration that valine can be attached by a covalent bond to the 3'-extremity of the viral RNA by a reaction catalyzed by ValRS was presented by Pinck et al [10] and Yot et al [11]. The immediate consequence of this discovery was the expectation that the viral RNA contained a structure mimicking a tRNA. Soon afterwards, it was also discovered that this tRNA mimicry is not restricted to aminoacylation, but concerns interaction of the viral RNA with other tRNA-specific proteins like elongation factors [12, 13], tRNA nucleotidyl-transferase [14], and RNase P.
Fig 1. Folding of the 3'-end of TYMV RNA. A. Secondary structure of the last 110 nucleotides of the viral RNA, including the tRNA-like domain. Numbering starts at the 3'-end with negative numbers (for simplicity the minus signs are omitted). Plain lines highlight long range tertiary interactions, leading to the formation of pseudoknots. B. Secondary structure highlighting the pseudoknots and the general L-shaped conformation. C. Graphical three-dimensional computer model of the 86 3'-nucleotides [27]. In B, residue U21 from loop L1 of the pseudoknot is placed opposite to discriminator base A4.

[15]. The functional tRNA mimicry is however not complete, since TYMV RNA is not amino acid donor in protein synthesis [16].

In this paper we summarize the present view on the structure and function of the tRNA-like domain of TYMV RNA. We describe the mimicry of this structure for tRNA recognizing enzymes (RNAse P, tRNA nucleotidyl-transferase, aminoacyl-tRNA synthetases, and elongation factor) and discuss its role in the viral life cycle. We show in particular its involvement in the replication of the viral genome. Finally we summarize our knowledge on the tRNA mimicries in other plant viral RNAs and show that such mimicries are widespread events.

Structure of the 3'-non-coding region of TYMV RNA

Because TYMV RNA could be valylated by ValRS with kinetic characteristics similar to those of the charging of canonical tRNA [17], it could be anticipated that the viral RNA contains structural features resembling those of tRNAVal for the recognition and charging by the synthetase. The surprise was great that no obvious resemblance between a tRNA cloverleaf and the sequence of the 3'-end of the viral RNA was found [18, 19]. Thus the mimicry had to be searched for at the conformational level of the molecule. In what follows we describe the structural studies, mainly biochemical, leading to the present structural knowledge of the TYMV tRNA-like domain. It is worth emphasizing the importance these studies had for understanding RNA, because they led to the discovery of a new folding principle in these molecules, the pseudoknot, which is one of the major recent discoveries in the RNA field [20, 21]. Indeed, pseudoknots appear widespread in nature and are found to be involved in an increasing number of biological processes (reviewed by ten Dam et al [22] and Westhof and Jaeger [23]).

Secondary structure

RNA fragments of 112 and 159 nucleotides, corresponding to TYMV RNA have been purified from partial RNAse digestes of the viral RNA, and their secondary structure determined experimentally and independently by two groups using enzymatic and chemical probing techniques [24–26]. Both approaches lead to the same folding. The longest of these fragments, containing the end of the coat protein gene, was shown to be as efficient a substrate for ValRS as the complete viral RNA [24] suggesting that this fragment contains all the chemical information required for recognition by the aminoacyl-tRNA synthetase. Figure 1A represents the secondary structure of the non-coding 110 last nucleotides including the stop codon for the coat protein cistron at position −107 to −109.
The secondary structure of the 110 3'-terminal nucleotides of TYMV RNA is divided in five well-organized loop regions. Three of them (II, III, IV) exist in the other sequenced tymoviral tRNA-like domains and one tobamoviral valylatable RNA resembles classical tRNAs. This secondary structure of the TYMV tRNA-like domain is supported by phylogenetic data, since five other valylatable tymoviral tRNA-like domains and one tobamoviral valylatable RNA fold similarly over their last 80 nucleotides [28].

Three-dimensional fold

Interpretation of experimental data on the reactivity of some nucleotides of loop I and of the single-stranded region between hairpins I and II to nucleases, and the observation of compensatory base changes in different TYMV tRNA-like domains is supported by phylogenetic data, since five other valylatable tymoviral tRNA-like domains and one tobamoviral valylatable RNA fold similarly over their last 80 nucleotides [28].

A three-dimensional model of the 3' 82 nucleotides has been built by computer modelling on a graphics system (fig 1C) [27]. This was the first computer modelling of an RNA molecule, a field which has since then developed rapidly [32] because of the difficulty in crystallizing RNA molecules [33, 34]. The amino acid acceptor arm portion of the model was constructed de novo. It shows that a pseudoknot can form according to the stereochemical rules known for RNAs. The rest of the molecule was constructed on the basis of the crystallographically solved structure of yeast tRNAAsp [35] and gives a general view of the tRNA-like structure of the tRNA-like region. This conformation mimics perfectly the fold of a classical tRNA, especially as far as the relative positioning of the amino acid end and the anticodon region is concerned. However, the network of tertiary interactions in the core of the molecule, although not determined with certainty, differs from that found in tRNA. This is also the case for the link of the remaining viral genome with the tRNA-like domain, which is different from that of the extra-RNA domains present in precursor tRNAs. While the viral RNA chain leaves the tRNA-like domain in the hinge region of the L-shaped domain [36], the extra-nucleotides in precursor tRNAs extend the molecule at their 5' and sometimes 3'-ends, which are in close vicinity to each other.

