P-site tRNA is a crucial initiator of ribosomal frameshifting

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ABSTRACT
The expression of some genes requires a high proportion of ribosomes to shift at a specific site into one of the two alternative frames. This utilized frameshifting provides a unique tool for studying reading frame control. Peptidyl-tRNA slippage has been invoked to explain many cases of programmed frameshifting. The present work extends this to other cases. When the A-site is unoccupied, the P-site tRNA can be repositioned forward with respect to mRNA (although repositioning in the minus direction is also possible). A kinetic model is presented for the influence of both, the cognate tRNAs competing for overlapping codons in A-site, and the stabilities of P-site tRNA:mRNA complexes in the initial and new frames. When the A-site is occupied, the P-site tRNA can be repositioned backward. Whether frameshifting will happen depends on the ability of the A-site tRNA to subsequently be repositioned to maintain physical proximity of the tRNAs. This model offers an alternative explanation to previously published mechanisms of programmed frameshifting, such as out-of-frame tRNA binding, and a different perspective on simultaneous tandem tRNA slippage.

Keywords: translation; recoding; kinetic model; frameshifting; ribosome

INTRODUCTION
A primary feature of the genetic code is its triplet character. During standard translation, codons do not overlap and are not separated by any nucleotides. This feature provides the basis for predicting coding sequences in mRNA. A sequence of triplets in mRNA not interrupted by a stop codon is defined as an open reading frame (ORF). Generally, a single ORF encodes a single protein. However, during translation of certain mRNAs, a proportion of ribosomes shift frame at a particular site, and a second product is synthesized. This product can have more or fewer amino acid residues than the product of standard decoding, depending on the existence of a partially overlapping ORF and its disposition. ORF fusion is widely used during the expression of viral genes and mobile elements, and several examples in decoding nonmobile chromosomal genes have been studied in detail (for reviews, see Brierley and Pennell 2001; Baranov et al. 2002a; Stahl et al. 2002; Baranov et al. 2003).

Programmed ribosomal frameshifting is an extension of genetic decoding that expands standard gene readout. More than one product can be synthesized from the same gene. For example, in Escherichia coli dnaX decoding, two proteins are synthesized from the same mRNA in an equimolar ratio, one by standard translation and the other one via frameshifting (Blinkowa and Walker 1990; Flower and McHenry 1990; Tsuchihashi and Kornberg 1990). Frameshifting also offers the prospect of conditional regulation of gene decoding: with bacterial release factor 2 (RF2), its own cellular concentration affects the efficiency of the frameshifting that is required for its biosynthesis (Craigie and Caskey 1986); whereas with mammalian antizyme, the efficiency of obligatory frameshifting depends on the concentration of polyamines whose synthesis is negatively regulated by antizyme (Matsufuji et al. 1995).

In addition to intrinsic interest, genes utilizing frameshifting for their expression provide an opportunity to understand the mechanism of open reading frame control. There have been several models dealing with the phase of the ribosomal cycle at which frameshifting occurs (Jacks et al. 1988a; Weiss et al. 1989; Horsfield et al. 1995; Farabaugh and Björk 1999; Harger et al. 2002) and others dealing with the influence of codon–anticodon interactions on frameshifting (Brierley et al. 1991; Tsuchihashi and Brown 1992; Curran 1993; Kolor et al. 1993; Sundararajan et al. 1999; Atkins et al. 2000). The relationship of these models to the present one, will be considered below.
out-of-frame, it should be rejected as a noncognate tRNA, because no base pairs are formed in the first position of the A-site codon. If this is true, and supportive evidence has been presented by Hansen et al. (2003), the incoming tRNA can be incorporated into the ribosome only after P-site tRNA re-pairing to mRNA at a codon in a new frame. There are two distinct elongation phases where repositioning of the P-site tRNA relative to mRNA can occur (Fig. 1). Repositioning involves codon:anticodon dissociation, relative movement of the anticodon to mRNA, and re-pairing to mRNA at a new codon.

