Ribosome dynamics during decoding

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Elongation factors Tu (EF-Tu) and SelB are translational GTPases that deliver aminoacyl-tRNAs (aa-tRNAs) to the ribosome. In each canonical round of translation elongation, aa-tRNAs, assisted by EF-Tu, decode mRNA codons and insert the respective amino acid into the growing peptide chain. Stop codons usually lead to translation termination; however, in special cases UGA codons are recoded to selenocysteine (Sec) with the help of SelB. Recruitment of EF-Tu and SelB together with their respective aa-tRNAs to the ribosome is a multistep process. In this review, we summarize recent progress in understanding the role of ribosome dynamics in aa-tRNA selection. We describe the path to correct codon recognition by canonical elongator aa-tRNA and Sec-tRNASec and discuss the local and global rearrangements of the ribosome in response to correct and incorrect aa-tRNAs. We present the mechanisms of GTPase activation and GTP hydrolysis of EF-Tu and SelB and summarize what is known about the accommodation of aa-tRNA on the ribosome after its release from the elongation factor. We show how ribosome dynamics ensures high selectivity for the cognate aa-tRNA and suggest that conformational fluctuations, induced fit and kinetic discrimination play major roles in maintaining the speed and fidelity of translation.

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1. Decoding and recoding

Translation of the genetic information is one of the fundamental processes in living cells. In each round of translation elongation, mRNA triplets are decoded by aminoacyl-tRNAs (aa-tRNAs) which are delivered to the A site of the ribosome in the complex with elongation factor Tu (EF-Tu; in bacteria, or eEF1A in eukaryotes) and GTP. EF-Tu and eEF1A bring all canonical elongator aa-tRNA to the A site where they decode sense codons of the mRNA. However, one uncommon amino acid, selenocysteine (Sec), which is carried by tRNASec, requires a specialized translation factor, SelB in bacteria or its homologue eEFSec in eukaryotes [1,2]. Sec-tRNASec recodes a stop codon UGA in a programmed fashion: in bacterial mRNAs, the stop codon reassigned for Sec is followed by a hairpin structure known as selenocysteine insertion sequence (SECIS) [1]. Recoding of UGA by Sec, as well as of another stop codon UAG by pyrrolysine are bona fide recoding events that occur in the A site of the ribosome. By contrast, other types of ‘recoding’, such as frameshifting and bypassing, appear to be more related to tRNA–mRNA translocation from the A to P and P to E sites or the sliding of the ribosome along the mRNA [3–6]. These phenomena are considered ‘recoding’ in the sense that the information coded by the mRNA is read in an alternative way, but do not involve unconventional codon reading in the A site. In this review, we focus on the EF-Tu- and SelB-mediated decoding/recoding at the A site.

EF-Tu and SelB are translational GTPases that share homologous domains 1–3; SelB has an additional domain 4 that binds to the SECIS [7]. Despite the similar function, the nucleotide-binding properties of EF-Tu and SelB are quite different. EF-Tu binds GDP tighter than GTP and requires a nucleotide exchange factor, EF-Ts, to regenerate the active form of the GTPase [8], whereas SelB has an intrinsically high affinity for GTP and does not need a nucleotide exchange factor.
[9,10]. Both factors in their GTP form interact with their respective aa-tRNAs. Because SelB has evolved to bind a single aa-tRNA, it can recognize Sec and tRNA^Sec by a network of precisely tuned interactions [11–13], resulting in a very high binding affinity, 0.2 pM [9]. By contrast, EF-Tu, which has to bind different aa-tRNAs, employs an affinity compensation mechanism that relies on the relative contributions to binding of the amino acid and tRNA and leads to a uniform overall affinity of about 10 nM regardless of aa-tRNA [14,15].