Interaction with proteins from the tRNA metabolism and function: in vitro aspects

Interaction with RNase P

Ribonucleases P are involved in the maturation of pre-tRNAs. These enzymes, composed of a catalytic RNA moiety and a protein subunit, generate mature 5'-ends of tRNAs (reviewed by Altman [37] and Brown and Pace [38]). The tRNA-like structure from TYMV is substrate for RNase P from Escherichia coli [39] and Bacillus subtilis [40]. Cleavage occurs at a position stereochemically equivalent to the cleavage site in canonical tRNAs, namely at the end of the amino acid acceptor arm, at the 3'-end of loop L1. Cleavage does not require the anticodon region of the tRNA, and is dependent upon the presence of a pyrimidine residue at the cleavage site. Cleavage does not require the pseudoknot, although the cleavage rate and specificity are increased in the presence of such a special folding [41]. Additionally, the nature of the nucleotide 5' to the CCA-terminus, as well as the sequence of the first base-pair of the acceptor arm, are of importance.

Interaction with tRNA nucleotidyl-transferase

tRNA nucleotidyl-transferases are other enzymes involved in the maturation of tRNAs (reviewed by Deutscher [42]) and are responsible for the presence of an accurate 3'-end CCA sequence in these mole-
cules. The enzyme from *E coli* incorporates a terminal adenosine to the TYMV RNA deprived of the terminal A [14]. Efficient recognition by this enzyme requires about 75 nucleotides [43].

**Interaction with aminoacyl-tRNA synthetases**

**Valylation properties of the tRNA-like structure**

The tRNA-like domain of TYMV RNA is efficiently recognized by yeast ValRS. Indeed, the kinetic parameters of valylation of the genomic and the subgenomic RNAs of TYMV are very close to those measured for valylation of the cognate tRNA\textsubscript{Val} [17]. The efficiency in aminoacylation of the tRNA-like structure (estimated by the ratio between the kinetic parameters \(k_{cat}\) and \(K_m\)) is 20–30 times lower than that of the cognate tRNA\textsubscript{Val}. This is very efficient when compared to the loss in valylation efficiencies of tRNA\textsubscript{Ile} or tRNA\textsubscript{Met}, the best non-cognate substrates for yeast ValRS [44], which have reduced efficiencies of 4–5 orders of magnitude.

**Optimal length for aminoacylation activity**

Equivalent kinetic parameters of valylation have been measured for the complete genomic RNA and for a fragment of 159 nucleotides containing the end of the coat protein cistron. An unsolved question however, concerns the length of the minimal fragment capable of being aminoacylated. A qualitative answer to this question was obtained by assessing the aminoacylation abilities of fragments originating from nuclease cleavage of the viral RNA. The shortest aminoacylatable fragment was found to be 82 ± 2 nucleotides [43]. Interestingly, such a fragment corresponds to the size required for the formation of the L-shaped structure (see fig 1). A more precise analysis was based on the measurements of kinetic parameters of aminoacylation of a series of RNA fragments of different length [45]. In vitro transcripts corresponding to TYMV RNA 3'-end fragments of 88, 99, 115 or 264 nucleotides, respectively, showed that the shortest fragments are incompletely charged and that the aminoacylation rates were slower than the rate observed for

![Fig 2. Recognition of the TYMV tRNA-like domain with ValRSs (A, B). A, a1. Contact points with yeast ValRS (boxed regions) and identity nucleotides required for efficient aminoacylation by this enzyme (dashed nucleotides). A, a2. Contact points of yeast tRNA\textsubscript{Val} with its cognate synthetase (data are from [47]). B. Identity nucleotides for efficient aminoacylation by wheat germ ValRS. For pseudoknot residue U21, see legend to figure 1.](image-url)
the longest transcript. Thus it appeared that nucleotides upstream the core region formed by the 3' 82 nucleotides, are important for optimal valylation. As will be shown below, nucleotides of this upstream region interact with yeast ValRS. Another and non-exclusive explanation for the lower efficiency of short fragments concerns their potential to form alternative inactive conformations [46]. Indeed, the kinetic analyses were performed on in vitro transcripts presenting six non-viral nucleotides at their 5'-end [45]. Whereas these extra-nucleotides probably have no influence on the folding of the longest tRNA-like molecule, they may interfere with the folding of the shorter ones and thus hinder efficient aminocaylation [46]. Thus, the sequence of 82 nucleotides forming the L-shaped structure of the tRNA-like domain appears to possess the full potential for optimal aminocaylation, but interaction between ValRS and external 5'-sequences may help to stabilize binding of the tRNA-like domain with the enzyme and consequently facilitate its charging.

