Coupling of mRNA Structure Rearrangement to Ribosome Movement during Bypassing of Non-coding Regions

Graphical Abstract

Highlights

- A long, non-canonical rotated-state pause of the ribosome is a hallmark of bypassing

- Nascent peptide-ribosome interactions slow down the ribosome prior to the take-off codon

- mRNA structure rearrangements drive ribosome movement across the non-coding gap

- The ribosome scans mRNA a short distance in search of the optimal landing codon

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In Brief
The ribosome can “hop” over a section of phage mRNA while in the midst of translating it, and single-molecule techniques indicate that these dynamics require interactions between the mRNA secondary structure, the nascent peptide, and the ribosome, which advances in a non-canonical rotated state.
Coupling of mRNA Structure Rearrangement to Ribosome Movement during Bypassing of Non-coding Regions

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SUMMARY
Nearly half of the ribosomes translating a particular bacteriophage T4 mRNA bypass a region of 50 nt, resuming translation 3’ of this gap. How this large-scale, specific hop occurs and what determines whether a ribosome bypasses remain unclear. We apply single-molecule fluorescence with zero-mode waveguides to track individual Escherichia coli ribosomes during translation of T4’s gene 60 mRNA. Ribosomes that bypass are characterized by a 10- to 20-fold longer pause in a non-canonical rotated state at the take-off codon. During the pause, mRNA secondary structure rearrangements are coupled to ribosome forward movement, facilitated by nascent peptide interactions that disengage the ribosome anticodon-codon interactions for slippage. Close to the landing site, the ribosome then scans mRNA in search of optimal base-pairing interactions. Our results provide a mechanistic and conformational framework for bypassing, highlighting a non-canonical ribosomal state to allow for mRNA structure refolding to drive large-scale ribosome movements.

INTRODUCTION
Translation normally occurs sequentially in triplets of nucleotides (codons) with strict maintenance by the ribosome of fidelity and reading frame with error rates of 10−3 to 10−4 per codon (Dunkle and Dunham, 2015; Hansen et al., 2003; Jenner et al., 2010). There are cases in which this well-established rule breaks down, where the genetic code can be recoded and altered in an mRNA-specific manner (called “programmed”). During programmed frameshifting, a portion of translating ribosomes can be stochastically diverted to a different reading frame (Chen et al., 2014b; Márquez et al., 2004; Tinoco et al., 2013). Ribosomes can even be directed to bypass, hopping over a stretch of nucleotides to continue translating a contiguous polypeptide (Herr et al., 2000a). These events increase the richness of information encoded in DNA or RNA, where a coding sequence can specify additional protein products not predicted from the standard readout of the open reading frame, as well as adding a layer of translational control.

The best-documented case of programmed bypassing is the gene 60 mRNA of bacteriophage T4 that codes for a subunit of a viral DNA topoisomerase (Herr et al., 2000a; Huang et al., 1988; Weiss et al., 1990). During translation of the gene 60 mRNA, ribosomes translate the first 45 codons (excluding the initiator fMet tRNA, which we term codon 0) to a Gly GGA codon. Half of the translating ribosomes stop at the subsequent UAG stop codon, and the other half skips the next 50 nt and resumes translation from a downstream Gly codon (Maldonado and Herr, 1998). Instead of stopping at the stop codon, the anticodon of the peptidyl-tRNA Gly (Gly-2) (Herr et al., 1999) disengages from mRNA (in a process called “take-off”), the ribosome skips over the 50-nt gap, and the peptidyl-tRNA re-pairs to mRNA downstream at a GGA codon (called the “landing site”). As a result, translation resumes at codon 46 to create a single, continuous protein product from a discontinuous open reading frame (Wills, 2010) (Figure 1A).

Biochemical, genetic, and mutational analysis relying on detection of protein products, both in vitro and in vivo, have identified the essential stimulatory elements for programmed bypassing in gene 60: (1) the tRNA Gly (Gly-2) and the matching GGA take-off and landing sites bounding the non-coding gap, (2) an upstream nascent peptide sequence, (3) a stem loop consisting of the take-off codon and the adjacent UAG stop codon, and (4) a GAG Shine-Dalgarno-like sequence located 6 nt 5’ to the landing site to promote precision of landing (Figure 1A). With the matched take-off/landing pairs, bypassing is the most efficient for the wild-type (WT) GGA codon; other codons are possible, but codons with G or C in the first two positions yield more efficient bypassing (Bucklin et al., 2005). With unmatched take-off/landing pairs, for example, GGA/GCA or GCA/GGA, bypassing efficiencies were greatly reduced (Weiss et al., 1990). The take-off codon is located within a potential −UUCG− hairpin stem loop in the 5’ portion of the non-coding gap, which is
Figure 1. Dynamic Pathways of Gene 60 Bypassing

(A) The elements of the gene 60 bypass are labeled: (1) the UAG stop codon immediately 3’ to the take-off GGA site at codon Gly45, (2) the tRNA_Gly and the matching GGA take-off and landing sites, (3) an upstream nascent peptide signal, (4) a stem loop consisting of the take-off codon, and possibly (5) a GAG Shine-Dalgarno-like sequence located 6 nt 5’ to the landing site to promote precision of landing. The full sequence of the gene 60 mRNA is shown, where the first 42 codons are written as their amino acids (with Met being codon 0), and the remaining sequence is labeled with nucleotides. The coloring of the codon or nucleotide matches the coloring in (B) and (C).

(B) Representative traces of ribosomes Cy3B (green) fluorescent intensity for bypassed and non-bypassed ribosomes. For both cases, there is a phase with normal translation (labeled with a green line), a phase of slowdown (blue line), and either termination at a stop codon for non-bypassed ribosomes or entering a rotated-state pause at codon Gly45 for bypassed ribosomes. The state assignment is shown in red, with the codon counts above.

