RACK1 on and off the ribosome

ALEX G. JOHNSON,1,2,5 CHRISTOPHER P. LAPOINTE,1,5 JINFAN WANG,1 NICHOLAS C. CORSEPIUS,1 JUNHONG CHOI,1,3 GABRIELE FUCHS,4 and JOSEPH D. PUGLISI1

1Department of Structural Biology, Stanford University School of Medicine, Stanford, California 94305, USA
2Department of Chemical and Systems Biology, Stanford University School of Medicine, Stanford, California 94305, USA
3Department of Applied Physics, Stanford University, Stanford, California 94305, USA
4The RNA Institute, Department of Biological Sciences, University of Albany, Albany, New York 12222, USA

ABSTRACT

Receptor for activated C kinase 1 (RACK1) is a eukaryote-specific ribosomal protein (RP) implicated in diverse biological functions. To engineer ribosomes for specific fluorescent labeling, we selected RACK1 as a target given its location on the small ribosomal subunit and other properties. However, prior results suggested that RACK1 has roles both on and off the ribosome, and such an exchange might be related to its various cellular functions and hinder our ability to use RACK1 as a stable fluorescent tag for the ribosome. In addition, the kinetics of spontaneous exchange of RACK1 or any RP from a mature ribosome in vitro remain unclear. To address these issues, we engineered fluorescently labeled human ribosomes via RACK1, and applied bulk and single-molecule biochemical analyses to track RACK1 on and off the human ribosome. Our results demonstrate that, despite its cellular nonessentiality from yeast to humans, RACK1 readily reassociates with the ribosome, displays limited conformational dynamics, and remains stably bound to the ribosome for hours in vitro. This work sheds insight into the biochemical basis of RPs exchange on and off a mature ribosome and provides tools for single-molecule analysis of human translation.

Keywords: ribosome; translation; human; RACK1; IRES; single-molecule fluorescence

INTRODUCTION

Protein synthesis by the ribosome is a heterogeneous and dynamic process across all kingdoms of life. As such, translation systems are particularly well-suited for single-molecule analysis, given its ability to resolve kinetics and parse out heterogeneous pathways through real-time measurements. Although nearly two decades of studies have clarified and expanded the mechanisms of bacterial translation, very few tools exist to study this process in human systems. A critical absence is an approach for high-efficiency labeling of human ribosomes with fluorescent dyes or other labels with minimal perturbation of ribosomal function or structure.

The eukaryotic ribosome contains ~80 ribosomal proteins (RPs) and four strands of ribosomal RNA (rRNA) organized into large (60S) and small (40S) subunits (Melnikov et al. 2012). The ribosome is often considered a particle of stable composition, and it is thus surprising that nearly 20% of ribosomal proteins (RPs) are nonessential in budding yeast (Steffen et al. 2012), implying that some RP-deficient ribosomes are capable of supporting cellular life. While much less is known about RP essentiality in mammals, preliminary work suggests that several RPs are nonessential within mammalian cell lines (Supplemental Fig. 1A; Blomen et al. 2015). This realization was our gateway into the fluorescent labeling of human ribosomes, wherein we leveraged the cellular nonessentiality of eS25 to genetically engineer a cell line bearing ribosomes labeled at that protein (Fuchs et al. 2015). However, this method has several limitations: In particular it relies on a large protein tag (SNAP-tag) that more than doubles the size of eS25, which may obscure analyses of conformational dynamics and reduce ribosome labeling efficiency through misfolding or proteolysis of the fusion protein. We were further concerned that such a system might artificially perturb or fail to monitor some translation events given the location of eS25 near the ribosomal E site and that sub-populations of cellular ribosomes are sub-stoichiometric for eS25 (Shi et al. 2017; Van De Waterbeemd et al. 2018).

5 These authors contributed equally to this work.
Corresponding author: puglisi@stanford.edu
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To address these issues, our focus turned to another eukaryote-specific RP: receptor for activated C kinase 1 (RACK1). Given its prominent position at the surface-exposed region of the 40S ribosomal subunit (Fig. 1A), we predicted that labeling at RACK1 would be less likely to perturb translation events. Nonetheless, RACK1 has been implicated in diverse biological roles (Gibson 2012), and we were concerned that its labeling might inadvertently alter ribosome function. For one, RACK1 is a WD40-domain protein, a domain seen broadly in cellular signaling proteins and often implicated in a “scaffolding” role (Stirnimann et al. 2010). While RACK1 is nonessential for cellular life from yeast to humans (Jha et al. 2017) (although a full knockout is embryonically lethal in mice [Volta et al. 2013]), its loss or depletion leads to altered cellular signaling (Gibson 2012; Nielsen et al. 2017), translational rewiring (Thompson et al. 2016; Gallo et al. 2018), misregulated ribosome-associated quality control (Juszkwiecz and Hegde 2017; Matsuo et al. 2017; Sitron et al. 2017; Sundaramoorthy et al. 2017), and protection against multiple viruses (Majzoub et al. 2014; Hafirassou et al. 2017; Van De Waterbeemd et al. 2018). However, a notable exception is in Plasmodium falciparum (the malaria parasite) where RACK1 is expressed and required for intracellular survival (Blomqvist et al. 2017), but paradoxically not present in any ribosome structure (Wong et al. 2014; Sun et al. 2015). Thus, RACK1 embodies suggestive qualities of an exchangeable RP, and these properties could be intimately linked with its function.

Here our first goal was to establish a stable and stoichiometric human ribosome labeling site, but given the uncertain stability of RACK1 on the ribosome and the unique opportunity it provided, we sought to harness RACK1 to study the biochemical basis for RP exchange. Although a few studies have demonstrated the possibility for RP incorporation into ribosomes lacking an RP (Lieberman et al. 2004), and some of its properties listed above seem to suggest an extra-ribosomal role (Warner and McIntosh 2009). The potential for RACK1 exchange from the ribosome is also suggested by it being one of the final proteins to incorporate during ribosome biogenesis (Larburu et al. 2016), yet it is also a stoichiometric component of most eukaryotic ribosomes studied to date (Sen Gupta et al. 2004; Rabl et al. 2011; Shi et al. 2017; Van De Waterbeemd et al. 2018). Another notable exception is in Plasmodium falciparum (the malaria parasite) where RACK1 is expressed and required for intracellular survival (Blomqvist et al. 2017), but paradoxically not present in any ribosome structure (Wong et al. 2014; Sun et al. 2015). Thus, RACK1 embodies suggestive qualities of an exchangeable RP, and these properties could be intimately linked with its function.

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RESULTS

RACK1 is organized into a symmetric and well-folded structure when free in solution (Fig. 1A) as revealed by crystallographic studies on proteins expressed in and purified from E. coli (Bjørndal et al. 2003; Nery et al. 2006; Coyle et al. 2009; Ruiz Carrillo et al. 2012). Whereas most other RPs are small and very basic (pl ≈ 10–11),
RACK1 has a relatively large molecular weight and near neutral pI (Supplemental Fig. 1B,C). In our hands, the non-essential RPs eS25 and eL22 may be expressed in E. coli when fused to maltose-binding protein (MBP), but readily precipitate at physiological salt concentrations. Thus, based on its biochemical tractability and cellular nonessentiality, we developed a strategy to fluorescently label RACK1 on and off the ribosome as a strategy to label intact ribosomes and to monitor its ribosome association as a free protein. To ensure a compact labeling strategy, we used site-specific labeling with the 11 amino acid ybbR-tag (Fig. 1A; Yin et al. 2006).