The main challenge in understanding the decoding process is to uncover the mechanisms that ensure the high fidelity of decoding. Ultimately, the information on which aa-tRNA is cognate at a given cycle of translation elongation is defined by the complementarity between the mRNA codon and the anticodon of the aa-tRNA. However, it has long been recognized that the difference in the stability of correct and incorrect base pairs is insufficient to ensure the observed high fidelity of decoding and, furthermore, even the existing modest discrimination potential cannot be used in full due to the high speed of translation [16–18]. Thus, the complementarity of the codon–anticodon interaction must be sensed by the ribosome in some additional ways. We have shown that the ribosome employs the kinetic discrimination mechanism to select correct aa-tRNA from the bulk of cognate, near-cognate and non-cognate substrates and that the rates of EF-Tu GTPase activation and aa-tRNA accommodation in the A site are greatly accelerated when the anticodon of aa-tRNA matches the codon in the A site [17,19]. However, the structural basis of aa-tRNA selection remained poorly understood and in particular the mechanism of codon-dependent GTPase activation has been unclear for a long time. Pioneering work by Ramakrishnan and co-workers [20,21] has demonstrated how the ribosome recognizes correct codon–anticodon complexes. They showed that cognate codon–anticodon base pairing induces a local conformational change in the decoding centre of the small ribosomal subunit (SSU), where two universally conserved adenes, A1492 and A1493, change their position from the ‘flipped-in’ arrangement, pointing away from the mRNA codon, towards ‘flipped-out’ oriented towards the codon–anticodon complex. The two key adenes—with the help of several other residues at the decoding site—recognize the Watson–Crick geometry of the correct base pairs. The local rearrangement at the decoding centre leads to the global closure of the SSU head and body domains. Conformational dynamics of the SSU is crucial for the GTPase activation of EF-Tu and the subsequent aa-tRNA accommodation in the A site on the large ribosomal subunit (LSU) [22,23], suggesting an important role of ribosome dynamics and induced fit in tRNA selection. Our recent reconstruction of several intermediates on the route to UGA decoding by SelB–Sec-tRNA^Sec finally provides insights into the mechanism of how correct codon–anticodon recognition facilitates GTPase activation [13]. These concepts provide explanations for numerous existing observations and will stimulate new experiments and development of novel approaches to study decoding. In the following, we will summarize the current views on how the ribosome selects cognate aa-tRNA for translation.

2. The path to correct codon recognition
EF-Tu-mediated aa-tRNA delivery to the ribosome proceeds through a number of steps. The initial identification of several discrete states on the EF-Tu pathway was carried out by biochemical methods and ensemble kinetics [19]. Single molecule techniques suggested the existence of additional transient steps and underlined the importance of dynamic fluctuations [24]. The structures of several late intermediates have been solved by X-ray crystallography or cryo-electron microscopy (EM) [25–27]. Molecular dynamics simulations have been successful in modelling the detailed trajectories of motions along the tRNA accommodation pathway [28]. Finally, genetic analysis provided powerful tools to probe the interactions at each state [29–32], leading to a rather consistent picture of what happens during decoding.

The sequence of major steps in aa-tRNA delivery to the ribosome (figure 1) starts with the initial binding of EF-Tu–GTP–aa-tRNA [19,30,36], followed by attempts of the tRNA to read the codon [37,38]. After codon recognition, the GTPase of EF-Tu is activated leading to GTP hydrolysis [33,34,39–41]. GTP hydrolysis in EF-Tu, like in all other translational GTPases, requires docking of the GTP-binding domain onto the sarcin–ricin loop (SRL) of the LSU. The subsequent Pi release [42] and the rearrangement of EF-Tu into the GDP-bound conformation releases aa-tRNA from the factor. The tRNA moves to accommodate itself into the A site of the peptidyl transferase centre, whereas EF-Tu–GDP dissociates from the ribosome [40,43]. Each of these major steps most probably includes additional rearrangements and movements [43,44] which are often too rapid to be resolved. The sequence of events for the eukaryotic analogue of EF-Tu, eEF1A, is probably similar [45,46].

The pathway for the specialized Sec delivery complex, SelB–GTP–Sec-tRNA^Sec, starts with its recruitment to the SECIS. Because SelB–GTP–Sec-tRNA^Sec binds to the isolated SECIS very rapidly and tightly [10], the ternary complex binds to the SECIS even before the translating ribosome has arrived at the SECIS (figure 2). When the ribosome approaches the UGA codon during translation, SECIS-tethered SelB occupies a strategic position across the intersubunit space, blocking the approach to the A site of the release factor 2 (RF2) which normally reads the UGA codon, providing an explanation of why stop codon recognition by SelB does not compete with termination [47]. In the next steps, SelB has to dock onto the SRL of the LSU and hydrolyse GTP. Similarly to EF-Tu, GTP hydrolysis by SelB is necessary to release the tRNA from the factor prior to accommodation and peptide bond formation [9,48]. At all steps on the way to decoding, the ribosome, the elongation factor and aa-tRNA change their positions and conformations, and in the following we will discuss the dynamics of these changes from state to state.