(legend continued on next page)
important for bypassing: mutations that disrupted base pairing reduced bypassing, whereas compensatory double mutations restored it. Altering the –UUCG– tetraloop sequence at the top of the stem, extending the length of the stem, or increasing loop size also reduced bypassing (Herr et al., 2000b; Weiss et al., 1990; Wills et al., 2008). In addition to the hairpin, a “nascent peptide signal,” KKYKLGQNNRSLKSSS13-29, potentially interacts with the exit tunnel of the ribosome to stimulate bypassing (Herr et al., 2004; Maldonado and Herr, 1998; Weiss et al., 1990). Lastly, there is an alternative landing site at GGG within the non-coding gap near the top of the stem loop (positions 9–11 from the take-off codon); however, the bypassing ribosome always lands at the wild-type landing codon (positions 48–50 from the take-off codon). Thus, it has been proposed that the ribosome does not scan the full non-coding gap in search of a potential landing site, but rather hops over the non-coding region (Wills et al., 2008).

How the ribosome traverses the gap remains unclear, and no definitive and testable model is proposed for the mechanism of such a large-scale movement. What stimulates the ribosome to initiate bypass, and what determines whether or not a ribosome bypasses? What are the roles of the nascent peptide and mRNA secondary structure in inducing bypass? What is the conformational state of the ribosome during bypassing? Prior investigations of frameshifting have underscored the importance of dynamics in translational recoding (Caliskan et al., 2014; Chen et al., 2014b). Here, we probe the dynamic and stochastic nature of bypassing using single-molecule fluorescence to track single translating ribosomes in real time, allowing us to define a global mechanism for bypassing.

RESULTS

Real-Time Observation of Ribosome Bypassing Dynamics

To monitor single Escherichia coli ribosomes progression on mRNAs in real-time, we used zero-mode waveguide (ZMW) instrumentation (Chen et al., 2014a, 2014b). In this study, we followed conformational changes underlying elongation, involving rotational movements of the small (30S) ribosomal subunit with respect to the large (50S) ribosomal subunit and correlated them with binding and departure of tRNAs and elongation factors. To observe rotational movement, the 30S subunit was labeled with Cy3B on helix 44, and a non-fluorescent quencher, BHQ-2, was placed on helix 101 of the 50S subunit, allowing fluorescence resonance energy transfer (FRET) between the two dyes (Chen et al., 2012b, 2013) (Figure S1A).

During one elongation cycle, the two subunits start in a non-rotated state (characterized by high FRET, low Cy3B intensity). The EF-Tu-GTP-aa-tRNA ternary complex (TC) binds to the vacant A site, followed by peptidyl transfer from P-site tRNA to the new A-site aa-tRNA. After peptidyl transfer, the ribosomal subunits rapidly rotate relative to each other (rotated state; lower FRET, higher Cy3B intensity). During this stage, the ribosome is “unlocked,” where the ribosome conformation and tRNA spontaneously fluctuate (Blanchard et al., 2004a; Chen et al., 2012a; Cornish et al., 2008), preparing for translocation. mRNA-tRNA interactions and ribosome-tRNA interactions are weaker at this stage (Liu et al., 2011; Valle et al., 2003). Upon translocation catalyzed by EF-G, the two subunits rotate back to their original high-FRET state and the ribosome is “relocked.” Thus, one round of high-low FRET (low-high Cy3B intensity) corresponds to a single ribosome translating one codon, allowing tracking of translation at codon resolution, and providing the timings of individual substeps at each codon (Chen et al., 2013) (Figure S1B). As opposed to previous smFRET studies with probes labeled at ribosomal proteins S6 and L9 showing spontaneous intersubunit rotations after peptidyl transfer (Cornish et al., 2008), our FRET probe positions possibly monitor a different intersubunit movement that occurs only one cycle per codon. Arrival and departure of the dye-labeled ligands, such as tRNAs, can be simultaneously observed as a sequence of fluorescent pulses (Chen et al., 2013) (Figure S1C). The correlation of single cycles of FRET to translation of a single codon has been confirmed in multiple studies (Aitken and Puglisi, 2010; Chen et al., 2012a; Marshall et al., 2008).

To follow translating ribosomes, we monitored the intersubunit conformational signal upon delivering total tRNA (tRNA tot) ternary complex (aa-tRNA-EF-Tu-GTP), EF-G, and BHQ-50S to immobilized Cy3B-30S preinitiation complexes on the bottom of the ZMWs, as done previously (Johansson et al., 2014; Tsai et al., 2014). Through statistical analysis of multiple translating single ribosomes, we obtained waiting times for the non-rotated and rotated states of each codon. Continuous translation can be observed for more than 50 codons, allowing us to profile the real-time dynamics approaching, during, and after bypassing.

Dynamic Pathways of Bypassing Show a Rotated-State Pause for Bypassed Ribosomes

Translation of the first 40 codons of wild-type gene 60 mRNA proceeds normally, with expected lifetimes of the rotated (waiting for translocation) and non-rotated (waiting for peptidyl transfer) states (2–5 s at 3 μM tRNA tot, TC and 240 nM EF-G), demonstrating a regular elongation rate at these codons. From codons 40 to 45, i.e., before the take-off site, translation gradually slows with an increase in both rotated and non-rotated states lifetimes to roughly 15 s for each state (3- to 7-fold increase). At the bypass site at codon Gly45, an exceptionally long rotated-state pause is observed, with a 20-fold higher mean lifetime of 40 s. For a subset of ribosomes paused at codon Gly45,
translation resumes instead of stopping at the UAG stop codon after Gly45, indicating that we observe bypassing (Figures 1B, 1C, S2A, and S2B).