We first tested whether ybbR-tagged RACK1 is functional on ribosomes in cell culture. Using lentiviral transduction, we expressed carboxy-terminally ybbR-tagged RACK1 in a previously generated knockout of RACK1 (ΔRACK1) in human HAP1 cell line (Jha et al. 2017), and examined wild-type, ΔRACK1, and RACK1-ybbR cell lines using cell proliferation and polysome profiling assays. Whereas the ΔRACK1 cell line grows slower relative to the wild-type cell line (Fig. 1B; Supplemental Fig. 2A), expression of RACK1-ybbR in the ΔRACK1 cell line recovered cell proliferation to wild-type levels. To probe its functionality, we examined the three cell lines using polysome profiling with western blot analyses. RACK1 and RACK1-ybbR were present in both monosome and polysome fractions, with minimal free protein detected from the post-nuclear cell lysate (Fig. 1C; Supplemental Fig. 2B). In the ΔRACK1 cell line, we observed an increase in the monosome peak relative to the polysome peaks, possibly indicating alterations in the pool of active ribosomes (Supplemental Fig. 2B). Thus, RACK1-ybbR rescues the ΔRACK1 growth defect and is incorporated into actively translating ribosomes.

**Fluorescent labeling of intact 40S ribosomal subunits via RACK1-ybbR**

RACK1 is a prominent feature on the surface-exposed region of the 40S subunit, where it contacts RPs uS3, uS9, and eS17, and helices 39 and 40 of 18S rRNA, and may facilitate interactions with numerous signaling proteins (Nielsen et al. 2017). It is proximal to the mRNA entry and exit channels (Fig. 2A), and adjacent to the region bound by the HCV IRES, which directly binds the 40S subunit with high-affinity (Johnson et al. 2017). The amino- and carboxy-termini of RACK1 are in close proximity and surface accessible, so the carboxy-terminal ybbR-tag on RACK1 should be accessible to SFP synthase (Yin et al. 2006) on the mature 40S ribosomal subunit (Fig. 2B,C). To test this, we purified ribosomes from RACK1-ybbR expressing cells, identified optimal conditions for labeling (Supplemental Fig. 3A,B), and routinely achieved labeling efficiencies >50% on purified ribosomal subunits.

To determine whether RACK1-ybbR containing 40S ribosomal subunits were biochemically intact, we characterized them alongside wild-type and ΔRACK1 subunits using denaturing and native gel assays. By SDS-PAGE, we observed a single fluorescent band in the 40S-RACK1-ybbR sample that matched the migration pattern of recombinant RACK1-ybbR (Fig. 2D; Supplemental Fig. 3C). This RACK1 band with an increased molecular weight was detected in 40S ribosomal subunits at a level similar to endogenous RACK1 in wild-type 40S subunits by immunoblotting (Supplemental Fig. 3D). As expected, no RACK1 was detected in the ΔRACK1 40S subunits. Several other RPs also exhibited similar levels across the three 40S samples; however, we observed a modest increase in a higher molecular weight band for uS3 in ΔRACK1 relative to wild-type 40S subunits, which was returned to near wild-type levels in 40S-RACK1-ybbR subunits (Supplemental Fig. 3D). The carboxyl terminus of uS3 directly contacts RACK1, and this band may represent a post-translational modification to uS3 that was increased in the absence of RACK1. Finally, using native-gel electrophoresis (Johnson et al.
2018), we found that 40S-RACK1-ybbR subunits bound the HCV IRES with similar efficiency as to WT 40S subunits, and they were competent to form Mg\(^{2+}\)-driven (Lancaster et al. 2006; Yamamoto et al. 2014) 80S complexes on the IRES (Supplemental Fig. 3E). In summary, 40S-RACK1-ybbR ribosomes are nearly identical in composition to wild-type subunits and functional in preliminary biochemical assays.

A reliable smFRET signal for the presence of RACK1 within 40S subunits

Given the high-affinity between the 40S subunit and HCV IRES and its utility for specific immobilization of human ribosomes (Fuchs et al. 2015; Johnson et al. 2018), we next examined the suitability of 40S-RACK1-ybbR ribosomal subunits for single-molecule analyses in the context of the HCV IRES. Domain II of the IRES is predicted to be within smFRET distance with the carboxyl terminus of RACK1, despite there being no direct RNA-RACK1 interaction (Fig. 3A). We therefore used segmental labeling to attach fluorophores directly to the HCV IRES to generate two versions of the IRES with Cy5 dyes at either the base (C44) or elbow (U56) of domain II (Supplemental Fig. 4A). These positions are predicted to be within the dynamic range of FRET (70 and 90 Å away from the carboxyl terminus of RACK1 for C44 and U56, respectively (Fig. 3A; Yamamoto et al. 2014; Quade et al. 2015), and allow an approximate triangulation of conformational dynamics within the 40S-IRES complex.

To detect smFRET between RACK1 and the HCV IRES, we immobilized complexes on a Neutravidin-coated quartz slide and imaged them using total internal reflection fluorescence microscopy (TIRFM) and laser excitation of native RACK1-ybbR-Cy3 on the 40S subunit (Fig. 3B,C; Supplemental Fig. 4B). We observed smFRET to Cy5-IRES (C44) or Cy5-IRES(U56) in 40S-IRES complexes (Fig. 3B). The resulting smFRET signals from both dye pairs were characterized at near- and above-physiological Mg\(^{2+}\) concentrations (2 and 5 mM, respectively) (Fig. 3D; Supplemental Fig. 4C–G) and in a Mg\(^{2+}\)-driven 80S-IRES complex (Lancaster et al. 2006; Yamamoto et al. 2015) (Supplemental Fig. 4H). Most of the 40S-IRES smFRET efficiency distributions were well-fit by single Gaussian functions, irrespective of the IRES labeling position, with a few best fit by a double Gaussian function (Supplemental Fig. 4I). However, there were relatively rare transitions from low to high FRET states for single complexes across all conditions (Supplemental Fig. 4D–H), some of which notably altered the appearance of the FRET efficiency histogram. As expected, the smFRET efficiency was higher when Cy5-IRES(C44) was the acceptor (∼0.28 in comparison to Cy5-IRES(U56) (∼0.17) (Fig. 3D). In both cases, the mean efficiencies were higher than predicted (Supplemental Fig. 4I), likely attributed to the uncertain, dynamic positions of the dyes on the flexible ends of RACK1. Thus, we conclude that RACK1-ybbR is present at its native position on the 40S subunit, and 40S-RACK1-ybbR ribosomes are amenable to single-molecule analysis.