3. Initial binding and codon sampling
The ternary complex EF-Tu–GTP–aa-tRNA is initially recruited by the C-terminal domain of the multimeric ribosomal protein L12 in an mRNA-independent fashion (figure 1) [30,49]. The rate of initial binding is about 100 μM^{-1} s^{-1} measured both by ensemble kinetics and single molecule fluorescence resonance energy transfer (smFRET) techniques [33,38]. From the initial recruitment of the ternary complex to the L12, aa-tRNA must move towards the SSU where it can sample the codon through reversible excursions of the anticodon in and out of the decoding site [33,37]. From smFRET experiments, the position of the tRNA in the...
Cognate (blue) and near-cognate (red) aa-tRNAs at 20°C [17,33–35]. The rate constants of the two irreversible steps, GTP hydrolysis and peptide bond formation, are limited by the respective preceding steps.

**Figure 1.** Schematic of the EF-Tu-dependent aa-tRNA delivery to the A site. The decoding pathway comprises two subsequent selection steps, initial selection and proofreading. The sequence of steps is based on ensemble kinetics and single molecule FRET experiments. Rate constants of kinetically important steps are shown for cognate (blue) and near-cognate (red) aa-tRNAs at 20°C [17,33–35]. The rate constants of the two irreversible steps, GTP hydrolysis and peptide bond formation, are limited by the respective preceding steps.

4. Codon recognition

Codon recognition (figure 1) was originally identified as a distinct kinetic step leading to an aa-tRNA conformational change and a strong stabilization of the cognate aa-tRNA binding in the complex with EF-Tu and the ribosome [34,40,50]. The stabilization is owing not only to the formation of hydrogen bonds in the codon–anticodon complex (and in some cases to interactions with the natural modifications at the tRNA anticodon loop), but is also caused by the interactions of the codon–anticodon complex with the SSU 16S rRNA. When a cognate aa-tRNA binds to the A site, be it either a canonical aa-tRNA in complex with EF-Tu or Sec-tRNA^Sec with SelB, the nucleotides A1492 and A1493 of 16S rRNA change their positions and assume a flipped-out conformation to interact with the second and third positions of the codon–anticodon complex, respectively [13,20] (figures 2 and 3). Because A1492 and A1493 sense the correct Watson–Crick geometry of the base pairs, the local conformational change in the decoding site has a key role in recognition of cognate aa-tRNA.

While it is clear how the cognate aa-tRNA is recognized at the decoding centre of the SSU, it is less clear how structural differences due to mismatches in the codon–anticodon complex result in selectivity and discrimination of incorrect aa-tRNAs. One model is that the near-cognate codon–anticodon complexes are not stabilized to the same degree as the cognate ones and are therefore preferentially rejected. However, comparison of the stabilities of the codon–anticodon complexes in solution [51] and on the ribosome [33,39] suggests that the cognate aa-tRNA is stabilized by about the same factor of approximately 100-fold [19]. Moreover, hydrogen bonds...
formed between the codon-anticodon complex and the elements of the decoding site of the SSU make a very small contribution to aa-tRNA stabilization and their removal has only modest effects on tRNA selection [52]. Inducing the flipped-out conformation of A1492/1493 with the antibiotic paromomycin, which facilitates the local conformational rearrangement in the decoding centre even in the absence of the cognate aa-tRNA [20], results in a 10- to 20-fold stabilization of aa-tRNA in the codon-recognition state and, importantly, both cognate and near-cognate aa-tRNA binding is stabilized [23]. Thus, somewhat counterintuitively, the stabilizing effect of the ribosome is not specific and as such does not contribute