Translating single ribosomes cluster into three major subpopulations: (1) ribosomes that translate 45 codons and stall at the stop codon; these ribosomes do not bypass and do not exhibit the long rotated pause at Gly45, (2) ribosomes that bypass and translate at least codon 46; these ribosomes ubiquitously exhibit a long rotated state at codon 45, or (3) ribosome traces showing end of Cy3B signal during the long rotated state due to photobleaching or end of movie. Combining the second and third clusters gives a bypassing efficiency of ~35%, consistent with our in vivo assays (33%) and prior studies (Maldonado and Herr, 1998; Samatova et al., 2014). These results also confirm that bypassing is programmed in mRNA itself (Samatova et al., 2014); no other auxiliary factors beyond the standard factors added here are required.

Bypassing and non-bypassing ribosomes show distinct dynamics. All ribosomes exhibit the gradual increase in non-rotated state and rotated lifetimes from codons 40 to 45. This increase in lifetime upon approaching the bypass site is reminiscent of the dynamic signatures observed for nascent peptide-ribosome interactions during SecM stalling (Tsai et al., 2014). The long rotated-state pause at the bypass site (Gly45) is observed only for ribosomes that undergo bypassing and is similar to the non-canonical rotated states observed in –1 frameshifting (Chen et al., 2014b). By parsing into two distinct populations of ribosomes, we obtain a more accurate mean lifetime for the rotated-state pause (88.2 ± 26.4 s) without the convolution of non-bypassed ribosomes (Figure 1D). Resumption of normal translation post bypassing is not immediate, and the ribosome translates slowly for a few more codons before the rotated state lifetimes return to normal (mean lifetime is 5 s), while the non-rotated state lifetimes remain higher (mean lifetime is 15 s) (Figures 1B and 1C; see Figure S3).

The Role of the Nascent Peptide Signal and Its Interaction with the Ribosome Exit Tunnel: Setting the Stage for Bypassing

We hypothesized that the general slowdown in translation observed for both non-bypassed and bypassed ribosomes is due to the nascent peptide signal, KYKLNVRSSKSSS14-29 (Weiss et al., 1990; Wills, 2010), which interacts with the ribosome exit tunnel. In vivo, deleting from codon 14 to codon 29 causes a 70% decrease in bypassing efficiency (Figure 2A). Mutational analysis of the nascent peptide highlighted the importance of a KKYK15-16 motif (Figure 2A). With our in vitro single-molecule system, deleting the sequence encoding KYKLNVRSSKSSS14-29 eliminates observable bypassing and pausing: translating ribosomes no longer exhibit the increase in rotated and non-rotated state lifetimes and ribosomes now translate 29 codons to the stop codon, with only 1% of the traces showing translation beyond the UAG stop codon (Figure S4). Mutating the critical KKYK motif to AAAA resulted in a similar behavior; the increases in non-rotated and rotated states lifetimes are no longer observed (Figure 2B). Deletion of the non-coding gap, while maintaining the nascent peptide signal, abrogates bypassing as expected, but increases in rotated and non-rotated state lifetimes approaching codon Gly45 are observed as for the wild-type sequence (Figure S4). These results indicate that the nascent peptide is responsible for the slowdown approaching the bypass site independent of the mRNA sequence and structure at the bypass site and that this slowdown is necessary for the ribosome to undergo the rotated-state pause at Gly45 for bypassing. Importantly, these interactions are different from SecM-induced stall (Figure 2C).

The Role of the mRNA Hairpin in Promoting Disengagement of Anticodon-Codon Interactions

The nascent peptide signal alone is not sufficient to promote bypass; the hairpin at the bypass site is required. Disrupting the potential base pairing in the hairpin stem abolishes the long rotated-state pause at Gly45, but we still observe the slowdown caused by the nascent peptide signal (Figure 3A). This demonstrates that the hairpin stem loop is required for the long pause in the rotated state prior to bypassing and that the action of the hairpin follows that of the nascent chain.

How the mRNA hairpin promotes such a pause is puzzling, since the –UUCG– hairpin stem loop should be fully melted by the ribosome within the mRNA channel at the take-off site, as the ribosome protects 9 nt subsequent to the P-site codon (Qu et al., 2011). We hypothesize here that the unusual stability of a UUCG tetraloop (Ennifar et al., 2000; Todd and Walter, 2013), which has a propensity to form a compact structure, may favor re-folding of the apical portions of the hairpin, providing a mechanism for the long rotated-state pause. If this hypothesis were correct, then the top portion of the hairpin would be sufficient for pausing and bypassing.

To test this hypothesis, we created two mutants, called Δtop hairpin (destabilized the three base pairs below the UUCG tetraloop) and Δbottom hairpin (leaving the three base pairs below the tetraloop intact but disrupting seven potential base pairs in the lower part of the stem) (Figures 3B and 3C). Translation of the Δtop hairpin mutant mRNA resulted in a decrease of ribosomes that enter the rotated-state pause (12%). Translation of the Δbottom hairpin mRNA remained similar to the wild-type sequence (36% compared to WT 35%). This highlights the importance of the UUCG tetraloop and the top portions of the hairpin in stimulating bypass, consistent with prior mutagenesis (Weiss et al., 1990; Wills et al., 2008) (~1%–30% of WT). Interestingly, destabilizing the three base pairs located 5 nt from the tetraloop did not significantly reduce bypass efficiency (~60%–90% of WT) (Weiss et al., 1990). Thus, the precise location of the UUCG tetraloop with respect to the ribosome during take-off is critical, in addition to the propensity of the tetraloop to re-fold (see the Discussion for speculations on where the hairpin refolds). It is likely that this propensity to re-fold induces a lateral tension on the mRNA–tRNA interaction, which combined with the vertical pull from the nascent peptide interaction, causes the disengagement of the anticyodon-codon interaction and “slippage” uncoupling ribosomal motions from tRNA–mRNA movement, causing the ribosome to be trapped in a non-canonical rotated state, reminiscent of the uncoupled translocation in –1 frameshifting (Chen et al., 2014b).
Second Hairpin 5’ to the Take-off Site Is Required for Bypassing