We previously observed conformational dynamics within the 40S-IRES complex at 5 mM Mg\(^{2+}\) that were absent at a lower Mg\(^{2+}\) concentration and in the presence of translation extracts (Fuchs et al. 2015). However, it was unclear whether these measurements were influenced by the bulky tags used for fluorescent labeling: eS25 was carboxy-terminally labeled with a SNAP tag, which is nearly twice its size; and the HCV IRES was labeled at the 5’ end by annealing a 24 nt DNA oligo. We thus completed similar analyses using our improved and much more compact RACK1-IRES smFRET signals for comparison. We observed that the smFRET efficiencies between RACK1 and both IRES constructs were modestly shifted when the Mg\(^{2+}\) concentration was raised from 2 to 5 mM (Fig. 3D; Supplemental Fig. 4). There was a minor population of higher FRET efficiency states (∼0.74) for the Cy5-IRES (C44) acceptor at 5 mM Mg\(^{2+}\) that was absent at the lower Mg\(^{2+}\) concentration. This minor population was also absent in the 80S-IRES complex, which instead had a major population of even higher FRET (∼1.00). Although we currently hesitate to conclude mechanisms from these high FRET states given their rarity and existence at higher-than-physiological Mg\(^{2+}\) concentrations, the observed changes occur at Mg\(^{2+}\) concentrations that are known to promote the transition from the 40S- to 80S-IRES complex (Lancaster et al. 2006), whereby higher Mg\(^{2+}\) concentration may be a surrogate for conformational transitions normally induced by protein factors. Collectively, our new labeling strategy revealed similar conformational dynamics as observed previously using bulkier labels, illustrating that the intrinsic flexibility of the 40S-IRES complex is still observed with a compact labeling scheme.

We subsequently probed the association of 40S subunits with the HCV IRES using the RACK1-IRES smFRET signals in real-time, and compared the measurements to those made with our prior labeling scheme (Fuchs et al. 2015). We applied a zero-mode waveguide-based (ZMW) system to make real-time smFRET measurements (Chen et al. 2014), and selected the IRES(C44) label since it had higher FRET efficiency with RACK1-ybbR in comparison to IRES(U56). To further improve the signal and ensure an unambiguous binding state, we tested both Cy3 and Cy3.5 as the donor fluorophore. The latter has greater distances. We prepared Cy3- and Cy3.5-IRES (C44) RNAs (donors), and native 40S-RACK1-ybbR-Cy5 ribosomal subunits (acceptor), and simultaneously immobilized the labeled IRESs within Neutravidin-coated ZMWs on a single chip with 150,000 ZMWs (RS chip) (Fig. 3E). 40S-RACK1-ybbR at 40 nM was added to the surface of the RS chip after the start of data acquisition, and its
association with a single immobilized HCV IRES was detected via appearance of smFRET (Fig. 3E; Supplemental Fig. 5A). As expected, the Cy3.5–Cy5 pair yielded increased smFRET efficiency (~0.23) relative to Cy3–Cy5 (~0.14) (Fig. 4F; Supplemental Fig. 5B–D), which was lower than that observed using TIRFM, possibly due to quenching of Cy5 by the aluminum ZMWs (Uemura et al. 2010). 40S association times, defined as the time between the start of the data acquisition and appearance of smFRET, and 40S residence times, defined as the length of the smFRET signal, were nearly identical for both Cy3- and Cy3.5-IRES(C44) molecules (Fig. 4D; Supplemental Fig. 5E,F), and similar to our prior measurements of 40S-IRES association kinetics (Fuchs et al. 2015). Thus, by dual-immobilization of Cy3- and Cy3.5-labeled HCV IRESs in ZMWs, we identified an optimal smFRET pair for our single-molecule analyses in ZMWs and validated a strategy that might enable simultaneous tracking of ribosome recruitment to multiple variant RNAs in a single experiment.

**RACK1 association with the 40S ribosomal subunit**

The above experiments established the functionality of RACK1-ybbR in native 40S ribosomal subunits and a

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**FIGURE 3.** RACK1-ybbR is proximal to domain II of the HCV IRES and provides a signal for 40S-IRES association and conformational dynamics on two single-molecule fluorescence platforms. (A) Structural model of the human 40S ribosomal subunit (tan) bound to the HCV IRES (gray), with RACK1 highlighted in green (PDB 5a2q). The carboxyl terminus of RACK1, and the locations of C44 and U56 in the HCV IRES are highlighted, with the measured distance between them indicated. (B) Representative single-molecule fluorescence trace and FRET conversions from TIRFM experiments, with FRET between RACK1-ybbR-Cy3 and IRES(C44-Cy5) highlighted in gray. smFRET was signified by anticorrelated changes in fluorescence intensity of the fluorophores, and single 40S-IRES complexes were identified via a single-step photobleach in the Cy3 donor signal, indicated by the arrow. (C) Schematic of surface immobilized 40S-RACK1-ybbR-Cy3:Cy5-HCV-IRES complexes on a quartz slide used for TIRFM. (D) Cumulative distribution plots of observed FRET intensities for 40S-RACK1-ybbR-Cy3 with IRES(Cy5-U56, purple) or IRES(Cy5-C44, orange) in the presence of 2 mM (solid lines) or 5 mM Mg2+ (dashed lines). The number of traces analyzed (n) for panels D are listed in Supplemental Figure 4I. (E) Schematic of 40S-RACK1-ybbR-Cy5 delivery to surface-immobilized HCV IRES labeled at C44 with Cy3 or Cy3.5 in zero-mode waveguides (ZMWs). The blue oval is the full view of an RS chip that has both IRES(Cy3-C44) and IRES(Cy3.5-C44) immobilized in individual ZMWs on the chip. Below the schematic are example single-molecule fluorescence traces that depict Cy3-Cy5 (left) or Cy3.5-Cy5 (right) upon delivery of 40S-RACK1-ybbR-Cy5 to dual-immobilized IRES(Cy3-C44) and IRES(Cy3.5-C44). The respective FRET events are highlighted in gray, and photobleaching of the donors are indicated by the arrows. (F) Schematic of an individual ZMW showing delivery of 40S-RACK1-ybbR-Cy5 to a Cy3.5 labeled HCV IRES. (G) Cumulative distribution plots of observed FRET efficiency from 40S-RACK1-ybbR-Cy5 FRET with either IRES(Cy3-C44) (green) or IRES(Cy3.5-C44) (yellow) after delivery to dual-immobilized IRESs. The number of traces analyzed (n) for panel G are listed in Supplemental Figure 5D.
smFRET pair between RACK1 and the HCV IRES suitable for real-time measurements. We next adapted these tools with a reconstituted system to study RACK1 flux on and off the ribosome. After expression in *E. coli*, we purified and fluorescently labeled ybbR-tagged RACK1, as well as mutant versions proposed to have reduced association with the ribosome. This included the D107Y mutation (Kuroha et al. 2010), and a phosphomimetic version (S278E) not expected to interfere with ribosome binding (Jha et al. 2017). Of note, the D107Y mutant was less stable in vitro and copurified with *E. coli* HSP60, suggesting a sub-population may be misfolded or unstable (Supplemental Fig. 6). Similarly, a second mutant proposed to disrupt the RACK1-ribosome interaction (R36D/K38E) (Sengupta et al. 2004; Coyle et al. 2009) was also recalcitrant to purification, again likely due to decreased stability (Supplemental Fig. 6). These mutants therefore were unsuitable for single-molecule analysis and suggest revisiting interpretations of the many experiments that have used the mutants to deconvolute ribosomal versus extra-ribosomal roles for RACK1, as previously cautioned (Thompson et al. 2016). To circumvent these issues, we hypothesized that the *S. cerevisiae* RACK1 ortholog, Asc1p, would bind the human ribosome with weakened affinity relative to RACK1. Asc1p has multiple substitutions in comparison to RACK1 along the interaction interface with the ribosome (Supplemental Fig. 7; Nielsen et al. 2017), and the human protein associated more weakly than Asc1p to ribosomes in live yeast (Gerbasi et al. 2004). We therefore purified and fluorescently labeled Asc1p (Supplemental Fig. 6), yielding a stable and potentially lower-affinity comparison to wild-type RACK1. Following a similar rationale, we also