The first occurs prior to the final (irreversible) acceptance of the A-site tRNA, that is, during the initial recognition of the A-site tRNA and/or the proofreading step (Fig. 1). If P-site tRNA repositioning occurs at this phase, then the next incorporated amino acid will be specified by the codon in the new frame. Repositioning can occur in both plus and minus directions, although in the case of the minus direction, the tRNA in the E-site could interfere (Horsfield et al. 1995; Baranov et al. 2002b; K.H. Nierhaus, pers. comm.). If the E-site plays such a role, E-site tRNA slippage or its relative dislocation should precede P-site tRNA repositioning. If the P-site tRNA can dissociate from its codon in the initial frame and bind to a codon in a new frame, then it may also be able to shift back to the initial frame, before the A-site tRNA is irreversibly incorporated into the ribosome. This would mean that the success of frameshifting largely depends on the availability of tRNAs or release factors competing for available A-site codons in different frames. This point is considered further below.

The second elongation phase to consider is after final acceptance of the A-site tRNA into the ribosome, but before translocation is complete (Fig. 1). In this case, the identity of the A-site tRNA corresponds to that specified by the zero frame codon. Because an irreversible choice of A-site tRNA has already been made, the concentration of potential competing tRNAs for the A-site is irrelevant for frameshift efficiency. Productive forward repositioning of the A-site tRNA is assumed to be unlikely because no base pairs would be formed with the first A-site codon base. In this scenario, P-site repositioning can occur only in the minus direction, because nucleotides 3' of the P-site codon are occupied by A-site tRNA that cannot be rejected. A substantial refinement of the presently accepted hypothesis of simultaneous slippage of two tRNAs suggested for this elongation phase by Jacks et al. (1988a) is provided below.

**FIGURE 1.** Simplified scheme of the translation elongation cycle. Two main types of ribosomal frameshifting are distinguished depending on the phase of the elongation cycle where tRNA repositioning occurs.
Effect of stability of the P-site codon:anticodon complex on reading frame maintenance

Selection of tRNA in the A-site is efficiently monitored by the ribosome. The geometry of base pairs in the first two positions of the A-site codon:anticodon duplex is controlled more precisely than the geometry of the base pair in the third position (for review, see Ramakrishnan 2002). This allows the ribosome to discriminate cognate tRNAs from near-cognate tRNAs that could form energetically stable complexes with the same codon in mRNA (i.e., although the codon UUU can form base pairs with both the AAG and GAI anticodons and energetically both complexes are nearly equal, only AAG is a cognate anticodon for UUU). Such discrimination of tRNAs in the A-site is vital for accurate translation. However, there is no need to control the geometry of the P-site tRNA codon:anticodon complex in the same way. First, the P-site tRNA correctness has already been monitored while it was in the A-site. If the identity of the P-site tRNA were controlled, its rejection would terminate translation because the tRNA is already attached to a growing polypeptide chain. (Indeed, studies by Potapov et al. [1995] have shown that DNA matrices may be used instead of mRNA for the P-site tRNA binding, but not for the A-site tRNA binding.) The only case in which recognition of mRNA occurs in the P-site is during translational initiation. In bacteria there are several other initiator codons that can be used instead of the standard AUG, for example, GUG, UUG, and in rare cases AUU. These codons are recognized by the same tRNA\textsuperscript{Met} (whereas elongation tRNA\textsuperscript{Met} recognizes internal AUG codons exclusively). This leads to the conclusion that ribosomes do not discriminate the positions of a codon:anticodon duplex in the P-site in the same way as they do in the A-site (e.g., tRNAs with the anticodons AAG and GAA may be equally stable in complex with the codon UUU).

If we consider the P-site tRNA repositioning as a chemical reaction, its rate may be expressed by the following equation:

$$\left( \frac{dR_0}{dt} \right) = -k_1[R0] + k_2[R1] = -\left( \frac{dR_1}{dt} \right)$$

where [R\textsubscript{0}] is the concentration of ribosomes with P-site tRNA in the complex with the codon in the zero frame, [R\textsubscript{1}] is the concentration of ribosomes with the complex in a new frame, k\textsubscript{1} is the rate constant of the forward reaction, and k\textsubscript{2} is the rate constant of the reverse reaction. The stage when P-site tRNA is entirely dissociated from the mRNA can be considered as the activation complex for this reaction. In this case, the value of k\textsubscript{1} and k\textsubscript{2} would depend primarily on the energetic barriers between the corresponding codon:anticodon complexes and dissociated complexes (Fig. 2). This means that the efficiency of frameshifting depends not only on the stability of the complex in a new frame, but also on the stability of the complex in the initial, or zero frame.