What provides the forward bias for the bypass? To answer this, we focused on a predicted hairpin 5’ to the bypass stem loop (Figures 4 and S5) (Samatova et al., 2014; Todd and Walter, 2013). We introduced synonymous mutations that disrupt this 5’ stem loop and preserve the amino acid identity and showed that the percentage of ribosomes that enter the rotated-state pause decreases to 11.8%, confirming the importance of this stem loop. Consistently, Samatova et al. (2014) showed that the synonymous mutations that disrupt this 5’ stem loop reduce bypassing efficiency in vitro (~10% of WT), while compensatory mutations partially restore bypassing. When the ribosome is positioned at the take-off Gly45 codon, the 5’ stem loop is likely partially formed, except for the bottom 3–6 base pairs. The refolding of the bottom three base pairs may provide a forward bias for the bypass movement. Alternatively, the directionality may be maintained through re-forming of the stem loop when

Figure 2. Mutation of the Nascent Peptide Interaction Abolishes the Slowdown

(A) In vivo analysis of bypassing with mutants of the nascent peptide. The absolute value of bypassing in these assays by WT (the second from the left) is 33%, and all other values are a percentage of it.

(B) Deleting the key interaction of the nascent peptide signal (KKYK) to AAAA did not increase non-rotated and rotated state lifetimes. Most ribosomes terminate at the stop codon after codon Gly45. An example trace is shown. The color scheme is the same as in Figure 1. n = 424.

(C) In vivo analysis of bypassing with fusions of gene 60/SecM nascent peptides. The cassette used to generate the result in the middle lane has gene 60 sequence encoding amino acids 32 to 46 in its native location 5’ adjacent to the gene 60 take-off codon. The SecM nascent peptide signal encoding sequence is 5’ adjacent to it. The right lane derives from a cassette with the SecM nascent peptide encoding sequence 5’ adjacent to the gene 60 take-off codon. See also Figures S2 and S4.
Figure 3. The –UUCG– Hairpin Stem Loop, Especially the Top Base Pairs, Is Important for the Rotated-State Pause
(A) The hairpin is shown in green, and the UUCG tetraloop is marked in red. To investigate the role of the mRNA hairpin, the base pairs were disrupted; the increase in the non-rotated state lifetime due to the nascent peptide signal is still observed, but a long rotated-state pause at Gly45, characteristic of bypassing, is no longer detected. n = 244.
(B) Mutation of 3 bp below the UUCG tetraloop decreased bypass efficiency to 12%. n = 442.
(C) Mutation of the bottom portion of the hairpin. The bypass efficiency remained the same at 36%. n = 349.
See also Figure S2.
the ribosome vacates the stem loop, acting as a block for backward movement.

**Take-Off and Landing Mechanisms: mRNA Refolding Causes Uncoupled Translocation**

The concerted effects from the nascent peptide interaction, refolding of the 5’ hairpin and the re-folding of the tetraloop, induce a long rotated-state pause characteristic of bypassing. Pausing may be caused by translocation that is uncoupled with the ribosome reverse rotation, similar to what was observed previously for −1 frameshifting in dnaX (Chen et al., 2014b). This leaves the ribosome in a non-canonical rotated state, resulting in the long rotated-state pause observed in frameshifting.

To test whether translocation occurs during the pause, we mutated Asp44 (the codon before Gly45) to Phe; this allows the use of Cy5-labeled tRNA^{Phe} to estimate when translocation occurs during the rotated-state pause (through the departure and disappearance of Cy5-tRNA^{Phe} with the Asp44Phe mutant) in correlation with the Cy3B ribosome conformational signal (Figure S1C). Translocation of the P-site tRNA to the E site is typically correlated with ribosome reverse rotation. Hence, the rotated state lifetime is equivalent to the time to departure of the P-site tRNA signal, and thus, ribosome reverse rotation and translocation are usually coupled. Here, we found the Cy5-tRNA^{Phe} departs 28.1 ± 8.5 s after the rotation of the ribosome at codon Gly45, which is much shorter than the lifetime of the rotated-state pause (88.2 ± 26.4 s) (Figure 5A), indicating that translocation precedes reverse rotation and that the two are now uncoupled. Uncoupled translocation results in a ribosome in a non-canonical rotated state with a peptidyl P-site tRNA and an empty A site. During this non-canonical state, recoding events can occur when ribosome-tRNA-mRNA interactions are weaker to allow for the “take-off” to occur and peptidyl-tRNA and mRNA to dissociate.

Similarly, the timing of “landing” was probed through the Leu46Phe mutant, with the codon after the landing codon mutated to Phe. The arrival of Cy5-tRNA^{Phe} after the intersubunit rotation at Gly45 indicates successful landing of the peptidyl-tRNA to the landing Gly45 codon, with an exposed Phe codon in the A site. The arrival time of Cy5-tRNA^{Phe} in this case is 67.3 ± 13.0 s after the rotation at Gly45. These results allow us to determine the timeline of the hop (Figure 5B)—it begins during the long rotated pause and ends within it.

After translational hopping to Gly46, tRNA^{Phe} arrives at the A-site codon 46 with the ribosome in the rotated state, unlike during normal translation when tRNAs usually bind to the non-rotated state. The binding of the tRNA^{Phe} is stable, with the lifetime comparable to the remaining lifetime of the rotated-state pause. During the remainder of the pause, tRNAs re-pair with the mRNA codon and peptidyl transfer occurs, returning the ribosome to the canonical rotated state with hybrid tRNAs. EF-G can then act on the ribosome and translocate the tRNAs, allowing for normal translation to resume (see Figure S6).

To decipher the mechanism of the bypass during the long rotated-state pause, we examined the effects of simultaneous mutations to the take-off and landing codons, as well as mutations of only the landing codon to create a mismatch (Figure 5C). For unmatched take-off and landing codons, the bypassing efficiency decreases to ~5% of wild-type (Weiss et al., 1990). Upon mutation of the landing codon to a GUA (Val) to create a mismatch, 36% of the ribosomes exhibit a pause at Gly45 with the slowdown approaching Gly45 due to the nascent peptide, similar to wild-type mRNA. Thus, the behavior up to the bypass is not affected by the mutated landing codon. However, only 4% of ribosomes in the landing site mutant resume translation after the pause (within observation window) compared to 67% of for wild-type mRNA.