Recognition by yeast and wheat germ valyl-tRNA synthetase

The signals within the tRNA-like domain recognized by ValRS and important for specific aminocaylation have been searched for using two approaches: the contact points with yeast ValRS have been defined by footprinting experiments and the nucleotides important for aminocaylation by yeast and wheat germ ValRS have been defined by the kinetic analysis of a series of mutated tRNA-like fragments, obtained by in vitro transcription.

The natural 159 nucleotide-long tRNA-like fragment is protected by yeast ValRS against digestion by nucleases and against chemical modification of phosphates by ethynitrosourea essentially in the anticodon and the acceptor arms [31] (fig 2). Interaction with the acceptor arm occurs in spite of the presence of the pseudoknot and shows that the yeast enzyme tolerates some structural variability in this part of its substrate. The global interaction domains are equivalent to the major sites of interaction of ValRS with its cognate tRNA\(^{val}\). These results were thus further evidence supporting the L-shaped folding proposed for the TYMV tRNA-like domain. Interestingly, yeast ValRS interacts also with the region upstream of the L-shaped part of the tRNA-like structure. This additional interaction may contribute to a better stabilization of the tRNA/synthetase complex [31].

Nucleotides important for valylation have been searched for by a mutational analysis of the tRNA-like domain [48, 49]. Variants of the tRNA-like structure were created via site-directed mutagenesis of a cDNA clone (covering the 258 last nucleotides of TYMV RNA) and in vitro transcription with T7 RNA polymerase. Two main loci were investigated for their importance in aminocaylation by both yeast and wheat germ enzymes, namely the anticodon loop and the 'discriminator' base in the acceptor arm (the fourth base starting from the 3'-end). Indeed, sequence comparisons with a consensus sequence of higher euksaryotic tRNA\(^{val}\) pointed to strong homologies with the anticodon and its 3'-flanking bases [28, 50]. Mutation of the central position of the anticodon triplet, replac- ing A-56 by G or U, resulted in a dramatic decrease in the efficiency of valylation by the yeast enzyme (decrease by four orders of magnitude [48]) indicating that this nucleotide is a very strong element, an identity nucleotide, for valylation. Mutation of the anticodon affects the catalysis in a kinetic- rather than affinity-based manner. Indeed, the mutants are still able to interact with the synthetase with affinities comparable to those of the wild-type transcript and behave as competitive inhibitors in the valylation reaction of yeast tRNA\(^{val}\). Minor negative effects on valylation efficiency were observed for mutants with substitutions at the discriminator base, suggesting some contribution to recognition and catalysis by yeast ValRS.

Further studies with wheat germ ValRS investigated in detail the contribution of the anticodon loop nucleotides (see table I) and of the discriminator base to valine identity [49]. Substitution of the discriminator base did not affect valylation, showing the non-importance of this position for wheat germ ValRS. These results suggest already a different recognition pattern by both types of eukaryotic ValRS. Most individual substitutions in the middle (A-56) or 3' (C-55) positions of the anticodon triplet resulted in very poor valylation (\(V_{\text{max}}/K_m\) values more than 1000-fold lower than wild-type); the smallest effect resulted from the C-55U substitution (a 59-fold decrease in \(V_{\text{max}}/K_m\)).

Nucleotides A-56 and C-55 are thus clearly strong valine identity determinants. A third determinant in the anticodon loop was identified from the study of double mutants. While individual substitutions of C-53 resulted in mild loss of valylation (3- to 11-fold decreases in \(V_{\text{max}}/K_m\)), the combined C-55U and C-53A substitutions resulted in drastically reduced valylation (2400-fold decrease in \(V_{\text{max}}/K_m\)). Residue C-53 is thus an important valine identity determinant recognized by wheat germ ValRS, but less potent than those in the anticodon triplet. Mutation of nucleotides A-54 or C-57 yielded RNAs with kinetic parameters not significantly different from those of wild-type RNA. Thus these nucleotides were not considered as valine identity elements. Conversely to what happens with yeast ValRS, the discrimination by wheat germ ValRS appeared to be more based on affinity than on kinetic events, since the valylation of most mutants was mainly affected at the level of \(K_m\).