The importance of the stability of the codon:anticodon complex in a new frame has been noted before (Jacks et al. 1988a; Weiss et al. 1990a; Tsuchihashi and Brown 1992). Curran (1993) performed a systematic analysis of frameshifting efficiency in a derivative of the release factor 2 (RF2) frameshifting cassette CUU UAG C with 32 different codons preceding the stop codon (the wild-type UGA stop codon was substituted with UAG). (Spaces are used to separate codons in the initial ORF, underlining shows the new frame.) He found that the most frameshift-prone codons are CUU > CCC > UUU > GUU > CUC, where CUU is the naturally occurring shifty codon in all bacterial genes encoding RF2 that use frameshifting for their expression. Curran (1993) was able to build an exponential correlation between frameshifting efficiency on these codons and the stability of the P-site tRNA in a complex with an overlapping +1 frame codon using simplified calculations of the energies of the codon:anticodon complex. However, in some cases no correlation was found. For instance, tRNA\textsuperscript{Phc}(3′-AAG-5′) should form a more stable complex with the +1 codon UUU in the sequence U UUU than would tRNA\textsuperscript{Luu}(3′-GAG-5′) in the sequence C UUU. Nevertheless, the observed frameshifting efficiency at the sequence U UUU was lower. Thus, it is not sufficient to estimate the stability of complexes relying solely on the energies of the base pairs involved. Certain mutations in tRNAs outside of the anticodon loop can affect the rate of P-site tRNA dissociation (e.g., Herr et al. 1999).

The effect of stability of the initial P-site tRNA:mRNA complex on frameshifting has been observed (e.g., Farabaugh et al. 1993) and experimentally studied in detail especially using the efficient ribosomal frameshifting that occurs during decoding of yeast Ty transposable element mRNAs (Vimaladithan and Farabaugh 1994; Pande et al.)
is UCC UGA U; however, several mutations in the UCC frameshifting site in mammalian antizyme 1 complex may be sufficient for the efficient frameshifting. Based on experiments with mutated mammalian antizyme +1 frameshifting cassettes, Ivanov et al. (2003) invoked structural obstacles to explain frameshifting on such codons. However, these several considerations indicate that there is no need to invokes structural obstacles to explain frameshifting in the P-site tRNA-anticodon complex. This led to the proposal that there are special features in those tRNAs that promote frameshifting by interfering with the binding of the incoming A-site tRNA (Vimaladithan and Farabaugh 1994). In this model, A-site tRNA recognition occurs in the alternative frame without a shift of the P-site tRNA. In their latest review, this group even considered the more radical hypothesis that in yeast, frameshifting occurs only caused by the out-of-frame binding of tRNA in the A-site (Stahl et al. 2001). However, if the initial codon:anticodon complex is weak, P-site tRNA repositioning to a codon in a new frame may be efficient even if the stability of the new complex is also weak (Fig. 2; kinetic model below); in this case, there would be a correlation between the propensity for re-pairing in the new frame and frameshifting efficiency, but it is not direct. The idea that near-cognate tRNAs are prone to slippage was also proposed by Gallant et al. (2000). Furthermore, recent work (A.J. Herr, N.M. Wils, C.C. Nelson, R.F. Gesteland, and J.F. Atkins, in prep.) has shown that in a case of “extreme” P-site repositioning—hopping—ribosomes carrying P-site tRNA can, in certain circumstances, re-engage with mRNA, in positions where no Watson-Crick base pairs can be formed between the codon and anticodon. Taken together, these several considerations indicate that there is no need to invoke structural obstacles to explain frameshifting on such codons. Based on experiments with mutated mammalian antizyme +1 frameshifting cassettes, Ivanov et al. (2003) proposed that <2 bp in the new, shifted codon:anticodon complex may be sufficient for the efficient frameshifting. The wild-type frameshifting site in mammalian antizyme 1 is UCC UGA U; however, several mutations in the UCC codon that diminish base-pairing of tRNA anticodon with mRNA codon in the +1 frame do not eliminate frameshifting (Matsusufi et al. 1995).