We determined the fate of ribosomes during the pause by examining the ending state of each ribosome at the pause. For wild-type mRNA, the vast majority of the traces (>90%) show ribosomes either resuming translation or the movie acquisition ends during the pause; for the landing site mutant, the majority of ribosomes (55%) show loss of a Cy3B signal at Gly45. This loss of signal is not due to photobleaching, since for the wild-type mRNA, under the same experimental conditions, only 5% of the traces showed a loss of Cy3B signal at Gly45. Thus, loss of a 30S-Cy3B signal is due to ribosome drop-off on the mutant mRNA, where ribosomes that initiated bypass failed to find the correct landing codon. Since these ribosomes do not stably form peptidyl-tRNA-mRNA interactions, they dissociate from mRNA (Herr et al., 2001). Accordingly, the rotated-state pause lifetime decreased from 40 s to 15 s (Figure 5C).
Figure 5. The Timing and Mechanism of Take-Off and Landing

(A) Using the Asp44Phe and Leu46Phe mutant mRNAs introduced in Figure S2, the timing of bypass was probed. Using the Asp45Phe mutant, we can get the timing of when the Cy5-tRNAPhe (red) departs relative to the start of the rotated-state pause at Gly45. This gives an upper estimate of when translocation occurs (legend continued on next page)
Mutation of both the landing and take-off codons from a GGA to GUA (Val) was previously shown to drop the bypass efficiency to 7% of wild-type (Bucklin et al., 2005). Similar results were observed in our experiments if both the take-off and landing codon are changed from GGA to GUA (Val) (Figure 5E). These results suggest that the identity of the take-off tRNA is not critical to start the process of bypassing; the nascent peptide and hairpins induce take-off and the rotated-state pause. The identity of the tRNA is important for successful landing; it must match the landing codon, but stable G-C-rich pairing is important for successful recognition and re-pairing of the peptidyl-tRNA and mRNA, consistent with earlier data (Bucklin et al., 2005).

Directly Monitoring the Timing of the Hop with Ribosome-mRNA FRET
To probe ribosome movement directly during bypassing, we used FRET between the ribosome and mRNA: mRNA was labeled downstream of the landing site by annealing a Cy5-labeled DNA oligonucleotide complimentary to mRNA (termed +15 Cy5-oligo, 15 nt downstream of the landing GGA codon), and 3OS subunits were labeled with Cy3B on helix 33a near the beak domain, which is close to the mRNA entrance channel (Figures 6A and S1). The bypass will bring the Cy3B dye on the ribosome close to the Cy5 dye with the simultaneous appearance of FRET. Translation is followed by stable binding of Cy3.5-labeled Phe-tRNA^{Phe}. Before bypassing, we observed no FRET between translating ribosomes and downstream labels in mRNA; ribosome-mRNA FRET is thus a hallmark of attempted bypassing.

Using the Asp44Phe mutant and Leu46Phe mutant mRNAs, we can use Cy3.5-labeled tRNA^{Phe} to score for the translation of Phe44 prior to the take-off or Phe46 after successful landing. This allows us to monitor the time between uncoupled translocation (departure of Cy3.5-tRNA^{Phe} from Phe44) and bypassing (appearance of FRET), and also the time between bypassing and successful landing (arrival of Cy3.5-tRNA^{Phe} at Phe46). The hop occurs shortly after uncoupled translocation, on average after 3.4 ± 0.9 s (Figure 6B). The ribosome quickly lands near the landing Gly codon, as demonstrated by the 1–2 frame FRET transition at 100-ms frame rate. After landing, the resume codon in the A site (Leu in the wild-type and Phe in the Leu46Phe mutant) is not immediately available for binding. Instead, Cy3.5-tRNA^{Phe} binds on average 50.5 ± 13.0 s after the hop (Figure 6C).

Does the ribosome land directly on the landing site, or does the ribosome land upstream and scan to find the optimal landing site? To distinguish between these possibilities, we note that the FRET average lifetime for the +15 Cy5-oligo is 72.3 ± 20.0 s. If we move the Cy5-oligonucleotide to 3 nt downstream of the take-off GGA codon (called +3 Cy5-oligo), such that the ribosome footprint is blocked upon landing, the FRET average lifetime decreases significantly to 10.2 ± 4.5 s (Figure 6D). Thus, even when the landing site is blocked, we still see a stable FRET signal, indicating that ribosomes land upstream and then scan before photobleaching or contact quenching the Cy5 dye. Thus, bypassing occurs in two steps: a hop in the 3′ direction, followed by scanning, which is associated with finding the best stable landing site to resume translation.

DISCUSSION
By tracking single ribosomes translating in real time, we delineate here the dynamic events underlying bypassing. All determinants for bypassing are specified by the gene 60 mRNA itself. Translation of the gene 60 sequence results in a branchpoint stimulated by the nascent peptide signal and hairpin. At the take-off codon Gly45, the nascent peptide and the hairpin induce a fraction of the ribosomes (35%) to undergo a long rotated-state pause, similar to what was observed for –1 frameshifting (Chen et al., 2014b). In this state, the ribosome-tRNA-mRNA interactions are weaker, which allows for unusual and large-scale ribosome reconfiguration events to occur for bypassing. Non-bypassed ribosomes terminate at the stop codon without the pause. In this mechanism both the nascent peptide and the hairpin (especially the UUGC tetraloop with three flanking nucleotides) are critical for bypassing. A recent study by Samatova et al. (2014), as well as our findings, confirms the importance of another 5′ stem loop, which provides directionality for the bypass. Here, we propose a model for bypassing that involves the sequential coupling of the re-folding of the two hairpins to ribosome movement, allowing the ribosome with weakened ribosome-tRNA-mRNA interactions induced by the nascent peptide to bypass the non-coding mRNA region.