Analysis of the behaviour of multiple mutants within the anticodon loop allowed to define the relationship between the three identity nucleotides.
Table I. Correlation between valylation and replication of TYMV RNAs.

| Mutant | Anticodon loop sequence$^a$ | Relative efficiency of in vitro valylation | Relative replication in protoplasts |
|--------|----------------------------|------------------------------------------|-----------------------------------|
| Wild-type | 59-CCCACAC-53               | 100                                      | 100                               |
| TY-U-57 | . . . U . . . . . .         | 114                                      | 1.5                               |
| TY-G-56 | . . . G . . . . . .         | 0.02                                     | $\leq 0.2$                        |
| TY-A-55 | . . . . A . . . . .         | 0.46                                     | 2.0                               |
| TY-C-54 | . . . . . . . . . . . . .   | 22                                       | 24                                |
| TY-A-53 | . . . . . . . . . . . . A   | 9.3                                      | 0.12                              |
| TY-U-57/A-55 | . . . . . . . . . . . U . . . . . . A . . . . . | 7.5 | 85 |

$^a$Anticodon positions are underlined.

Indeed, if the effects are additive, the relative specificity constants (relative $V_{\text{max}}/K_m$) of a double mutant equals the product of the relative specificity constants of the single mutants [51]. In cases of cooperative or anti-cooperative effects, the calculated values of the relative specificity constants will deviate from the experimental determinations. This approach allowed to detect a mild cooperative effect between nucleotides C-53 and C-55. Moreover, they revealed an unexpected contribution of nucleotide A-54. Indeed, multiple mutants involving changes at the level of these nucleotides in combination with changes at nucleotides(s) -55 and/or -57, had relative $V_{\text{max}}/K_m$ 1.9-2.6 times lower than expected if the individual mutations were additive.

Conversely, the double mutant U-57/A-55 is a much better substrate than expected and shows compensatory effects between both nucleotides. This anti-cooperativity may reflect that ValRS is sensitive to the conformation of the phosphate backbone as well as to functional groups on the bases of the anticodon [49]. Interestingly, this anti-cooperative behaviour was also observed in vivo [52]. Indeed, a TYMV genomic RNA variant, mutated in the tRNA-like domain (mutation C-55 to A) and thus impaired in its aminoacylation and replication capacities, inoculated to plants, became capable of systemic infection after several weeks in host cells. Sequencing of the progeny viral RNAs revealed the occurrence of a compensatory mutation (C-57 to U). This is an example, at the molecular level, showing that alternative solutions exist for a same biological recognition problem, likely involving compensatory effects within RNA/protein contacts.

Aminoacylation of TYMV RNA is sensitive to local conformations in the RNA [36]. Mutating residue U-43 to C-43, at the junction of the amino acid accepting and anticodon branches of the tRNA-like domain, drastically impairs the valylation capacity of the variant, presumably because it allows the formation of a new G-83/C-43 base-pair that brings the remaining viral RNA closer to the tRNA-like domain and consequently hinders interaction with ValRS. An extensive study of variants mutated in the pseudoknotted region showed that as long as the pseudoknot can form, there is no dramatic change in the valylation of the tRNA-like fragment. However, if the pseudoknot cannot be formed, valylation is suppressed [29].

Degenerated specificity
The specificity of tRNA aminoacylation is often not absolute since synthetases can catalyze mischarging reactions (reviewed by Ebel et al [53] and Giegé et al [54]). This possibility was also suggested for the aminoacylation of TYMV RNA after detection of the binding of amino acids other than valine to viral RNA samples by E coli aminoacyl-tRNA synthetases [28]. However, in these experiments it was not excluded that these amino acids were bound to contaminating cellular tRNAs. Charging of the tRNA-like domain with non-cognate amino acids was explicitly demonstrated by aminoacylation studies of in vitro RNA transcripts [55]. Whereas the 254 nucleotide-long TY-Sma transcript is neither recognized by ArgRS, AspRS nor by PheRS, it is an efficient substrate for yeast HisRS. The efficiency of histidinylation however, is dependent upon the length of the transcript.
The 88-nucleotide long TY-AIu transcript is the best suggesting that the upstream sequences hinder recognition by HisRS, increasing the length of transcripts decreases their ability to be recognized by HisRS, suggesting that the upstream sequences hinder recognition by steric effects. Mischarging occurs not only with wild-type RNA transcripts but also with variants mutated at valine identity positions, thus demonstrating that valine and histidine identities do not overlap, and more specifically, that the middle position of the valine anticodon does not participate in histidine identity.

Histidinylation of TYMV RNA fragments of different length pointed to an efficient histidinylation of a minihelix containing a pseudoknot (fragment TY-AA). This minihelix corresponds to the amino acid acceptor arm of the tRNA-like structure, thus convincingly supporting the view that the major nucleotides responsible for histidine specificity are located in the amino acid accepting arm (fig 3). Likely, in this molecule, as in canonical tRNAHis from E. coli [56] and in minihelices derived from this tRNA [57], an additional nucleotide at position -1 is present for identity. This nucleotide would be the first nucleotide of loop L1 from the pseudoknot in the case of the tRNA-like molecule. Interestingly this nucleotide has been shown to be stacked on top of the acceptor arm helix as is residue -1 in tRNAHis [27, 55] (fig 1C, D). Whereas aminocoylation of classical minihelices has been demonstrated in a number of cases [58], studies on the tRNA-like molecule led to the first evidence for the aminocoylation of a pseudoknoted minihelix.