In summary, the stability of P-site codon:anticodon complex formation in the initial and new frames should greatly influence to the efficiency of frameshifting.

**Effect of competitors in the A-site**

As considered above, when P-site tRNA repositioning occurs during tRNA recognition before the peptidyl transferase reaction, the efficiency of A-site tRNA acceptance should depend on the concentration of tRNAs or release factors corresponding to A-site codons in the different frames. Despite the potential ability of P-site tRNA to be repositioned on certain overlapping codons, realization of this potential depends on several features, among which relative tRNA concentration and affinity for mRNA are prominent. The kinetic model described below indicates how these features affect P-site tRNA repositioning and consequently frameshifting.

Let us consider a sequence of the general type 000 111 2, where the tRNA that recognizes codon 000 is capable of being repositioned to codon 001 when it is in the P-site. These processes can be considered as a combination of five reactions:

\[
  \begin{align*}
    k_0 & \; R \rightarrow R_0 \\
    k_1 & \; R_0 \rightarrow R_1 \\
    k_2 & \; R_1 \rightarrow R_0 \\
    k_3 & \; R_0 + T_0 \rightarrow R_0T_0 \\
    k_4 & \; R_1 + T_1 \rightarrow R_0T_1
  \end{align*}
\]

where \( R \) is the ribosome incoming into this particular sequence; \( R_0 \) is the ribosome with its P-site tRNA base-paired to the 000 codon; \( R_1 \) is the ribosome with its P-site tRNA base-paired to the 001 codon; \( T_0 \) is a tRNA corresponding to the A-site codon 111; \( T_1 \) is the A-site tRNA corresponding to the 112 codon; \( R_0T_0 \) is a complex of the ribosome with irreversibly bound P-site and A-site tRNAs in frame 000 111; and \( R_0T_1 \) is a complex where P-site and A-site tRNAs are irreversibly bound in the +1 frame 001 112. The rates of all reactions can be expressed as follows:

\[
  \begin{align*}
    d[R_0]/dt &= k_0[R] - k_1[R_0] + k_2[R_1] - k_3[R_0][T_0], \\
    d[R_1]/dt &= k_1[R_0] - k_2[R_1] - k_4[R_1][T_1], \\
    d[R_0T_0]/dt &= k_3[R_0][T_0], \\
    d[R_0T_1]/dt &= k_4[R_1][T_1],
  \end{align*}
\]

where we assume that the concentration of the incoming ribosomes \([R]\) is large and does not change significantly in the course of this reaction, so that ribosomes arrive at a given sequence with a constant speed \( k_0[R] \). In the stationary regime, from

\[
  d[R_1]/dt = k_1[R_0] - k_2[R_1] - k_4[R_1][T_1] = 0,
\]
we get a value for \([R_0]/[R_1]\), which is

\[
\frac{[R_0]}{[R_1]} = \left( k_2 + k_4[T_1] \right)/k_1,
\]

where we also assume that the concentration of ribosomes at this particular sequence is significantly lower than of tRNAs, so that \([T_0]\) and \([T_1]\) can also be considered as constants.

Next, from the constant \([R_0]\) and \([R_1]\) of the stationary regime, we obtain

\[
d[R_0]/dt = k_3[T_0][R_0] = \text{const., i.e.,} \]

\[
[R_0] = k_3[T_0][R_0]t + \sigma_0,
\]

and

\[
d[R_1]/dt = k_4[T_1][R_1] = \text{const., i.e.} \]

\[
[R_1] = k_4[T_1][R_1]t + \alpha_1,
\]

where \(\sigma_0, \alpha_1\) are integration constants representing the concentrations \([R_0]\) and \([R_1]\) achieved during the initial equilibration stage of the reaction. After some time, these constant terms become negligible as compared with the terms linear in time, \(t\), and then we can write for this later stage,

\[
\frac{[R_0]}{[R_1]} = k_3[T_0][R_0]/k_1 k_3[R_1][T_1].
\]

Substituting the stationary value of \([R_0]/[R_1]\), we finally obtain

\[
\frac{[R_0]}{[R_1]} = k_3[T_0]/k_1 k_4[T_1] + 1.
\]

This equation illustrates the dependence of frameshifting efficiency \([R_1]/[R_0]\) on the concentration of “in-frame” and “out-of-frame” A-site tRNAs. The graphical representation of this dependence is shown in Figure 3.