During the pause. The mean departure time was 28.1 ± 8.5 s, which is a lot shorter than the mean lifetime of the pause (90 s), indicating that the translocation is uncoupled with reverse rotation. This gives an estimate of when the launch occurs.

(B) With the Leu46Phe mutant, A-site accessibility could be probed with Cy5-tRNA^{Phe}, giving an estimate of when landing is completed. The mean arrival time was 67.3 ± 13.0 s, which is also during the pause. Thus, bypass and landing is completed during the rotated-state pause, making the A site available for tRNA binding.

(C) The landing site was changed from GGA(Gly) to GUA(Val), and the mRNA sequence is shown. The increase in the non-rotated state lifetime due to the nascent peptide signal can be seen. The rotated-state pause at Gly45 is shorter than for wild-type. This is due to the lost Cy3B signal during the rotated-state pause, when the ribosome fails to find the correct landing codon after launching the bypass and drops off. Thus, matching take-off and landing codons are required. Consistent with this, the percentage of ribosomes undergoing the rotated-state pause at Gly45 is the same as wild-type. However, the percentage of ribosomes that resume translation (departure of Cy3.5-tRNAPhe from Phe44) and bypassing (appearance of FRET), and also the time between bypassing and successful landing (arrival of Cy3.5-tRNAPhe at Phe46). The hop occurs shortly after uncoupled translocation, on average after 3.4 ± 0.9 s (Figure 6B). The ribosome quickly lands near the landing Gly codon, as demonstrated by the 1–2 frame FRET transition at 100-ms frame rate. After landing, the resume codon in the A site (Leu in the wild-type and Phe in the Leu46Phe mutant) is not immediately available for binding. Instead, Cy3.5-tRNAPhe binds on average 50.5 ± 13.0 s after the hop (Figure 6C).

Does the ribosome land directly on the landing site, or does the ribosome land upstream and scan to find the optimal landing site? To distinguish between these possibilities, we note that the FRET average lifetime for the +15 Cy5-oligo is 72.3 ± 20.0 s. If we move the Cy5-oligonucleotide to 3 nt downstream of the take-off GGA codon (called +3 Cy5-oligo), such that the ribosome footprint is blocked upon landing, the FRET average lifetime decreases significantly to 10.2 ± 4.5 s (Figure 6D). Thus, even when the landing site is blocked, we still see a stable FRET signal, indicating that ribosomes land upstream and then scan before photobleaching or contact quenching the Cy5 dye. Thus, bypassing occurs in two steps: a hop in the 3′ direction, followed by scanning, which is associated with finding the best stable landing site to resume translation.

See also Figures S2 and S6.
The nascent polypeptide of gene 60 causes slowdown in translation as the ribosome approaches the take-off Gly45 codon (from codons 40 to 45), which is a required prelude to bypassing. The interaction causing slowdown begins after a ribosome translates 40 codons when the key KKYK portion of the nascent peptide is ~25 amino acids from the P-site tRNA. The slowdown is defined by the increased lifetimes of both the non-rotated and rotated states, indicating increased barriers to tRNA selection/accommodation and translocation, respectively. These barriers increase progressively during translation from codon 40 to 45. At the take-off site, the KKYK portion of the nascent peptide is ~30 amino acids from the P-site tRNA (as opposed to interaction of SecM, which is 17 amino acids (Nakatogawa and Ito, 2002; Tsai et al., 2014). Thus, even though the dynamic signatures are similar to other stalling sequences, the interaction in bypassing is different from that of SecM; the SecM stalling mechanism does not promote bypassing. We recently showed that co-translational folding of a short peptide sequence upstream of the SecM sequence in the exit tunnel beyond the constriction point “pulls” on the peptide releaving the stall (Nilsson et al., 2015). Since the nascent peptide signal sequence from codons 14–29 in bypassing has been predicted to fold into a α-helical structure (Bhushan et al., 2010; Samatova et al., 2014) (see Figure S7), it may play a similar role in “pulling” on the peptidyl-tRNA to cause the disruption of anticodon-codon interactions necessary for take-off. This suggests that for efficient bypassing, stall is insufficient; the specific interaction and force direction from the traditional SecM stall may not be conducive for bypassing (Goldman et al., 2015). The precise interactions of the nascent peptide with the tunnel will require further study using structural methods. Nonetheless, this nascent peptide interaction is a prerequisite for the ribosome pausing in the rotated state at codon Gly45.

Ribosomes at the bypass site stochastically continue translating or bypass. We propose that the –UUCG– hairpin is the origin for this branchpoint of pathways, similar to the role played by a helical stem loop for frameshifting (Chen et al., 2014b). The role of the bypass hairpin, however, is puzzling, since at the take-off site, the hairpin has been melted by the ribosome. However, the stability of the UUCG tetraloop (Ennifar et al., 2000; Todd and Walter, 2013), which has a propensity to form a compact structure, may cause the apical portion of the hairpin stem loop to refold. In addition, the recent work by Samatova et al. and our work have identified a previously uncharacterized 5′ stem loop that also contributes to bypassing (Samatova et al., 2014). When tRNA<sub>Gly</sub> at codon 45 accommodates into the ribosome and the ribosome rotates, the A site is over codon 45, which places the ribosome such that the UUCG tetraloop is just within the 3′ mRNA channel and the 5′ stem loop is mostly folded except for the bottom base pairs (Figure 7). The tendency for the 5′ stem loop and the UUCG tetraloop to re-fold, in addition to the “pull” on the tRNA through the cascade of nascent peptide
At take-off Gly45 (GGA) codon, ribosome in rotated state after peptide bond formation. The nascent peptide signal interaction pulls on the peptidyl-tRNA, as indicated by the red arrow. The UUCG tetraloop is shown in red. EF-G catalyzes translocation, moving the GGA codon to the P site. Combined with the propensity of the UUCG tetraloop to re-fold, the ribosome slips forward and leads to uncoupled translocation, allowing the UUCG tetraloop to refold. Since the 5' stem-loop blocks backward movement, relaxation of the unstable state threads mRNA in the 5' direction. Refolding of bypass stemloop launches ribosome forward. The ribosome then scans mRNA to find the optimal base pairing, assisted by the GAG Shine-Dalgarno-like sequence and a possible 3' stem-loop. Upon arriving at the landing site, the next tRNA accommodates into the rotated ribosome to help re-define the reading frame and translation is resumed.