The mischarging of TYMV RNAs and the derivatives mentioned above has also been studied with wheat germ HisRS (Tsai and Dreher, unpublished). Under a variety of conditions, a maximal charging level of 0.08 mol histidine/mol RNA was observed for RNAs containing the entire tRNA-like structure, and 0.15 mol/mol for TY-AA RNA. These results imply that TYMV RNA would only become moderately histidinylated in its higher plant host cells, and the mischarging with histidine thus appears not to be of primordial biological importance. The different behaviours of the yeast and wheat germ HisRS enzymes also suggest that they utilize different minor identity determinants in selecting their RNA substrates.

Recognition by elongation factors

Elongation factors sequester aminoacylated tRNAs and carry them to the ribosome where the adaptor function of tRNA occurs (reviewed by Riis et al. [59]). The aminoacylated TYMV tRNA-like structure, like charged tRNAs, interacts with elongation factors EF-Tu from E. coli [12], from Thermus thermophilus [43], and EF 1α from higher organisms [13]. The minimal sequence required for interaction with the E coli factor was determined by retention experiments testing the binding of aminoacylated fragments to immobilized elongation factor [43]. A fragment of 47 nucleotides, corresponding to the 3'-end of the viral RNA was the shortest RNA segment able to bind to the elongation factor. This fragment corresponds to the acceptor arm of the tRNA-like fragment, suggesting that the pseudoknot does not interfere with the binding events.

Biology of the TYMV tRNA-like structure

Role in protein synthesis or in viral genome replication?

Soon after the discovery of the efficient aminocoylation of plant viral genomes such as TYMV RNA, enquiries on the role of tRNA mimicry in the viral life cycle focussed on interactions with the ribosomes. It was thought that if interaction of charged viral RNA with the A site occurs in much the same way as that of tRNA ternary complexes, preferential translation of viral RNA may result. This might occur by displacement of host mRNAs, with or without actual donation of the amino acid charged on the viral RNA to the nascent polypeptide. However, no convincing evidence was ever obtained for amino acid donation [16]. Similar conclusions arose from studies on other aminocoylatable viral genomes [60, 61].

Subsequently, the focus has shifted towards an involvement of the tRNA-like structure directly in viral RNA replication. This was prompted by the expectation that RNA promoter sequences directing (−) strand synthesis are likely to be located near the 3'-end of the genomic RNA, and by the adoption of the replicase of the RNA bacteriophage Qβ as a paradigm for the replicases of the plant viruses with aminocoylatable RNAs [62]. Core Qβ replicase is comprised of a virus-encoded polymerase subunit and three host subunits, EF-Tu, EF-Ts and the ribosomal protein S1.

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Minihelix mimicking the amino acid acceptor arm of the TYMV tRNA-like structure, with residue U22 opposite to discriminator base A4 (for numbering see legend to figure 1).
It has been shown that the TYMV tRNA-like structure can be valylated in vivo, either in *Xenopus laevis* oocytes [64] or in Chinese cabbage leaves during a natural infection [65]. In *Xenopus* oocytes, the genomic RNA became fragmented, and the valylated species was a 4-5S RNA fragment of the genome [64]. The form of TYMV RNA valylated in Chinese cabbage plants was not determined, but since 3'-fragments and the entire genomic RNA can be valylated in vitro with similar efficiencies [24], it is assumed that at least some of the genomic RNAs are valylated during infection. The genomic and subgenomic RNAs in the virion lack the 3'-A, and terminate in 3'-CCOH [18, 19], and so the encapsidated RNAs cannot be directly valylated. This implies that most of the newly synthesized viral RNAs – those destined for encapsidation – do not become valylated. Valylation is thus restricted to those viral RNAs that are engaged in translation or those that are participating in replication as templates. No way has yet been devised to directly assess the aminoacylation status of the viral RNAs engaged in these different functions.

The relevance of valylation to TYMV replication has been studied genetically using TYMV RNAs with mutations in the valine identity elements of the anticodon region. The same mutations used in TY-Sma-derived RNAs to map the valine identity determinants [49] were transferred into genome-length transcripts, derivatives of the infectious TYMC RNAs made by transcription with T7 RNA polymerase from cloned DNA from the TYMV Corvallis strain [8]. A direct correlation was observed between the replication of the genomic transcripts in Chinese cabbage protoplasts and the valylation efficiencies of the related 3'-fragments as charged by wheat germ ValRS in vitro [66] (table 1). A low basal level of RNA replication, detected as (-) strand synthesis in inoculated protoplasts, was observed for mutants with $V_{\text{max}}/K_m$ values for *in vitro* valylation that were $<10^{-2}$ relative to wild-type (mutants with substitutions at the middle or 3'-positions of the anticodon). These mutants were unable to infect whole plants. By contrast, mutants such as those with substitutions in the wobble position (-57), which could be efficiently valylated, replicated to levels similar to wild type in both protoplasts and whole plants. Mutants with intermediate levels of valylatability (C-54 and A-53 mutants) replicated to intermediate levels *in vivo* (table 1).