Figure 2. Mechanism of UGA recoding to Sec by SelB–Sec-tRNA^Sec as delineated by cryo-EM. Labels: pep-tRNA, peptidyl-tRNA; d, domains of SelB; dark red shading, GTPase-activated state of SelB; arrows 'open' and 'close', domain opening and closure of the 30S shoulder (sh). States ● ○ ● correspond to structural intermediates resolved by cryo-EM; state ○ was modelled based on the structural data [13]. ● The initial complex contains an mRNA with a UGA stop codon in the A site and a SECIS element exposed for SelB recruitment. The ribosome is in the classical state with a peptidyl-tRNA in the P site and a vacant A site, while the universally conserved bases A1492 and A1493 of 16S rRNA fluctuate between the flipped-in and flipped-out conformations. ● The recruitment state is formed upon binding of SelB–GTP–Sec-tRNA^Sec to the SECIS. The contact between SelB domain 4 and the SECIS is maintained in all subsequent steps. Sec-tRNA^Sec can spontaneously sample a broad range of different conformations. ● In the transient initial binding state, Sec-tRNA^Sec binds to the SRL and SelB to the shoulder domain of the 30S subunit. The shoulder domain moves apart stabilizing A1492 and A1493 in a flipped-in conformation. ● The distance to the UGA codon decreases as tRNASec attempts to read the codon in the transient codon reading state. ● GTPase-activated pre-hydrolysis state. Codon recognition by tRNASec induces a local closure of the decoding centre with A1492 and A1493 flipping out. The resulting global closure of the shoulder domain leads to repositioning of the tRNA and docking of SelB on the SRL. The docking leads to a codon-dependent GTPase-activated state of SelB (dark red shading) and GTP hydrolysis. Notably, SelB, which initially interacts with the 30S shoulder only, does not interact with the SRL until the final GTPase-activated state is reached.

(a)

Figure 3. Conformational rearrangements of the SSU upon UGA codon recognition by Sec-tRNA^Sec (modified from [13]). (a) Local rearrangements of the decoding centre on the SSU. IC, IB, CR and GA are the initial complex prior to SelB-Sec-tRNA recruitment, initial binding, codon reading and GTPase-activated states, respectively. In the IC state, A1492 and A1493 switch between the flipped-in (in) and flipped-out (out) conformations (two-headed arrow); G530 adopts the anti-configuration (anti). (b) Global changes of the SSU. Left: 16S rRNA with landmarks indicated; h44, helix 44 of 16S rRNA. Right: Displacements of the 16S rRNA backbone phosphates in the IB, CR or GA states with respect to the IC as obtained by superposition on 16S rRNA; negative values denote domain opening and positive values domain closure of the 30S subunit.
to the discrimination between cognate and near-cognate aa-tRNAs. Nevertheless, steric complementarity in the decoding centre provides the structural basis for the kinetic enhancement of discrimination by induced fit, as described below.

Correct codon–anticodon interaction induces further conformational changes of the complex, including the domain closure of the SSU and distortion of aa-tRNA. Domain closure entails an inward rotation of the SSU shoulder towards the SSU head and the LSU (figure 3). Kinetically, codon–anticodon recognition, rearrangements in the SSU and aa-tRNA distortion occur at the same time and are probably rate-limited by the adjustment of the codon–anticodon duplex in the decoding site. Based on the comparison of the SSU complexes with the cognate and near-cognate ASLs, Ramakrishnan and co-workers [21] suggested that SSU domain closure is a key determinant of tRNA selection. In the following years, the closed SSU conformation was found in a large number of ribosome-bound structures of EF-Tu with cognate aa-tRNA in pre- and post-hydrolysis states [25–27,53–55]. However, high-resolution structures of near-cognate complexes are not available; therefore, the exact details of how mismatches in codon–anticodon complexes affect domain closure remain obscure. Furthermore, some aspects of the domain closure model were questioned based on crystal structures of 70S ribosomes in complex with full-length tRNAs bound to both P and A sites [54]. In particular, A1493 appeared in the flipped-out position also in the absence of the A-site ligand [54,56]. Such differences are generally difficult to explain, as they may stem from the use of 30S subunit versus 70S ribosomes or the absence or the presence of a tRNA in the P site. The recent cryo-EM structures of ribosome-bound SelB–Sec-tRNA complex argue in favour of the conformational mobility of the two adenines and provide a detailed sequence of conformational changes at the SSU upon decoding [13] (figure 3). As long as the A site is not occupied with aa-tRNA, A1492 and A1493 fluctuate from the flipped-in to flipped-out positions even with a tRNA bound in the P site. One such dynamic conformational state might have been captured in the crystal [54]. In the complexes where SelB–Sec-tRNA^Sec has bound, but codon–anticodon complex is not yet formed, the SSU is in an open conformation with A1492 and A1493 in the flipped-in (that is inactive) conformation (figure 3) [13]; one can hypothesize that these states are also sampled by non-cognate EF-Tu–aa-tRNA complexes attempting to read the codon. Cognate codon–anticodon interaction facilitates or stabilizes SSU domain closure, and notably, the magnitude of the movement (figure 3) is much larger than that described for the isolated SSU [21]. One can envisage that near-cognate complexes do not induce the same closed conformation and are trapped in an open or partially closed state. The cryo-EM structure of the ribosome-bound EF-Tu with a near-cognate aa-tRNA suggests that the global domain closure did not occur [57]; however, the resolution of that structure might be insufficient to draw the final conclusions.