See also Figure S7.

Figure 7. A Model of Translational Bypassing
interactions, likely creates a tension in the system. Thus, the refolding of the 5′ stem loop and the ribosome stochastically encountering a folded or unfolded UUUCG tetraloop may cause the initial branchpoint.

EF-G catalyzed translocation occurs, and combined with the 5′ stem-loop refolding, we propose that the ribosome slips forward in the 3′ direction, allowing for the 5′ stem loop to completely refold. This is consistent with the observation that the lower part of the secondary structure is important for bypassing (Wills et al., 2008; Samatova et al., 2014). Simultaneously, the –UUUCG– tetraloop becomes positioned such that it is able to refold. In one model this is within the A site of the ribosome. The folding of a tetraloop or hairpin within the A site is not without precedent: a crystal structure of the 70S ribosome showed mRNA forming a hairpin with a 4 base pair stem and a tetraloop in the A-site, overlapping the natural codon-anticodon interaction region (Yusupova et al., 2001). Along similar lines, a previous model of bypassing suggested that the hairpin re-folds within the A site of the ribosome (Wills et al., 2008). Alternatively, mRNA may be forced a short distance in the forward direction before the tetraloop hairpin forms, perhaps even in the ribosomal E site, with formation of the stem loop it nucleates enhancing forward mRNA to position the ribosome to a more 3′ position on mRNA. Here, we propose that the tetraloop hairpin forms within the A site, though only the top base pairs of the stem are formed.

We further propose that the slip caused by the refolding of the 5′ stem loop and tetraloop uncouples anticodon-codon interactions and translocation from ribosome reverse-rotation. This non-canonical conformation may be hyper-rotated (Qin et al., 2014) or represent a translocation intermediate (Tourigny et al., 2013). The rotated state, with its weakened ribosome-tRNA-mRNA interactions, is key to allowing the mRNA rearrangements that promote bypassing. This ribosome state with a hairpin within the A site may be unstable, and relaxation of this unstable state threads mRNA in the 5′ direction. This forward bias is due to the 5′ stem loop preventing backward movement (Figure 7).

The bypass begins and ends during the long rotated-state pause, with the movement occurring in two steps. First, as soon as the tetraloop clears the ribosome on the 5′ side, the high tendency for the hairpin to refold may cause mRNA to fold directionally in the 5′ direction and the hairpin to fold 5′ of the ribosome. This launches the ribosome forward toward the landing site. However, even with hairpin folding 5′ to the ribosome, the distance threaded is not sufficient to place the ribosome over the landing codon. Thus in the second step, as we have demonstrated, the ribosome scans a short distance to find the optimal landing codon, possibly with the aid of the internal Shine-Dalgarno sequence. This is consistent with the delay between mRNA rearrangement and resumption of translation as measured here.

A combination of mRNA rearrangement-induced movement with processive scanning builds upon and reconciles inconsistencies in earlier models of bypassing (Samatova et al., 2014; Wills et al., 2008). The model proposed here, although still speculative in some aspects, explains many outstanding questions and provides a testable model for future studies. In our model, the re-pairing of the peptidyl-tRNA to the correct position on mRNA during scanning may be stabilized by the SD-like sequence or a possible downstream 3′ stem loop (Samatova et al., 2014); the SD-like sequence has a moderate effect on bypassing but may be important for the fidelity of landing site selection (Herr et al., 2004; Wills et al., 2008). All of these events happen during the rotated-state pause; the majority of the pause is the ribosome sampling and exploring the reading frame widely, with movements possibly similar to the excursions and sliding behaviors observed previously (Koutmou et al., 2015; Yan et al., 2015). In the mechanism proposed here, bypassing is not induced by A-site (UAG stop codon) starvation, explaining why the absence of RF1 did not significantly affect the bypassing efficiency (Herr et al., 2000b). Bypassing induced by A-site starvation may follow a different mechanism (Lindsley et al., 2005a; Lindsley et al., 2005b).

How does the ribosome resume translation? After successful landing and initial contact of the peptidyl-tRNA in the P site with the mRNA codon, the ribosome remains in a non-canonical rotated state with an exposed A site, similar to what was observed for frameshifting. The subsequent tRNA can bind to the ribosomal A site, which may help the ribosome re-define the correct reading frame. Peptidyl transfer in this state is slow, since the rotated ribosome may not position the A- and P-site tRNAs correctly for peptidyl transfer to occur efficiently. Subsequent to peptide bond formation, EF-G can then act on the ribosome and translocate the tRNAs, allowing for normal translation to resume. However, normal rates are not immediately resumed. The nascent peptide is major contributor to this slowdown, suggesting that it still inhibiting subsequent peptidyl transfer and slowing non-rotated state lifetimes until the key sequences leave the ribosomal exit tunnel. This is consistent with the inference from mutagenesis experiments that the nascent peptide signal also has affects at the completion stage (Herr et al., 2000b).

Our results provide a glimpse of an unexpectedly versatile translation scheme with widespread implications. Bypassing may be more widespread than previously thought, suggesting that this phenomenon is not limited to gene 60 (Lang et al., 2014; Nosek et al., 2015). Furthermore, the issue of a fidelity check may be significant for bypassing. Any mismatches upon codon-anticodon re-pairing during reading frame sampling before landing would not be susceptible to the fidelity controls governing proper mRNA decoding (Yan et al., 2015). Lastly, the mechanisms presented here may have parallel in eukaryotic scanning during initiation or other recoding events.