The correlation between replication and valylatability was further strengthened by the characterization of a mutant that arose spontaneously in a plant inoculated with TYMC-(A-55) RNA, which does not support systemic infection in plants. Apparently as a result of polymerase error during replication, the plant became systematically infected with progeny having a U-57/A-55 genotype. This genotype was reconstructed in TYMC-(U-57/A-55) RNA, which was shown to replicate to levels comparable to TYMC *in vivo* [52]. Besides replicating efficiently, this mutant RNA could also be relatively efficiently valylated: the RNA could be valylated to completion, and 3'-transcripts with the U-57/A-55 mutations were valylated with a $V_{\text{max}}/K_m$ value 0.075 relative to wild-type. The C-57U mutation has thus clearly suppressed the effect of the C-55A mutation in both the valylation and replication properties. The above data imply that valylation of the viral RNA by host ValRS is a requisite for the efficient replication of TYMV RNA.

The mechanistic basis of the requirement for valylation is not known at present, nor is it known whether efficient charging with an amino acid other than valine is compatible with efficient replication. It is likely that the valylation status of a molecule of TYMV RNA influences its potential as a template for (-) strand synthesis, the first stage of viral RNA replication, the effect being exerted at the level of (-) strand promoter selection. It will be important to identify the host subunits of TYMV replicase, and so determine the validity of the Q8 replicase as a paradigm. Elongation factor EIF1α would be a logically expected subunit, compatible with the correlation discussed above, but two studies have failed to detect this protein in preparations of TYMV replicase [13, 67].

**The role of a pseudoknot upstream of the tRNA-like structure**

Nucleotides between the CP-ORF termination codon and the tRNA-like structure can be folded into a pseudoknot (see fig 1), a structural element commonly found in viral RNAs and which has also been implicated in translational frameshifting in RNA viruses [68]. Because of their distinctive structure, pseudoknots may function as protein recognition elements involved in the regulation of translation and/or RNA replication.

In the course of studies on the role of the 3'-non-coding region of TYMV RNA, it was observed that mutation of U-96G, which destabilizes the base-pairing needed for the formation of one of the helical segments of the pseudoknot upstream of the TYMV tRNA-like structure, resulted in a poor replication rate and inability to infect plants systematically [52].
used to inoculate Chinese cabbage plants, some plants did develop symptoms, but these infections were all due to progeny with altered sequences. Most plants contained progeny with simple G-96U reversions, but in a few cases, second-site suppressing mutations were responsible for the ability to develop systemic symptoms. In one such case, the suppressing mutation was found to be the A-107C substitution, which recovered the ability to form a pseudoknot similar to that present in the wild-type RNA (see fig 1) [52]. This mutation also abolished the normal CP-ORF termination codon, such that the extended CP-ORF terminated five codons downstream. TYMC-(C-107/G-96) was able to produce normal systemic symptoms in plants, but the virions were less stable than normal, resulting in the degradation of virion RNA in mature, infected leaves as virions dissociate. This suggests that the extended coat protein plays no role in rescuing the poor replication caused by the G-96 mutation, and that the rescue is due to the fact that the U-107C mutation recovers the ability to form a pseudoknot. The pseudoknot may represent a binding element recognized by the replication machinery in assembling a (−) strand replication complex. It is interesting that the mutated nucleotides are protected when yeast ValRS binds TYMV RNA [31], but no altered valylation properties were observed for 3′-transcripts with the G-96 mutation [52]. The presence of one or several pseudoknobs in the 3′-non-coding region just downstream of the stop codon is a common feature of plant viral RNA [69]. In the case of TMV RNA, these pseudoknots were shown to be necessary for the replication of the viral RNA [70].

Possible role of the TYMV tRNA-like structure as the 3′ telomere

As mentioned above, TYMV virion RNA lacks the complete 3′-CCA₀ required for aminoacylation, yet viral RNAs are known to be valylated in vivo. It is presumed that host tRNA nucleotidyl-transferase functions to add the missing 3′-A residue to disencapsidating RNAs after inoculation. tRNA nucleotidyl-transferase serves to maintain intact 3′-termini on cytoplasmic tRNA molecules, whose 3′-ends are constantly at risk from exonuclease attack and which turn rapidly over in the cell [42]. The site of (−) strand initiation on the tRNA-like structure template has not been determined for TYMV, but it is likely to be within one or two nucleotides of the 3′-end, as has been shown with BMV [71]. Thus, loss of 3′-nucleotides could cause the loss of the (−) strand initiation site and abortion of the entire replication cycle. Thus, the role of host tRNA nucleotidyl-transferase in maintaining intact 3′-ends is probably a crucial one, and is a role analogous to that of the telomerase enzymes that add T/G-rich sequences to the ends of linear DNA chromosomes to prevent the loss of terminal sequences during replication [72]. Since the tRNA-like structure is a feature that makes the viral RNA a substrate for tRNA nucleotidyl-transferase, the tRNA-like structure may be considered the telomere of the linear viral RNA genome. While this functional role is supported by experimental evidence in the case of BMV RNA [73], it remains to be demonstrated for TYMV RNA.