Knowing the structure of near-cognate aa-tRNA on the ribosome would be a key to understanding the ribosome fidelity. This prompted Yusopova, Yusupov and co-workers [54,55,58] to solve the structures of the ribosome–tRNA complexes with diverse mismatches in the codon–anticodon complex in the A site. However, those complexes turned out to represent snapshots of the ribosome’s failure to select a cognate aa-tRNA, rather than pictures of the near-cognate complexes that will be rejected [59]. Because the geometry of such incorrect codon–anticodon complexes is identical to the cognate ones [54,55,58], the ribosome responds by undergoing the same local and global rearrangements as if the complex was a cognate one. The complexes represent those tRNAs that escaped all anticodon complementarity check-points and thus show how incorrect tRNA can mimic the correct one to bypass the discrimination mechanisms of the ribosome [59]. It remains unclear whether such codon–anticodon structures can form during initial selection, as they were prepared in the absence of EF-Tu; rather, the complexes may reflect errors of the proofreading step after the release of EF-Tu. Furthermore, while the efficiency of complex formation during crystallization must be very high, in the cell such complexes are extremely rare, which is evident from the very low measured error frequency, which is between 10^{-3} and 10^{-8} [60,61]. Molecular dynamics calculations also showed that tautomerization as such does not cause high codon reading error frequencies, as the resulting tRNA binding free energies are significantly less favourable than for the cognate complex [62]. Thus, the structures of the near-cognate ribosome–tRNA complexes do not tell us how and why incorrect aa-tRNA are selected. Nevertheless, these structures show that the ribosome recognizes the correct geometry of the complex, rather than the number and positions of hydrogen bonds, and demonstrate that from the structural point of view, decoding errors arise when mismatched codon–anticodon duplexes adopt a Watson–Crick shape, which circumvents the discrimination mechanisms of the ribosome [54,55,58].

5. GTPase activation

The local changes in the SSU not only stabilize the cognate tRNA in the decoding site, but also initiate transitions that lead to the GTPase activation. Until recently, the mechanism of GTPase activation was not clear. We and others suggested that the tRNA plays a crucial role by helping to induce the catalytically active conformation of EF-Tu. This is supported by crystal structures and biochemical and genetic data suggesting that the integrity, structure and rigidity of the tRNA affect the rate of GTP hydrolysis [26,31,63,64]. In principle, the communication between the decoding centre on the SSU and the GTPase centre of EF-Tu or SelB on the LSU could propagate through tRNA, resulting in a series of conformational changes. However, the recent structures of the ribosome–SelB–GTP–Sec-tRNA^Sec complexes suggest a different scenario (figure 4) [13]. We show that codon recognition is communicated by the resulting transition of the decoding site and SSU shoulder domain from an open to a closed conformation which, in turn, moves the tRNA and SelB onto the SRL of the LSU. In the states prior to codon recognition, SelB does not contact the SRL; instead, the SRL interacts with the tRNA, which may even protect the GTPase from premature GTP hydrolysis. GTPase activation occurs upon docking of the GTPase domain of SelB onto the SRL. The docking is assisted by a network of interactions between the SRL and SelB which stabilize the active site conformation of SelB, thereby further facilitating GTP hydrolysis (figure 4) [13]. In the GTP-activated state, the GTP-binding domain 1 of EF-Tu is bound to the SRL, whereas domain 2 is in contact with the shoulder of the SSU [25–27]. The head and shoulder domains of the SSU are in the closed conformation and the tRNA is strongly distorted. The arrangement of the SSU domains, the contacts of domains 1 and 2 with the
SRL and the SSU shoulder and the distortion of the tRNA are very similar for EF-Tu and SelB, suggesting the evolutionary conservation of the GTPase activation mechanism [13].