FIGURE 4. Monitoring RACK1 flux by real-time smFRET in ZMWs. (A) Schematic of RACK1 delivery to surface-immobilized ΔRACK1 40S in complex with the HCV IRES labeled at C44 with Cy3.5 in ZMWs. (B) Representative single-molecule fluorescence trace for RACK1-ybbR-Cy5 binding to a single ΔRACK1 40S:Cy3.5-IRES complex indicated by Cy3.5-Cy5 FRET, highlighted in gray. (C) Cumulative distribution plot of observed RACK1 association times from 0 to 60 sec. The labels on the right indicate RACK1-ybbR-Cy5 concentration (nM). The lines represent fits to double-exponential functions. (D) Plot of observed RACK1 association rates ($k_{\text{on}}$) of the fast phase at the indicated concentrations. Error bars represent the 95% confidence interval for each value. The line represents a fit to a linear function, with equation $y = 0.0017x + 0.0149$ and $R^2 = 0.9225$, which yielded the indicated association rate ($k_{\text{on}}$) for the fast phase. The nonzero intercept may be due to experimental factors such as the presence of unlabeled RACK1-ybbR that competes with the labeled protein (RACK1-ybbR had ~70% labeling efficiency). (E) Plot of observed association rates ($k_{\text{on}}$) for the fast phase at the indicated temperatures upon delivery of RACK1-ybbR-Cy5 at 75 nM. Error bars represent the 95% confidence interval for each value. (F) Plot of the observed number of binding events on a single ΔRACK1 40S-IRES complex within a single experiment, indicated by smFRET, for RACK1-ybbR-Cy5 and Asc1p-ybbR-Cy5 delivered at 15 or 75 nM at the indicated temperatures. (G) Native gel electrophoresis analysis of 40S-RACK1-ybbR-Cy5 at 40 nM following competition for 24 h with recombinant unlabeled RACK1-ybbR and Asc1p-ybbR at 20-fold excess, and recombinant RACK1-ybbR-Cy3 at 10-fold excess. The gels were analyzed in parallel and are displayed as an overlay of Cy5 (red) and Cy3 (green) fluorescence. The integrated density of each band was quantified using ImageJ, normalized to the respective control sample (lane 1, from left), and the mean ($n = 3$) Cy5 fluorescence was plotted. Error bars represent standard deviation, and circles represent individual data points. (H) Native gel electrophoresis analysis of 40S-RACK1-ybbR-Cy5 at 40 nM following competition after the indicated times with recombinant unlabeled RACK1-ybbR at 20-fold excess. A representative gel is shown and displays Cy5 fluorescence. Each point on the graph below represents the mean fluorescence intensity remaining relative to the 0.08 h time point, and error bars represent standard deviation. The line represents a fit to an exponential function ($R^2 = 0.93$), which yielded $t_{1/2} = 15 \pm 5$ h. The number of traces analyzed ($n$) for panels C, E, and F are listed in Supplemental Figures 9E, 10C.
attempted to purify the RACK1 ortholog from *P. falciparum*, which is absent from structures of its ribosome, but we were unable to purify the protein for similar reasons to the human mutants.

With a small panel of RACK1 proteins predicted to have a range of affinities with the human ribosome in hand, we next tested whether the different RACK1 proteins associated with wild-type and RACK1-deficient 40S subunits in vitro. We first purified ΔRACK1 40S ribosomal subunits and determined that they bound fluorescently labeled HCV IRES with a similar efficiency to wild-type subunits and were competent in Mg²⁺-driven 80S-IRES formation (Supplemental Fig. 8A,B). These analyses further validated the stability of RACK1-deficient ribosomes and implied that any requirement for RACK1 in HCV IRES activity is independent of mRNA recruitment (Majzoub et al. 2014). Recombinant RACK1 incorporated into ΔRACK1 40S subunits, and it did so far more efficiently than to wild-type 40S subunits (Supplemental Fig. 8C,D). The small amount of RACK1 that incorporated into wild-type ribosomes (∼20% relative to the RACK1-deficient subunits), which presumably have native RACK1 present, likely reflects a small population that lacked RACK1 or exchange of the native protein with the labeled recombinant version over the time course of the experiment. As hypothesized, the D107Y mutant RACK1 failed to incorporate into WT and RACK1-deficient 40S subunits, whereas S278E RACK1 incorporated at similar levels as the wild-type protein in both ribosomes (Supplemental Fig. 8D). Although Asc1p incorporated into ΔRACK1 40S subunits, it failed to incorporate into wild-type ribosomes, supporting a lower-affinity than RACK1 for the human ribosome. Furthermore, in vitro reconstituted 40S-RACK1-ybbR ribosomal subunits were competent to form 80S complexes on the HCV IRES without notable RACK1 dissociation (Supplemental Fig. 8E), contrasting a previous structural study of the complex in which ∼50% of particles lacked RACK1 density (Boehringer et al. 2005).

To confirm that recombinant RACK1 incorporates into the native RACK1 binding site, we examined in vitro reconstituted 40S ribosomal subunits bound to labeled HCV IRES using single-molecule FRET. In complex with HCV IRES(C44-Cy5), in vitro reconstituted 40S-RACK1-ybbR subunits yielded a broad smFRET efficiency distribution with a mean centered at ∼0.31 (Supplemental Fig. 8F,G). This was remarkably similar to the mean intensity observed with native 40S-RACK1-ybbR subunits (∼0.28) (Supplemental Fig. 4I). However, the smFRET efficiency distribution of the reconstituted complex had a broader standard deviation than that of native 40S-RACK1-ybbR (∼0.30 versus ∼0.21), potentially resulting from multiple, heterogeneous FRET states that are difficult to distinguish. These modest differences between native and in vitro reconstituted 40S-RACK1-ybbR subunits may represent differences in post-translational modifications of RACK1 (e.g., amino-terminal acetylation; Van De Waterbeemd et al. 2018) or modifications of other RPs as discussed above (Supplemental Fig. 3D).

We also found that human 40S subunits containing Asc1p-ybbR yielded a smFRET distribution similar to both in vitro reconstituted and native 40S-RACK1-ybbR subunits, albeit with a small population of high FRET states (mean ∼1.00) (Supplemental Fig. 8F,G). Of note, following reconstitution, Asc1p had to be included in the TIRFM imaging buffer to facilitate observation of enough smFRET instances for analysis, which was not required for human RACK1 and preliminarily suggested that Asc1p has a shorter residence time on the human 40S subunit. Thus, our findings demonstrate that recombinant RACK1-ybbR incorporated into its native binding site on the ribosome in vitro and the reconstituted ribosomes behave similarly to those assembled in cells.