Function of the tRNA-like structure as promoter for (−) strand synthesis

Studies with TYMV are not yet sufficiently advanced to provide as convincing a case as with BMV for the involvement of the tRNA-like structure in promoting (−) strand synthesis. Nevertheless, data consistent with such a role for the TYMV tRNA-like structure exist. Morch et al [74] used short RNA transcripts from the 3′-end of the genome as templates in vitro for a TYMV replicase preparation, and as inhibitors of the copying of TYMV RNA. Despite the presence of a 3′-tract of 50 adenosine residues and 31 further heterologous nucleotides, an RNA containing the 100 3′-nucleotides of TYMV RNA (ie including the tRNA-like structure) was able to function as a template for RNA synthesis [74]. This RNA was also able to inhibit the copying of TYMV RNA when present at 2.5- to 25-fold molar excess. A related RNA containing a longer segment from the 3′-end of TYMV RNA had similar properties, but these were abolished by the removal of the tRNA-like structure. In further studies, shorter fragments containing only part of the tRNA-like structure were tested in the above assays. RNAs containing as few as 38 nucleotides from the 3′-end of TYMV RNA (ie the amino acid acceptor stem and part of the T-stem analogue) as well as eight unspecified nucleotides at the 5′-end and 13 at the 3′-end, were able to inhibit TYMV RNA copying in vitro, and appeared to act as templates for the TYMV replicase.

In preliminary experiments using a TYMV replicase preparation, we have obtained results consistent with the above observations (Georgel and Dreher, unpublished). Transcripts with correct 3′-CCA termini and containing variable lengths of sequence from the 3′-end of TYMV RNA were used as templates for TYMV replicase. RNAs derived from TY-Sma, TY-Dde, TY-Dra and TY-Alu clones and containing 258, 109, 93 and 82 nucleotides of TYMV RNA (ie the amino acid acceptor stem and part of the T-stem analogue) as well as eight unspecified nucleotides at the 5′-end and 13 at the 3′-end, were able to inhibit TYMV RNA copying in vitro, and appeared to act as templates for the TYMV replicase.
Fig 4. Other tRNA mimicries. A. Plant viral tRNA-like domains from TMV (a1) and BMV (a2). B. tRNA-like region with the regulatory region of the messenger RNA of E coli ThrRS. C. tRNAs with minor structural changes: (c1) tRNA Gly from S epidermidis and (c2) tRNA Sec (specific for selenocysteine) from E coli. D. Bizarre mitochondrial tRNA sequences from nematodes (obtained from gene sequencing) (d1 and d2) and from bovine heart (d3). In C, structural features in tRNA sequences deviating from standards are shown.
Sma containing 18 additional 3'-nucleotides (GGGGGUACCAGGCUCGAAU), an RNA that was itself not copied by the TYMV replicase (Georgel and Dreher, unpublished).

The above results thus are consistent with the localization of the (-) strand promoter to the 3'-tRNA-like structure of TYMV RNA. We remain concerned, however, about the specificity of the in vitro studied TYMV replicase, and the relevance of the results obtained with TYMV infection processes.

Other functional possibilities

Besides the replication-related roles discussed above, it is likely that other, perhaps more subtle roles for the tRNA-like structure will emerge in the future. It has been suggested that the sequences forming the TYMV tRNA-like structure may assume an alternative conformation involving base-pairing with 5'-terminal sequences [75]. Such pairing is suggested by light-scattering experiments [76], and would prevent recognition of both the initiation codons of ORF-69 and ORF-206, and of the tRNA-like structure. This may play a role in the regulation of the infection process. The 3'-non-coding regions of viral RNAs (especially TMV RNA) have been shown to increase mRNA expression in a number of cell types [77, 78], and may thus be important in ensuring adequate viral gene expression. Other regulatory roles involving RNA binding to the ribosome along the lines of the suggestions discussed earlier should not be disregarded. Ribosomes are known to play a role in the active disencapsulation of viruses [79, 80], and it would be interesting to determine with modern binding studies whether the tRNA-like structure has any affinity for ribosomes.