One remaining question is why the mismatches in the codon–anticodon complex impede GTPase activation. The rate of GTP hydrolysis increases by three to seven orders of magnitude in response to the correct codon–anticodon interaction in the decoding centre [33,34,39,41,65]. Also the smFRET data on decoding suggest that the reactions leading to GTP hydrolysis are more rapid with the cognate than near-cognate aa-tRNA [37,38]. The initial source of the selectivity of GTPase activation must come from the dynamics of the decoding site. This is demonstrated by the finding that inducing the ‘correct’ conformation of the decoding site by paromomycin increases the GTPase rate in the near-cognate ternary complex by a factor of 100, whereas in the cognate complex the rate is even slightly reduced; as a result, near-cognate aa-tRNA is incorporated more efficiently resulting in a higher error frequency [23]. Additionally, the effects of paromomycin on the GTPase activation may be due to its effect on the position of helix 69 of 23S rRNA [54]. The global dynamics of the SSU is crucial, because restricting the conformational flexibility of the SSU by the antibiotic streptolydmycin alters the rates of GTP hydrolysis in such a way that they became almost identical on cognate and near-cognate codons, resulting in virtually complete loss of selectivity [22]. These results, as well as genetic epistatic analysis of ribosome ambiguity mutation (ram) at the h12/S4/S5 region and at the intersubunit bridge B8 [66] are in line with the notion that the SSU domain closure is important. Also the existence of negative regulatory elements in the SSU is consistent with the importance of coordinated, strictly defined motions of the SSU [29]. However, the distortion of the tRNA plays a key role. Mutations in the tRNA D-arm of tRNA^^{Sec} accelerate the rates of GTP hydrolysis independently of codon–anticodon pairing [31]. Mutation A9C tRNA^^{Sec} accomplishes this by increasing tRNA flexibility, whereas G24A tRNA^^{Sec} allows the formation of an additional hydrogen bond that stabilizes the distortion [63]. However, the tRNA is distorted when bound to the ribosome either on cognate or near-cognate codons, as shown both by cryo-EM and fluorescence quenching studies [57,67]; the latter work indicates a somewhat larger distortion of the tRNA in a near-cognate complex. It is difficult to speculate how the structure of the near-cognate GTPase-activated state may look. Codon–anticodon mismatches change the accessible range of global SSU fluctuations, so that EF-Tu does not reach the SRL.

The lower rate of GTP hydrolysis may be a result of a misalignment at the active site or of a lower success rate of the fluctuations from the codon recognition to GTPase-activated state.