It is clear that the effect of A-site tRNA corresponding to the +1 frame codon and the effect of stability of P-site tRNA in a shifted complex are limited. As seen in Figure 3B, a low concentration of this tRNA would favor standard triplet translation. A high concentration should increase the efficiency of frameshifting; however, this effect is limited so that the efficiency cannot exceed the limit \(k_3/k_4[T_0]\). On the contrary, the concentration of tRNA corresponding to the in-frame A-site codon (Fig. 3A) plays a more significant role. Its high concentration favors triplet decoding, its lower concentration favors frameshifting. Depletion of this tRNA should bring frameshifting efficiency to virtually 100%. Figure 3C and D, illustrates the effect of forward and reverse P-site tRNA repositioning rates on frameshifting. \(k_1\) is the rate of forward repositioning from the zero-frame codon to the plus-frame codon. This rate depends on energetic differences between the zero-frame complex and the dissociated complex (Fig. 2). Decreasing its stability, for example, by incorporating a near-cognate tRNA in the P-site, should increase frameshifting efficiency (Fig. 3C). \(k_2\) corresponds to the rate of reverse P-site tRNA repositioning. It depends on energetic differences between the complex of P-site tRNA with the plus-frame codon and the dissociated complex (Fig. 2). Low affinity of the P-site tRNA to the plus-frame codon favors triplet decoding, whereas its high affinity favors frameshifting. However, even if P-site tRNA forms a perfect complex with the plus-frame codon, frameshifting efficiency cannot exceed the \(k_1/k_4[T_0]\) value (Fig. 3D). These considerations illustrate the parameters, cellular tRNA concentration and their codon affinity, that are necessary for standard triplet decoding. If these requirements are not fulfilled for tRNAs involved in decoding a particular combination of codons, this combination of codons would be shift-prone. A later paragraph deals with such examples.

There are several cases of experimentally analyzed +1 frameshifting that support this kinetic model. The Ty1 and Ty3 frameshifting sites are CUU AGG C (Belcourt and Farabaugh 1990) and GCG AGU U (Farabaugh et al. 1993), respectively. In both cases, the frameshifting rate should depend on the concentration of tRNAs that recognizes the zero- and +1-frame A-site codons. Under normal conditions in yeast, the cellular concentration of tRNAs depends on their gene copy numbers (Percudani et al. 1997). With Ty1, the zero-frame codon is recognized by tRNA\(^{\text{Asg}}\)(3’-UCC-5’), which is encoded by only a single gene. The +1-frame codon is recognized by tRNA\(^{\text{Gly}}\)(3’-CCG-5’), which is encoded by 16 identical genes. In decoding Ty3 mRNA, the zero-frame codon is recognized by tRNA\(^{\text{Val}}\)(3’-UCG-5’), which is encoded by 14 gene copies. In both cases, misbalance between the concentra-
tions of the corresponding tRNAs is exceptional, and favors frameshifting (Sundararajan et al. 1999), as it should according to Figure 3, A and B. Indeed, following the finding of earlier experiments (Atkins et al. 1979; Weiss and Gallant 1983), manipulations of the concentration of the corresponding tRNAs should affect frameshifting efficiency. Belcourt and Farabaugh (1990) have shown that overexpression of tRNA<sup>Arg</sup>(3'-UCC-5'), corresponding to the zero-frame codon, reduces Ty1 frameshifting about 40-fold (Fig. 3A). Deletion of the only gene encoding this tRNA increases frameshifting and as a result decreases transposition of Ty1 (Kawakami et al. 1993). Furthermore, overexpression of tRNAs corresponding to the +1-frame codon can increase frameshifting efficiency (Pande et al. 1995) as it should according to Figure 3B. Both these effects are evident in Figure 3, A and B. At the same time, mutations of the last C in the Ty1 frameshifting site (CUU AGG C) to any other nucleotide reduces frameshifting efficiency ~40-fold (Belcourt and Farabaugh 1990). The tRNA that normally recognizes the potential +1-frame codons GGA or GGG, is significantly less abundant than tRNA<sup>Arg</sup>(3'-CCG-5'), which decodes the wild-type +1-frame codon GGC. The codon GGU is presumably recognized by the same tRNA<sup>Arg</sup>(3'-UCC-5'), corresponding to the zero-frame codon, reduces Ty1 frameshifting about 40-fold (Fig. 3A). Deletion of the only gene encoding this tRNA increases frameshifting and as a result decreases transposition of Ty1 (Kawakami et al. 1993). Furthermore, overexpression of tRNAs corresponding to the +1-frame codon can increase frameshifting efficiency (Pande et al. 1995) as it should according to Figure 3B. Both these effects are evident in Figure 3, A and B. At the same time, mutations of the last C in the Ty1 frameshifting site (CUU AGG C) to any other nucleotide reduces frameshifting efficiency ~40-fold (Belcourt and Farabaugh 1990). The tRNA that normally recognizes the potential +1-frame codons GGA or GGG, is significantly less abundant than tRNA<sup>Arg</sup>(3'-CCG-5'), which decodes the wild-type +1-frame codon GGC. The codon GGU is presumably recognized by the same tRNA<sup>Arg</sup>(3'-UCC-5'), but it may have lower affinity because of the G:U pair at the third position of the codon.