Other tRNA-like molecules and mimicries with tRNAs

The presence of a tRNA-like domain within the RNA for TYMV is not an exception in the constitution of viral genomes. Several other plant viruses present similar functional domains at the 3'-end of their RNA genomes. Besides tymoviruses, whose RNAs specifically charge valine, there are two other large groups of viruses presenting an aminoacylatable domain. Tobamoviruses are histidinylatable (with one exception, the CcMV strain, charged with valine) and bromo-, cucumo-, and hordeiviruses are all tyrosylatable. All these viral RNAs are also recognized by a series of tRNA-specific proteins like elongation factors and tRNA nucleotidyl-transferases (reviewed by Mans et al [50] and Florentz and Giegé [81]). Their structural mimicry with tRNAs is much more complicated than that found for TYMV (fig 4A). The extreme case is RNA from BMV that presents the most intricate sequence (fig 4a2), and nevertheless is well charged by TyrRS. As for the TYMV tRNA-like domain, the biological role of these structures is now being deciphered [50, 81], and involvement during replication is also strongly suggested in the case of BMV. But in contrast to TYMV RNA, aminoacylation is not required for BMV RNA replication [71, 82].

tRNA mimicries have also been found in several other biological systems. Thus, in messenger RNAs coding for aminoacyl-tRNA synthetases (E. coli HisRS, ThrRS and MetRS), the regulatory region of the mRNA contains tRNA-like elements [83-85]. For ThrRS mRNA, the structure of the tRNA-like domain is well established (fig 4B) [86] and it was shown that interaction with the synthetase regulates the level of transcription of the gene [86, 87]. Interestingly, even for natural tRNAs, a large variability in structure has been revealed (fig 4C, D). Small sequence variations have been found in tRNAs involved in specialized biological functions. For example, tRNA\textsubscript{Gly} from Staphylococcus epidermidis, which participates in cell wall synthesis (but not in classical protein synthesis) is devoid of the classical TVC sequence as well as of canonical D-loop characteristics (fig 4Cl) [88]. Noticeable are the sequence similarities between this glycine-specific tRNA and the homologous regions in the TYMV tRNA-like domain (see fig 1) [25], which also is not involved in ribosome directed protein synthesis. The recently discovered tRNA\textsubscript{Sec} species, incorporating the 21th amino acid selenocysteine into a limited number of proteins has an acceptor arm with 13 base-pairs and presents peculiar tertiary interactions [89, 90] (see fig 4c2 for the secondary fold of E. coli tRNA\textsubscript{Sec}). Large structural changes can also occur in other tRNAs participating in protein synthesis. This was first found in mammalian mitochondrial tRNA\textsubscript{Ser} species lacking the entire D-loop and stem [91, 92] (fig 4D). More recently a series of other mitochondrial tRNAs have been discovered with bizarre structures, in particular in nematodes (fig 4D) [93], in which D-stems, T-stems, or both of them show strong structural deviations and can even be absent.

What is the evolutionary significance of the great structural variability within RNA molecules that can be recognized by tRNA specific proteins? While a common origin can be postulated for tRNAs presenting a cloverleaf structure (even if parts of the cloverleaf are missing), there are strong theoretical arguments for a different evolutionary origin of tRNA-like molecules in which the tRNA mimicry results from different architectural principles (as in plant viral RNAs where pseudoknots are required). Thus, it is likely that molecular tinkering events occurred in the early development of life when strategies for both
transcriptional and translational mechanisms had to be found by nature. Because these processes are so basic in all forms of life, they could have involved common molecules, but also alternate structures able to fulfill similar functions. This scenario is not unlikely, since in vitro or in vivo genetic selection methods have recently generated RNA molecules with functional tRNA characteristics but presenting novel structural features not found in known tRNAs (or tRNA-like molecules) [94, 95].

As a conclusion, and from a more practical point of view, we emphasize that tRNA-like molecules are excellent tools to understand tRNAs. Studies on TYMV RNA valylation have shown that the functional strategy used by nature to ensure the aminoacylation identity of this RNA is the same as for canonical tRNAs. Indeed, the identity nucleotides responsible for valylation are located at topologically similar places in the structure of both types of molecules [48, 49, 81]. Extending this concept, it should be possible to engineer molecular substrates of synthetases in which identity elements are carried by adequate structural frames allowing them to interact with their counterparts on the synthetases. A first application came from the engineering of an RNA minihelix derived from the TYMV tRNA-like domain, in which the histidine identity element is mimicked by a residue from the terminal pseudoknot that structurally mimics the major identity nucleotide from histidine specific tRNAs (see above). As anticipated, the minihelix became an efficient substrate for HisRS [55].

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We dedicate this paper to the memory of Professor Léon Hirth (1920–1991). He contributed enormously to the development of plant virology in France and promoted studies on TYMV RNA. He encouraged our projects on the aminoacylation of TYMV RNA at a time when this phenomenon was not well understood. We acknowledge the many discussions we had with him on this topic and on its relevance to virology.

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