Enabled by our reconstituted system, we next measured the kinetics of RACK1 association with ΔRACK1 ribosomes using real-time smFRET analyses in ZMWs. We immobilized preformed complexes of the HCV IRES, labeled at C44 with Cy3.5, and ΔRACK1 40S ribosomal subunits in ZMWs (Fig. 4A). RACK1 association with the 40S-IRES complex was observed via appearance of smFRET (Fig. 4B; Supplemental Fig. 9A). As predicted for a single-step bimolecular association, RACK1 displayed concentration-dependent association times (Fig. 4C; Supplemental Fig. 9B). After fitting data from the 15, 25, 50, 75, and 100 nM RACK1 deliveries with double-exponential functions, we found that both the rates and amplitudes were concentration dependent (Fig. 4D; Supplemental Fig. 9C–E). The fast second-order rate constant was approximately 2 µM⁻¹ sec⁻¹ at 20°C (Fig. 4D), which increased linearly with temperature in a range from 20°C to 37°C (see Materials and Methods for more details; Fig. 4E; Supplemental Fig. 9E–G). The slow second-order rate constant was approximately 20-fold slower (∼0.08 µM⁻¹ sec⁻¹) and plateaued at 30°C (Supplemental Fig. 9C–E). The slow phase might represent a sub-population of RACK1-deficient 40S subunits that inhibits RACK1 reincorporation, perhaps due to a post-transcriptional modification on or near the RACK1 binding site, a conformational rearrangement of the 40S-IRES complex, and/or a missing factor that orients the ribosome in a RACK1-accessible conformation. However, the slow phase could instead represent an experimental artifact arising from a source such as nonuniform mixing across the ZMW chip upon delivery of RACK1-ybbR (Chen et al. 2014). Regardless, the fast association rate we observed for RACK1 incorporation is similar to association rates observed for protein–protein (median ≈ 6.6 µM⁻¹ sec⁻¹) and protein–RNA (median ≈ 6.3 µM⁻¹ sec⁻¹) interactions (Schreiber 2002; Gleitsman et al. 2017), and it provides a useful baseline rate for incorporation of a core RP into mature human ribosomes.

To test the role of RACK1 sequence on ribosomal association and dissociation kinetics, we probed binding of
RACK1 mutants using single-molecule FRET. We first delivered the RACK1 mutant (D107Y) to immobilized ΔRACK1 40S-IRES complexes in ZMWs as above. Consistent with a weaker interaction, the D107Y substitution reduced the rate of RACK1 association (Supplemental Fig. 9H); however, it is due to the reduced stability we observed in vitro (Supplemental Fig. 6), we suspect that this effect could be due to an overestimation of the folded and active RACK1 concentration. We therefore examined yeast Asc1p association with the human ribosome given its in vitro stability and our prior results suggesting a dynamic ribosome interaction. Upon its delivery to immobilized ΔRACK1 40S-IRES complexes, the rate of Asc1p association was dependent on its concentration and temperature (Supplemental Fig. 10A–C). The observed fast association rate for Asc1p at 75 nM (~0.2 sec⁻¹ at 20°C) was similar to the observed RACK1 association rate (~0.2 sec⁻¹) in identical conditions (Supplemental Fig. 10B), as estimated via double-exponential functions (Supplemental Figs. 9E, 10C). However, in contrast to RACK1, we observed multiple Asc1p binding events separated by as little as 1-3 min on a single 40S-IRES complex, which increased with Asc1p concentration and reaction temperature (Fig. 4F; Supplemental Fig. 10D–F). Thus, our real-time data suggest that perturbing the RACK1-ribosome interface alters the stability of the interaction.

We next applied single-molecule and bulk approaches to determine the rate of dissociation of wild-type and mutant RACK1 from ribosomes. Wild-type RACK1 and Asc1p residence times on the 40S-IRES complex, defined as the duration of smFRET events, were both concentration-independent (Supplemental Fig. 11A–C) and dependent on the power of the excitation laser (Supplemental Fig. 11D). Thus, our single-molecule residence time measurements were limited by photobleaching of the dyes, suggesting that they underestimated the lifetime of RACK1-ribosome interactions. To overcome this obstacle, we applied an ensemble approach using native gel electrophoresis. We incubated native 40S-RACK1-ybbR-Cy5 ribosomal subunits with a 20-fold excess of recombinant, unlabeled RACK1-ybbR and quantified the departure of RACK1-ybbR-Cy5 from the ribosome by resolving 40S subunits on a native gel. In this assay, ribosome binding of the unlabeled competitor prevents reassociation of the labeled protein upon its dissociation from the ribosome, yielding a time-dependent decrease in fluorescence signal. After 24 h at 37°C, nearly all of the Cy5 fluorescent signal was lost upon competition with excess RACK1-ybbR (Fig. 4G). Furthermore, there was near-complete exchange of RACK1 upon competition with 10-fold excess RACK1-ybbR-Cy3, indicating that the RACK1 binding site and ribosomal subunits remain intact throughout the experimental window. In contrast, we observed minimal exchange of RACK1 at 20°C or upon competition with 20-fold excess Asc1p at both temperatures. We therefore measured the half-life of RACK1 on the ribosome at 37°C, which we determined to be ~15 h using this bulk exchange assay (Fig. 4H; Supplemental Fig. 11E). Together with the observed fast association rate estimated from the 37°C single-molecule experiments, we estimated an equilibrium dissociation constant of ~10 pM for the RACK1-40S interaction. These results suggest that RACK1 will be stably associated with the ribosome for well over the timeframe of translation in the cell, a process that is typically completed within a few minutes.

**DISCUSSION**

Single-molecule fluorescence strategies are powerful approaches to examine the inherent dynamics of protein synthesis. In this work, we establish RACK1 as a new location to label the human ribosome site-specifically with fluorescent dyes via a compact peptide tag and SFP synthase (Yin et al. 2006). By leveraging the proximity of HCV IRES domain II to RACK1 within the 40S-IRES complex, we applied a smFRET signal to examine the kinetics of the RACK1-ribosome interaction. This integration of biochemical, biophysical, and cell-based approaches revealed the remarkable stability of RACK1 on the ribosome. Unlike prokaryotic systems for which ribosome assembly has been reconstituted (Traub and Nomura 1968; Nierhaus and Dohme 1974), such kinetic parameters of RP-ribosome interactions have remained elusive in humans.