This model is also valid for the frameshifting observed during expression of Escherichia coli release factor 2 (RF2), where the efficiency of frameshifting is highly dependent on the concentration of RF2 itself competing for the A-site, UGA, in the zero frame [shift site CUU U(GA)]. However, its precise description in the model shown above is complicated, because the efficiency of frameshifting depends on the rate of Shine Dalgarno/anti-Shine Dalgarno complex formation. This complex formation is important for destabilization of P-site tRNA interactions with the zero-frame codon.

This model indicates that there are certain concentrations of tRNAs that are necessary for the maintenance of accurate triplet decoding.

**Caution about the phrase “simultaneous slippage” for tandem A-site and P-site anticodon repositioning**

All the cases considered so far have involved P-site tRNA repositioning before the A-site tRNA has been irreversibly incorporated into the ribosome. When the A-site tRNA cannot be rejected, a different consideration applies (this arises after the last step of proofreading, during the peptidyl transferase reaction or during translocation as suggested earlier [Weiss et al. 1989], but certainly before translocation is complete). The cellular concentration of tRNA corresponding to the codon in the A-site does not play any role. The success of frameshifting now depends on the ability of the A-site tRNA to be repositioned to a new frame. This kind of frameshifting can occur only in the minus direction, because, as discussed above, the P-site tRNA cannot pair with the codon, 1 nt of which is already occupied by the A-site tRNA. The ability of P-site tRNA to pair with the new codon most probably depends primarily on the energies of the corresponding codon:anticodon complexes in the same manner as during recognition of tRNA in the A-site. However, for A-site tRNA repositioning, the effect of stability of the corresponding complexes is more difficult to estimate.

The anticodon of this tRNA is still located in the A-site, where ribosomal monitoring of the correct geometry of the codon:anticodon dupplex may still occur. This implies that correct interactions at the first two positions in the codon: anticodon dupplex play a bigger role than the correct interactions at the third position. On the other hand, because normally there are no unoccupied nucleotides in the mRNA between the A-site and P-site codons, positioning an A-site tRNA should be more favorable at a codon immediately adjacent to the P-site codon. P-site tRNA repositioning in this case does not necessarily lead to frameshifting. Whether a ribosome will shift depends also on the efficiency of A-site tRNA repositioning.

The classical and important example of such frameshifting is tandem repositioning on the sequence X XXY YYZ. The P-site tRNA that forms the initial complex with XXY can also form a complex with XXX, and the A-site tRNA that recognizes YYZ can also pair with YYY (underlined). Jacks et al. (1988a) suggested that this kind of frameshifting is caused by simultaneous slippage of two tRNAs. This has been interpreted by many to mean relative movement of two tRNAs as a single unit with respect to mRNA. If so, one would expect such movement in the plus direction by the same mechanism, for example, XXX YYY Z. However, no examples of this have been found so far, whereas the number of the examples of minus frameshifting on such sequences is increasing. The hypothesis of simultaneous tandem slippage does not explain the perceived directional selectivity.