### 6. GTP hydrolysis

EF-Tu and SelB have a nucleotide binding pocket which is conserved in the majority of translational GTPases [68]. Two universally conserved residues in the Pro-Gly-His (PGH) motif of the switch II region and in the P loop, His84 and Asp21, respectively, in EF-Tu, are crucial for GTP hydrolysis on the ribosome (figure 5) [69]. By contrast, the slow intrinsic GTPase activity of EF-Tu proceeds through a mechanism that is independent of His84 and Asp21. Instead, a monovalent K⁺ ion coordinated by Asp21 has a small stimulatory effect. Upon binding to the ribosome the mechanism of GTPase reaction changes [69–72]. The key ribosome element responsible for the activation is the SRL of the LSU [26,73]. Ribosomal protein L12 also contributes to the GTPase activation through a yet unknown mechanism [49,74]. The rate of GTP hydrolysis is independent of pH, which indicates that neither His84 nor Asp21 acts as a general base or acid catalyst [69,75]; the kinetic solvent isotope effect is small, suggesting that proton transfer in the transition state is not rate-limiting [69]. Both observations are consistent with the computer simulations of the reaction pathway [70–72]. The predicted pKₐ of His84 is shifted upwards so that the sidechain is positively charged at neutral pH [72]. The most probable role for His84 is thus to contribute to catalysis by positioning of the nucleophilic water molecule for attack, which is triggered by the interaction of EF-Tu with A2662 of the SRL upon codon recognition. The conserved Asp21 may contribute to the acceleration of GTP hydrolysis by providing an optimal orientation of the γ-phosphate. Computer simulations suggest a concerted mechanism of hydrolysis with early proton transfer from water to the γ-phosphate group of GTP, followed by nucleophilic attack by hydroxide ion. His84 and the backbone of the PGH peptide act to ‘pull’ at the γ-phosphate negative charge, while Asp21 acts as to ‘push’ it in the same direction [72]. Thus, GTP hydrolysis is largely governed by the electrostatics of the reaction centre [69–72]. The Asp-bound Mg²⁺ ion is not resolved in EF-Tu, but is clearly seen in the ribosome-bound structures of SelB and EF-G (Asp10 and Asp22, respectively) [13,76,77]. Interestingly, some details of interactions at the GTPase active site appear different in EF-Tu and SelB (figure 5). Crystal structures suggest
that in EF-Tu (and EF-G) the main chain phosphate oxygen of
the SRL A2662 interacts with the NE2 group of His, whereas
in SelB the 2OH of SRL G2661 interacts with ND1 of His.
This does not change the overall mechanism, but may in part
account for differences in the rates of GTP hydrolysis by
EF-Tu and EF-G (more than 100 s$^{-1}$) and SelB (10 s$^{-1}$).

7. Accommodation of aa-tRNA in the A site
Accommodation comprises the movement of aa-tRNA after its
release from EF-Tu into the peptidyl transferase centre on
the LSU [40]. The tRNA CCA end moves by about 100 Å
through the so-called accommodation corridor [28]. Molecular
dynamics simulations suggest that tRNA accommodation
proceeds in three steps, with sequential movements of aa-tRNA
elbow, acceptor arm and then 3'-CCA end entry into the PTC
(figure 6). Unperturbed accommodation is very efficient,
with virtually all of cognate aa-tRNA entering the peptidyl
transferase centre and undergoing peptide bond formation
[39,40]. However, when peptide bond formation is abolished
owing to the presence of a deacylated tRNA in the P site,
only 50% of aa-tRNA reaches the final accommodated step as
estimated by smFRET experiments [38]. Upon accommodation,
aa-tRNA undergoes fluctuations observed as reversible move-
ments of the A-site tRNA elbow region closer to the P-site
tRNA [38,78]. These movements likely reflect the first step of
accommodation with the elbow of tRNA moving into the
close proximity of H89 of LSU [28,44]. Interestingly, smFRET
experiments also report on the second step of accommodation,
which does not lead to a FRET change but could be extracted
from the kinetic analysis of the fluctuations [38]. While the

![Figure 5. The active site of EF-Tu and SelB and the mechanism of GTP hydrolysis. Close up of the GTPase centre of (a) EF-Tu (PDB: 4V5 L) and of (b) SelB (PBD: 5LZD) in the activated state (modified from [13]). (c) and (d) The mechanism of GTP hydrolysis in translational GTPases. Two universally conserved residues, an Asp in the P-loop and a His in the switch II motif contribute to proper positioning of the γ-phosphate of GTP and of the water molecule, respectively. The side chain of the catalytic His is protonated upon contact with the SRL. The role/position of the Mg ion is based on the GTPase-activated pre-hydrolysis structures of EF-G (PDB 4V90, 4JUW, REF) in (c) and SelB (PDB 5LZD [13]) in (d).]