Our initial goal was to engineer human ribosomes for improved labeling with fluorescent dyes amenable to single-molecule analysis. RACK1 was an ideal candidate, as it is dispensable for growth in select cell lines, located between the mRNA entry and exit channels, and proximal to binding sites of translation-related factors and viral IREs. We therefore fused the 11 amino acid ybbR tag onto RACK1, which is among a small group of peptide tags used to label ribosomes in other species (Kaiser et al. 2011; Wang et al. 2012). However, such a short tag has yet to be applied to human ribosomes, and RACK1 is linked to a myriad of translation-related cellular processes. We therefore characterized RACK1-ybbR tagged 40S subunits using orthogonal lines of experiments in cells, and with bulk and single-molecule analyses to ensure their functionality. We found that ribosomes with RACK1-ybbR are functional in cells, biochemically intact, active in simple functional assays, and amenable to single-molecule assays on multiple platforms. In tandem with the segmental-labeling strategy of the IRES, they yielded an unambiguous smFRET signal for the presence of RACK1 within the 40S-IRES complex and enabled us to refine our previous analyses on the dynamics of HCV IRES domain II within the 40S-IRES complex. We also demonstrate their amenability to real-time measurements in ZMWs with complete translation systems. Together, this work highlights the utility of compact labeling schemes for large RNAs and
RACK1 on and off the ribosome

ribbonucleoprotein complexes in single-molecule assays. It also illustrates that ybbR and other small peptide tags are prime candidates for tagging human RPs at their endogenous genomic loci, expanding the candidates for fluorescent labeling to include essential RPs.

Our in vitro experiments collectively demonstrate that RACK1 is a rigid and stable component of the human ribosome. Motivated by the rarity of information on the biochemistry of RP-deficient ribosomes and the rates of in vitro RP exchange (Kossinova et al. 2008; May et al. 2012), we leveraged the smFRET signal described above with ΔRACK1 40S subunits and RACK1-ybbR to monitor RACK1 binding directly to the ribosome. We showed that RACK1-deficient 40S subunits are functional in simple biochemical assays and nearly identical in composition to wild-type ribosomes. Recombinant RACK1 specifically and unambiguously reincorporates into human ΔRACK1 ribosomes at the expected location and we observe limited conformational dynamics. The fast second-order rate constant for RACK1 incorporation is ~4 μM⁻¹ s⁻¹ at 37°C, similar to that of Asc1p incorporation (k₀n ≈ 2 μM⁻¹ s⁻¹ at 37°C), which served as our best available proxy for a stable RACK1 mutant. Using a native gel shift assay, we found that RACK1 has a long half-life on the human ribosome in vitro (t₁/₂ ≈ 15 h), consistent with the expectation from most structural analyses. Substitutions along the RACK1-ribosome interface, as demonstrated by the distance of the distantly related Asc1p, led to multiple dissociation and reassociation events on the 40S subunit over the short time window of our single-molecule experiments. Thus, the high-affinity RACK1-ribosome interaction (K_D ≈ 10 pM) is specified by the slow rate of RACK1 dissociation that is likely mediated by the extensive interface (~1800 Å²) between RACK1 and the ribosome, which includes contacts with three RPs and 18S rRNA (Sengupta et al. 2004; Nielsen et al. 2017).

Both in our single-molecule assays and in cells, the ribosome is often assumed to have infinite stability over its lifetime. Recent cellular studies have challenged this paradigm and have proposed that variations in the composition of the ribosome can yield specialized functions (Gilbert 2011; Xue and Barna 2012). To overcome the intrinsic stability of the RACK1-ribosome interaction that we observed, however, regulatory control via RACK1 ribosome occupancy would require hours for its stochastic departure from the ribosome or active mechanisms to release RACK1. Indeed, our findings are consistent with the possibility that the ribosome-bound and small pool of free RACK1 can exchange stochastically over the lifetime of a ribosome (t₁/₂ ≈ 5–7 d) (Dice and Schimke 1972; Nikolov et al. 1987; Defoiche et al. 2009; Mathis et al. 2017). Such exchange has been proposed for other RPs and may take place during viral infections in bacteria (Mizuno et al. 2019) and neuronal functions of mammals (Shigeoka et al. 2018). On the other hand, rapid changes to mature ribosomes could be achieved through the activated release of RACK1, or another RP, perhaps triggered by covalent modifications or other ribosome rearrangements. The RP eS10 may be an example of such a phenomenon, as it has been observed at sub-stoichiometric levels in the human ribosome (Shi et al. 2017; Van De Waterbeemd et al. 2018) and is a target for regulatory ubiquitination (Sundaramoorthy et al. 2017; Juszkiwicz et al. 2018). Our in vitro findings therefore provide an initial kinetic benchmark for the stability of an integral RP on mature human ribosomes, lending biochemical insight for complementary cellular investigations on the biological consequences and timescales of RP exchange.

The results presented here establish RACK1 as an advantageous location to install a compact fluorescent label on the human ribosome. We demonstrate the utility of this labeling position by defining baseline kinetic parameters for RACK1 association with the ribosome, revealing the remarkable stability of the interaction. We therefore expand the toolkit available for single-molecule studies required to reveal the kinetics that govern human translation and its regulation (Sokabe and Fraser 2018), and provide a biochemical benchmark for the flux of RPs on and off the mature ribosome.

MATERIALS AND METHODS

Molecular cloning

To isolate human RACK1 cDNA for cloning, total RNA from HeLa cells was extracted using TRizol (ThermoFisher, cat.# 15596026), reverse transcribed and amplified with gene-specific primers, transformed into TOP10 cells, and validated by Sanger sequencing. To make the RACK1-ybbR lentiviral expression construct, RACK1-ybbR was amplified and inserted into pENTR/D-TOPO (ThermoFisher) and then cloned into the lentivirus expression plasmid pLenti CMV Puro DESt (w118-1) (Addgene) by Gateway cloning (ThermoFisher) and validated by sequencing. The open-reading frames (ORFs) for RACK1 proteins were inserted into a pET28c-6xHis-MBP bacterial recombinant expression plasmid backbone downstream from MBP using a construct derived from a human eIF1A expression construct (Fraser et al. 2007). The ORF from human RACK1 was subcloned from the lentivirus expression construct, the ORF for P. falciparum RACK1 (PIRACK1) was subcloned from P. falciparum asexual stage cDNA from the D10 strain (a gift from Michael Boucher, Ellen Yeh laboratory), and the ORF for S. cerevisiae Asc1p was subcloned from a cloned cDNA construct purchased from GenScript (ORF clone OSi04112D). At the 3’ end of each construct was sequence encoding for the ybbR peptide tag and a stop codon (GATTCTCTTTATATGGCGATGAAGTTCCGAGTA). We refer to the proteins encoded by these plasmids as 6xHis-MBP-RACK1/Asc1p-ybbR fusion proteins. The D107Y, R38D/K40E, and S278E amino acid substitutions were introduced into human RACK1 using the QuickChange II XL site-directed mutagenesis kit (Roche). All plasmids are available upon request.
Cell growth

HAP1 cells were grown at 37°C with 5% CO₂ in Iscove’s Modified Dulbecco’s medium (IMDM) (Gibco, cat.# sh30228) supplemented with 10% (v/v) fetal bovine serum, 2 mM l-glutamine (Gibco), and 1X penicillin/streptomycin (Gibco). As indicated, wild-type and ΔRACK1 (clone E3-AS) HAP1 cells were used in our studies (Carette et al. 2011; Jha et al. 2017).