In addition, simultaneous slippage of tRNAs presumes something that will coordinate their simultaneous movement with respect to mRNA (and perhaps interactions between these tRNAs). Otherwise, it is not clear what would simultaneously dissociate two codon:anticodon complexes. (Although 3’ stimulatory elements can greatly elevate the level of frameshifting, shifting can occur at a significant level in their absence.) Ironically, before structural data emerged, simultaneous slippage was invoked as an argument in support of the ‘R’-configuration of A- and P-site tRNAs inside the ribosomes, which would allow such interactions between codon–anticodon dupplexes of two tRNAs in contrast to the ‘S’-configuration (Lim 1997). It is well known now that the mutual orientation of tRNAs is closer to the ‘S’-configuration (e.g., Stark et al. 1997; Cate et al. 1999). Therefore, contemporary structural data do not provide any information about possible interactions between A- and P-site tRNA that might be responsible for coord-
nated dissociation of these tRNAs from mRNA. Although it is possible that the ribosome is responsible for coordinated and simultaneous movement of two tRNAs, it is not obvious why such a feature would have evolved in the ribosome. Moreover, there is at present no data supporting such a hypothesis.

**One mechanism of frameshifting versus two**

Why do some cases of frameshifting involve P-site tRNA repositioning that takes place during recognition of a new tRNA in the A-site and other cases involve the repositioning of P-site tRNA at the stage when A-site tRNA is irreversibly incorporated? Is there anything special that determines at what elongation phase repositioning should take place?

The site of frameshifting required for synthesis of the HIV-1 GagPol precursor polyprotein is UUUUUA. Translation studies in reticulocyte lysates show that there are two products. In ~70% of the synthesized polypeptide, the amino acid incorporated after the UUU-encoded Phe is Leu, and this is explicable by tandem repositioning. In the remaining 30% of the synthesized product, there is tandem Phe (Jacks et al. 1988b), and this corresponds to single P-site tRNA repositioning in the posttranslocated complex. (Studies in heterologous translation systems provided further evidence for two different products; Yelverton et al. 1994.) Horsfield et al. (1995) proposed that E- and P-site slippage take place in this case. If E-site slippage indeed occurs in this situation, than it should precede P-site repositioning. Thus, it is likely that repositioning of P-site tRNA in the HIV frameshifting cassette occurs in both elongation phases.

The discrimination between these two phases becomes even more problematic at a very shift-prone heptamer in E. coli, AAAAAG. The explanation for the high frameshifting activity is dependent on E. coli lacking a tRNA<sup>lys</sup>(3'-UUC) that would exclusively recognize AAG. Uridine in the wobble position of tRNA<sup>lys</sup> is modified to 5-methylaminomethyl-2-thiouridine (3'-UUmms<sub>2</sub>U-5'), which pairs poorly with the codon's third base, G. The original model for this frameshifting was that anticodon complex precedes A-site tRNA repositioning. The only difference is that in the case of the "single A-site tRNA slippage hypothesis" (Mejlhede et al. 1999), P-site tRNA is required to rearrange the form of the complex with the same initial codon, but in a different manner.