![Figure 6. Model of the aa-tRNA accommodation pathway (from [28]; copyright 2005 National Academy of Sciences USA). Stage 1 represents the hypothetic state after the dissociation of EF-Tu, but before the onset of tRNA accommodation. Stage 2 shows the movement of the aminoacyl-tRNA body (except for the 3'CCA portion of tRNA) towards the PTC. In stage 3, the aminoacyl-tRNA body is accommodated. During stage 4, the aminoacyl-tRNA 3'-CCA end is accommodated into the PTC. The schematics depict the process of accommodation.]
original assignment of this step as peptide bond formation [38] seems unlikely, because all experiments in that work were carried out with a deacylated tRNA in the P site, this step may reflect the accommodation of the acceptor arm and the CCA end. Final adjustment of the CCA end in the PTC may proceed via multiple local pathways owing to the flexibility of the single-stranded CCA end [44].

The biological significance of tRNA accommodation is in providing the mechanism for the aa-tRNA proofreading [39]. When the aa-tRNA is cognate, it is rapidly accommodated [31,33,34,39,79,80]. With a near-cognate aa-tRNA, the rate of accommodation is reduced and the rejection becomes predominant [31,33,34,39,80]. smFRET experiments suggest a similar trend for the accommodation, with the forward fluctuations faster and backward fluctuations slower for the cognate than for near-cognate aa-tRNA [38]. If our assignment of the two accommodation steps observed by smFRET is correct (see above), a 10-fold faster forward movement of the cognate compared to near-cognate aa-tRNA [38] can be interpreted as a crucial role of the acceptor arm accommodation in aa-tRNA discrimination.

Similarly to the initial selection step, the proofreading rates are defined by the codon–anticodon base pairing. The interactions between the cognate codon–anticodon complex and the ribosome elements that resulted in global SSU domain closure during the initial selection step are likely maintained during proofreading, because structures of the complexes in the A/T and A/A states indicate identical interactions in the decoding centre. By contrast, it is less clear whether the rearrangements induced by the near-cognate codon–anticodon complexes are the same or different at the initial selection and proofreading stages; ribosome mutations can differentially affect the aa-tRNA binding stability at the initial selection and proofreading stages [32]. Those near-cognate aa-tRNAs that escaped rejection and have been accommodated into the peptidyl transferase centre adopt geometry mimicking the cognate Watson–Crick base pairs at the first and second codon–anticodon position [54,55,58]. Base pairing at the third codon–anticodon position may adopt unusual geometries, particularly when nucleotide 34 is modified [53,55]. However, the bulk of near-cognate aa-tRNA does not reach the accommodated state. From the structural point of view, the higher rate of near-cognate aa-tRNA rejection may be explained by the reduced stability of the mismatched codon–anticodon complex and the unfavourable geometry imposed by the ribosome, which probably comes at an energetic cost [55]. Whether this leads to the selective destabilization of near-cognate complexes as suggested [55] remains to be tested; as described above, interactions with the decoding site do not selectively destabilize near-cognate codon–anticodon complexes during the initial selection stage [19]. The structural basis for the different accommodation rates of cognate and near-cognate aa-tRNA is not clear, but may result from the misalignment of the near-cognate aa-tRNA [28]. The proofreading step is particularly important when the efficiency of initial selection is low, as it allows reduction of the potentially adverse error of protein synthesis [81]. The remaining errors may be removed by post-transfer editing which employs non-coded termination events [82].

8. Concluding remarks

Structural and kinetic work of the past two decades suggested an important contribution of ribosome dynamics, induced fit and kinetic discrimination in the mechanism of decoding and recoding. Recent breakthrough developments in cryo-EM now allow us to follow the ribosome as it selects aa-tRNA in each round of translation. We can now see more clearly how translational GTPases work and how the ribosome discriminates between the correct and incorrect substrates. Knowing the rate constants of decoding by different aa-tRNAs provides the numbers to build predictive models for translation and to explain the existence of natural translational pauses. The mechanism of stop-codon recoding by Sec emerges in great molecular detail. Understanding the structural basis of tRNA selection will require visualizing the analogous sequence of decoding intermediates for EF-Tu-mediated aa-tRNA delivery. Recent work on deciphering the binding pathway of SelB–Sec-tRNA^[Sec] to the ribosome demonstrates that we can now tackle such complex heterogeneous dynamic systems by both structural and biophysical approaches. The ribosome is a dynamic molecular machine; understanding how spontaneous and induced fluctuations of the ribosome and other translational players are rectified into rapid and accurate translation will answer fundamental questions about the movement and selectivity of molecular ensembles.

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