Lentiviral transductions

Lentivirus packaging was conducted either at the Neuroscience Gene Vector and Virus Core of Stanford University or performed in-house using pLenti constructs by cotransfection with ΔVPR, VSV-G, and pAdVAntage packaging plasmids into HEK293FT cells using FuGENE HD (Promega) (Campeau et al. 2009). Cells were transduced with lentivirus-containing media 24 h after seeding in six-well plates at 250,000 cells per well. Lentivirus-containing media was replaced with fresh media after 24 h, and an additional 24 h, transduced cells were selected by the addition of 1 µg/mL puromycin. Following 5 d under puromycin selection, surviving cells were expanded in culture for characterization and ribosome purifications.

Polysome profiling

HAP1 cells were grown by routine passaging and cells were brought to ~80% confluence in 150-mm culture dishes, receiving fresh IMDM media 4–6 h before harvesting. Immediately before harvest, 100 µg/mL cycloheximide (CHX) was added to media, and cells were incubated for 3 min at 37°C. Media was removed by aspiration, adherent cells were washed twice with ice-cold PBS (containing 100 µg/mL CHX). PBS was aspirated from cells, which were subsequently dissociated from plates by scraping and transferred to Eppendorf tubes in residual PBS. An equal volume of 2X cell lysis buffer (30 mM Tris-HCl pH 7.5, 150 mM NaCl, 2% Triton X-100, 10 mM MgCl₂, 2 mg/mL heparin, 0.2 mg/mL CHX, and 2 mM DTT) was mixed with cell suspensions, and lysis proceeded for 10 min on ice. Lysate was clarified to remove nuclei at 8400g for 5 min at 4°C with a microcentrifuge, and lysates were measured with a nanopod to normalize gradient inputs by absorbance at 260 nm. Using 14 × 89 mm thin-wall polypropylene tubes (Beckman Coulter ref. 331372), 10%–60% sucrose gradients were prepared in 20 mM Tris-HCl pH 7.5, 150 mM NaCl, 5 mM MgCl₂, 0.1 mg/mL CHX, and 1 mM DTT, and linearized using a BioComp 107 Gradient Master. Post-nuclear lysate samples from a single 150-mm dish (500 µL of ~1000 ng/µL RNA) was loaded onto each prechilled sucrose gradient and centrifuged at 150,000g (35,000 rpm) using a SW41 Ti rotor for 2 h 45 min at 4°C. Gradients were fractionated with a Brandel gradient fractionation system using a 5 mm UV cell detector, pumping at 0.75 mL per minute, and collecting fractions across the length of the gradient. Protein was extracted from each 0.75 mL fraction by methanol extraction. Briefly, 0.1 mL of each fraction was mixed with 0.4 mL methanol, vortexed, and spun for 10 sec at 9000g in a microcentrifuge. To each sample, 0.1 mL chloroform was added, followed by additional vortexing. Next, 0.3 mL nanopore H₂O was added to each fraction with vortexing for phase separation. The upper phase was discarded, and 0.3 mL was added to the lower phase with more vortexing. The protein was then pelleted by centrifuging at max speed in a microcentrifuge for 2 min, from which the supernatant was removed, and the pellet was further dried using a SpeedVac. Dried protein pellets were resuspended in 0.01 mL 8 M urea/ 100 mM Tris-HCl pH 8.0. Solubilized proteins were then used for SDS-PAGE and western blotting.

Proliferation assay

Proliferation assays were performed as described previously (Johnson et al. 2018). Briefly, wild-type, ΔRACK1, and ΔRACK1 + RACK1-ybbR HAP1 cells were dissociated from culture plates with trypsin and counted for live cells by trypsin blue. Cells were then seeded into 96-well plates at 10,000 live cells/well with 10 replicates per cell line, assaying over 4 d of growth using the Vybrant MTT Cell Proliferation Assay Kit (ThermoFisher). Absorbance values at 570 nm were determined using a Synergy Neo2 instrument (BioTek).

Immunoblotting

Western blotting was performed as described previously (Johnson et al. 2018). When reprobing blots, HRP-conjugated secondary antibodies were either inactivated by incubation with sodium azide in 5% skim milk/TBST or stripped with Restore Stripping Buffer (ThermoFisher). The following antibodies were used in this study: anti-RACK1 (Cell Signaling, ref. 4716S, 1:1000), anti-RACK1 (Santa Cruz, ref. 17754, 1:500), RPL5 (Genetex, ref. 101821, 1:1000), uS10/RPS20 (abcam, ref. 133776, 1:1000), eS10/RPS10 (Genetex, ref. 101836, 1:1000), eS5/RPS2 (Santa Cruz, ref. 130399, 1:500), and uS3/RPS3 (Bethyl, ref. 303-840A, 1:1000).

Ribosome purifications

40S and 60S ribosomal subunits from wild-type, ΔRACK1, and ΔRACK1 + RACK1-ybbR HAP1 cells were purified as described previously with a few optimizations (Fuchs et al. 2015). A total of 20–40 × 150-mm culture dishes were grown to ~80% confluency. Media was removed, cells were washed with ice-cold PBS, and PBS was removed via aspiration. In the residual PBS, cells were scraped to release them from the dish, combined into a 15 mL culture tube, and pelleted by centrifugation at 1000g for 10 min in an Eppendorf 5810 R centrifuge equipped with a swinging bucket rotor. Cell pellets were resuspended in ice-cold PBS and repelleted via centrifugation to wash. PBS was removed via aspiration, and cell pellets were resuspended in an equal volume of fresh, ice-cold PBS (typically 1–2 mL). An equal volume of 2X lysis buffer (30 mM Tris-HCl pH 7.5, 300 mM NaCl, 20 mM MgCl₂, 2% (v/v) Triton-X 100, 4 mM DTT, 2 mg/mL heparin, and 1X Complete Mini protease inhibitor cocktail (Roche)) was added, cells were mixed gently by multiple inversions of the culture tube, and cells were incubated on ice for 10 min to complete lysis. Cell lysates were cleared of debris and nuclei via centrifugation at 42,000g (10,000 rpm) at 4°C for 10 min in a Fiberlite F21 rotor. The supernatant was layered directly onto a high-salt sucrose cushion (20 mM Tris-HCl pH 7.5, 500 mM KCl, 30% (v/v) sucrose, 10 mM MgCl₂, 0.1 mM EDTA pH 8.0, and 2 mM DTT).
were pelleted via centrifugation at 63,000 g for 16–18 h at 4°C using a Type 80 Ti rotor (Beckman Coulter). Ribosome pellets were dissolved in resuspension buffer (20 mM Tris-HCl pH 7.5, 500 mM KCl, 7.5% [v/v] sucrose, 2 mM MgCl₂, 75 mM NH₄Cl, 2 mM puro-mycin, and 2 mM DTT) by gentle pipetting. To complete resolubilization and splitting of the ribosomal subunits, ribosomes were incubated at 4°C for 1 h followed by an incubation at 37°C for 1.5 h. To isolate 40S and 60S ribosomal subunits, the solution was layered directly onto a prechilled linear 10%–30% sucrose gradient (with 20 mM Tris-HCl pH 7.5, 500 mM KCl, 6 mM MgCl₂) in 25 × 89 mm polypropylene centrifuge tubes (Beckman Coulter, ref. 326823) prepared using the BioComp 107 Gradient Master. Gradients were centrifuged at 49,123g for 16 h at 4°C using an SW32Ti rotor. Gradients were fractionated with a Brandel gradient fractionation system using a 1 mm UV cell detector, pumping at 1.5 mL per minute, and collecting 0.75 mL fractions across the length of the gradient. Fractions containing 40S and 60S ribosomal subunits as identified via absorbance at 254 nm were pelleted via centrifugation at 63,000g for 20 h at 4°C using a Type 80 Ti rotor (Beckman Coulter). Ribosomal subunits were resuspended in storage buffer (30 mM HEPES-KOH pH 7.4, 100 mM KOAc, 5 mM Mg(OAc)₂, 6% [v/v] sucrose, and 2 mM DTT) aliquoted, flash frozen in liquid N₂, and stored at −80°C. Ribosome concentrations and labeling efficiencies were determined by measuring absorbance at 260 nm (total ribosome concentration) and at 548 or 646 nm (Cy3 or Cy5 dyes, labeled ribosome concentration) using a NanoDrop and the appropriate extinction coefficients. Ribosomal subunits were characterized by SDS-PAGE (Bio-Rad, cat.# 7322010) equilibrated in TEV buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% [v/v] glycerol, 10 mM imidazole, and 5 mM β-mercaptoethanol). Dye-labeled proteins and ribosomes were characterized as indicated by SDS-PAGE (Bio-Rad Any KD precast gels), fluorescent scanning using a Typhoon instrument, the Pierce Silver Stain Kit (ThermoFisher), western blotting, and mass-mapping with the Stanford PAN facility.