As originally interpreted, one unusual case of noncognate tRNA-mediated −1 frameshifting is an exception to the model proposed. In E. coli cell-free protein synthesis with the normal balance of tRNAs, phase MS2 RNA directs the synthesis of two forms of its synthetase (viral-encoded replicase component). Synthesis of the longer form is due to tRNA<sup>Thr</sup>(3'-UGG-5') reading a CCA or CGG proline codon and causing a shift to the −1 frame by making the third codon base available for pairing with anticodon base 34 of the following tRNA (Bruce et al. 1986; Dayhuff et al. 1986). The original model for this frameshifting was that anticodon bases 34 and 35 pair with the second and first codon bases, respectively, by implication in the ribosomal A-site (Atkins et al. 1979). However, this explanation seems unlikely. When in the A-site, the third base of the proline codon is likely occluded by tRNA<sup>Thr</sup>(3'-UGG-5') or involved in pairing with anticodon base 33, so that it is unavailable for pairing with anticodon base 34 of the next incoming tRNA. Rearrangement of tRNA<sup>Thr</sup>(3'-UGG-5') in the P-site likely permits pairing by the next tRNA with the third codon base of the "proline" codon (Atkins et al. 2000). Thus, this case is unlikely to be an exception to the model proposed here.
All known cases of frameshifting can be explained within the framework described here. In all cases, rearrangements in the P-site precede slippage of A-site tRNA or its incorporation in a new frame. Rearrangements in the P-site include standard repositioning (but not necessarily completed), dissociation of initial complex, or conformational changes within tRNA. The identity of the incorporated amino acids, direction of the frameshifting, and the effect of tRNA concentrations depend on the elongation stage, where the P-site tRNA rearrangements occur.

There are several different stimulatory signals that can stimulate frameshifting through direct or indirect influence on one or several parameters (stability of P-site tRNAs in the ribosome, affinity of A-site ligands to a codon in the A-site, etc.) described in the present model. These stimulatory signals are highly diverse and include secondary RNA structures, such as pseudoknots (for review, see Brierley and Pennell 2001), stem–loops (Kollmus et al. 1994; Larsen et al. 1997) and RNA three-way junctions (Rettberg et al. 1999).

Some mRNA elements act through interactions with the ribosome (Weiss et al. 1988; Larsen et al. 1994; Li et al. 2001; Stahl et al. 2001). Particular sequence elements flanking the “shiftly” codon affect the affinity of incorporated tRNAs (Bertrand et al. 2002) or release factors (Tate et al. 1995). Among other elements are sequences that influence frameshifting (in this case translational bypass) via the encoded growing nascent peptide chain (Weiss et al. 1990b) and sequences located ~4000 nt downstream of the frameshifting site (Barry and Miller 2002). There are some cis-elements whose nature is unknown (Ivanov et al. 2000), and it is very likely that more new elements will emerge in the future.

CONCLUSIONS

The analysis presented results in several predictions:

1. For frameshifting, tRNA repositioning (or other rearrangements between codon and anticodon) in the P-site always paves the way to repositioning of A-site tRNA or to A-site tRNA binding in a new frame. Therefore, (a) out-of-frame binding of A-site tRNA does not occur without a preceding event in the P-site. (b) Tandem A- and P-site tRNA repositioning is unlikely to occur simultaneously. P-site tRNA rearrangement relative to mRNA should first provide a space for A-site tRNA repositioning.

2. The direction of frameshifting and its sensitivity to the intracellular concentration of tRNAs depends on the elongation phase, in which P-site tRNA repositions (or rearranges with mRNA in another manner).

3. The correct reading frame is controlled by an interplay of the ribosome, tRNAs and mRNAs. Ribosome and tRNAs maintain stable tRNA–mRNA interactions in the P-site. According to the kinetic model in the present work, proper cellular tRNA concentrations in combination with optimal codon usage in translated mRNAs are necessary for reading frame maintenance. Moreover, the effects of the relative concentrations of tRNAs competing for the initial, or new frame, A-site codons are not equivalent according to the kinetic model. Alterations in tRNA concentration of a given organism or expression of heterologous mRNA (with a very different codon bias) should result in elevation of frameshift errors.

ACKNOWLEDGMENTS

We are grateful to Olivier Fayet, Olga Gurvich, Michael Howard, Ivaylo Ivanov, Mikhail Khoudiakov, Knud Nierhaus, Marina Rodina, Petr Sergeiv, Andrey Sivachenko, Daniel Wilson, and particularly to Robert Thompson and Mike Yarus for discussion of the manuscript. This work benefited enormously from the stimulatory effect of others in the field, especially Phil Farabaugh, who are driving the evolution of ideas about the role of P-site tRNAs in framing. This work was supported by NIH (grant R01-GM61200 to R.F.G. and R01-GM48152 to J.F.A) and DOE (Grant DE-FG03-O1ER63132 to R.F.G.).

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Received July 7, 2003; accepted October 9, 2003.

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RNA 2004 10: 221-230

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