Here, we present a general mechanistic and conformational framework for ribosomal bypassing that may be applicable to different recoding signals. Many aspects of the framework are speculative and still require further investigation, especially the high-resolution structures of the many bypassing intermediates. Nonetheless, a long-lived, non-canonical translational state is the centerpiece of this mechanism and provides a window for reading-frame reset through mRNA structure rearrangement. This state, whose formation is driven by mRNA and nascent chain energy barriers in bypassing, may be universal for many recoding events and possibly a central feature of translational control.
EXPERIMENTAL PROCEDURES

Reagents and Buffers for Translation Experiments

_Escherichia coli_ translation factors (IF2, EF-Tu, EF-G, and EF-Ts) and initiator fMet-tRNA for the single-molecule experiments were prepared and purified as described before (Blanchard et al., 2004b; Marshall et al., 2008). Ribosome purification, tRNA aminocytlation, preparation of biotinylated mRNA, and in vitro bypass assays are described in the Supplemental Experimental Procedures.

All experiments were conducted in a Tris-based polymer mix buffer consisting of 50 mM Tris-acetate (pH 7.5), 100 mM KCl, 5 mM ammonium acetate, 0.5 mM calcium acetate, 5 mM magnesium acetate, 0.5 mM EDTA, 5 mM putrescine-HCl, and 1 mM spermidine. All single-molecule experiments had 4 mM GTP and were performed at 20°C.

Single-Molecule Profiling Experiments

Translation experiments with ribosome Cy3B/BHQ conformational FRET were performed as described (Chen et al., 2014b). Before each experiment, 30S (helix 44 mutant) and 50S (helix 101 mutant) ribosomal subunits (at 1 μM) were mixed in a 1:1 ratio with the 3' dye-labeled oligonucleotides specific for the hairpin extensions in each subunit and incubated at 37°C for 10 min and then at 30°C for 20 min in a polymer mix buffer system. 30S pre-initiation complexes (PICs) were formed as described (Marshall et al., 2008) by incubating the following at 37°C for 5 min: 0.25 μM Cy3B-30S, pre-incubated with stoichiometric S1, 1 μM IF2, 1 μM fMet-tRNA^Met^, 1 μM mRNA, and 4 mM GTP to form 30S PICs in the polymer mix buffer. Before use, mRNA was heated to 90°C for 1 min and then snap cooled to 4°C for 20 min to promote mRNA folding.

Before use, we pre-incubated a SMRT Cell V3 (Pacific Biosciences), a zero-mode waveguide (ZMW) chip, with a 1 mg/ml Neutreadin solution in 50 mM Tris-acetate (pH 7.5), and 50 mM KCl at room temperature for 5 min. The cell was then washed with the Tris-based polymer mix buffer. After washing, 40 μl of the buffer was left in the cell to keep the cell surface wet. We then diluted the 30S PICs with polymer mix containing 1 μM IF2 and 4 mM GTP down to 100 nM PIC concentration. A higher immobilization concentration compared to previous reports was used since not all the PICs have mRNA with biotin (Chen et al., 2014b). The diluted PICs are loaded into the SMRT cell at room temperature for 3 min to immobilize the 30S PICs into the ZMW wells. We wash away unbound material with polymer mix containing 1 μM IF2, 4 mM GTP, 2.5 mM Trolox, and a PCA/PCD oxygen scavenging system (2.5 mM 3,4-dihydroxybenzoic acid and 250 mM protocatechuic deoxy-nase [Atken et al., 2008]). After washing, 20 μl of the washing mix was added to the cell to keep the surface wet.

We formed ternary complexes (TCs) between total charged _E. coli_ tRNAs and EF-Tu/GTP as described (Marshall et al., 2008). Total or Δ(Phe) aminocytlation-TRNA EF-Tu/GTP ternary complexes were pre-formed by incubating (2 min at 37°C) the aa-tRNAs with 5-fold excess of EF-Tu, GTP (1 mM), PEP (3 mM), and EF-Ts (40 μM) in polymer mix. The ternary complexes (3-6 μM) were added to BHQ-50S (200 nM), EF-G (200-800 nM), IF2 (1 μM), GTP (4 μM), 2.5 mM Trolox, and the oxygen-scavenging system to form a delivery mix in polymer buffer. Experiments are done at 3 μM ternary complexes and 240 nM EF-G (chosen to have well-defined, detectable FRET transition signals), unless indicated otherwise. Before an experiment, the SMRT cell is loaded into a modified PacBio RS sequencer. At the start of the elongation experiment, the instrument illuminates the SMRT cell with a green laser and then automatically delivers 20 μl of a delivery mixture onto the cell surface at t ~10 s. Experiments involving labeled tRNAs and ribosome-mRNA FRET were performed similarly. See the Supplemental Experimental Procedures.

ZMW Instrumentation and Data Analysis

All single-molecule fluorescence experiments were conducted using a modified PacBio RS sequencer that allowed the collection of single-molecule fluorescence from individual ZMW wells in four dye channels corresponding to Cy3, Cy5.5, Cy5, and Cy5.5 (Chen et al., 2014a). The RS sequencer had 532- and 632-nm excitation lasers. In all experiments, data were collected at ten frames per second (100-ms exposure time) for 10 min. The energy flux of the green laser was 0.32 μW/μm², and the red laser was at 0.14 μW/μm².

Data analysis for all experiments were conducted with MATLAB (MathWorks) scripts written in-house (see the Supplemental Experimental Procedures). All error bars are SEM.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and seven figures and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2015.10.064.

AUTHOR CONTRIBUTIONS

J.C., J.F.A., and J.D.P. conceived of and designed the experiments. J.C. performed and analyzed the single-molecule experiments. A.C. and M.O'C performed the in vivo experiments. J.C. wrote the manuscript with input from all of the authors.

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