**SFP synthease labeling of ybbR peptides**

SFP synthease was expressed recombinantly in bacteria using the pET29-Sfp construct (a gift from Jun Yin at Georgia State University) (Yin et al. 2006). Protein expression was performed as described, and protein was subsequently used to catalyze ybbR labeling reactions. Sulfo-Cyanine3 (cat. # 11380) and Sulfo-Cyanine5-maleimide (cat. # 13380) were purchased from Lumiprobe. Coenzyme A trilithium salt (CoA) was purchased from Sigma-Aldrich (cat.# C3019). CoA was reacted with maleimide-dyes as described, and CoA-dyes were purified using a 5%–50% gradient of acetonitrile in water with 0.1% TFA using an HPLC equipped with a C18 column (Yin et al. 2006). Successful synthesis and purification of CoA-dye products was verified by LC-MS analysis. After confirming the published synthesis and purification scheme, we instead used the crude CoA-dye reactions without HPLC purification, by quenching excess maleimide with DTT and directly labeling ybbR-tagged proteins with SFP synthease.

40S ribosomal subunits containing RACK1-ybbR were fluorescently labeled with Cy3 or Cy5 by incubating 40S subunits with 2–4 µM excess SFP synthease and CoA-Cyanine dyes at 37°C for 2 h in ybbR-labeling buffer (50 mM HEPES-KOH pH 7.5, 100 mM NaCl, 10 mM MgCl₂, 10% [v/v] glycerol, and 1 mM DTT). To remove free dye and SFP synthease, labeled 40S subunits were layered onto a low-salt sucrose cushion (30 mM HEPES-KOH pH 7.5, 100 mM KOAc, 0.5 M sucrose, 5 mM Mg(OAc)₂, and 2 mM DTT), and centrifuged at 287,000 g (90,000 rpm) for 1 h at 4°C in a TLA100.2 rotor in 11 × 34 mm thick-wall polycarbonate ultracentrifuge tubes (Beckman, ref. 343788). Ribosome pellets were washed once and subsequently resuspended with ribosome storage buffer, aliquoted, flash frozen in liquid N₂, and stored at −80°C. Recombinant proteins were labeled with cyanine dyes by incubating 10 µM 6xHis-MBP-RACK1-ybbR proteins with 2 µM SFP synthease and 15 µM CoA-Cy3/5 dyes at 37°C for 2 h. Free dye was removed via purification over 10DG-desalting columns (Bio-Rad, cat.# 7322010) equilibrated in TEV buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% [v/v] glycerol, 10 mM imidazole, and 5 mM β-mercaptoethanol). Dye-labeled proteins and ribosomes were characterized as indicated by SDS-PAGE (Bio-Rad Any KD precast gels), fluorescent scanning using a Typhoon instrument, the Pierce Silver Stain Kit (ThermoFisher), western blotting, and mass-mapping with the Stanford PAN facility. Protein concentrations and labeling efficiencies were determined by measuring absorbance at 280 nm (total protein concentration) and at 548 or 646 nm (Cy3 or Cy5 dye, labeled protein concentration) using a NanoDrop and the appropriate extinction coefficients.

**RACK1 recombinant protein expression and purification**

Recombinant 6xHis-MBP-RACK1-ybbR fusion proteins were expressed and purified from E. coli BL21(DE3) cells. Two to four liters of cells were grown at 37°C in LB supplemented with 50 µg/mL kanamycin to OD₆₀₀ of 0.5, and cells were rapidly cooled on ice. Expression of the fusion proteins was induced by addition of 0.5 mM IPTG, and cells were maintained in a shaking incubator for 16 h at 17°C. Cells were harvested by centrifugation at 5000g for 10 min at 4°C in a Fibertite F9 rotor.

Cells were lysed by sonication in lysis buffer (20 mM Tris-HCl pH 8.0, 300 mM NaCl, 10% [v/v] glycerol, 10 mM imidazole, and 5 mM β-mercaptoethanol), and lysates were cleared by centrifugation at 38,000 g for 10 min at 4°C in a Fibertite F21 rotor followed by filtration through a 0.22 µm syringe filter. Clarified lysate was loaded onto a Ni-NTA gravity flow column equilibrated in lysis buffer, washed with 20 column volumes (CV) of lysis buffer, 20 CV of wash buffer (20 mM Tris-HCl pH 8.0, 1 M NaCl, 10% [v/v] glycerol, 25 mM imidazole, and 5 mM β-mercaptoethanol), and 10 CV of lysis buffer. Recombinant proteins were eluted with six sequential CV of elution buffer (20 mM Tris-HCl pH 8.0, 300 mM NaCl, 10% [v/v] glycerol, 250 mM imidazole, and 5 mM β-mercaptoethanol), and fractions with recombinant protein as identified by SDS-PAGE analyses were dialyzed overnight at 4°C into ybbR-labeling buffer (50 mM HEPES-KOH pH 7.5, 100 mM NaCl, 10 mM MgCl₂, 10% [v/v] glycerol, and 1 mM DTT) or TEV Buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 10% [v/v] glycerol, 10 mM imidazole, and 5 mM β-mercaptoethanol). Fluorescently labeled and unlabeled proteins were dialyzed overnight in the dark at 4°C into TEV buffer in the presence of 1–1.5 mg of TEV protease. To separate 6xHis-tagged proteins from RACK1-ybbR(Cy3/5), the proteins were passed through a second Ni-NTA gravity column equilibrated in TEV buffer to capture 6xHis-MBP, TEV protease, and SFP synthease. Flow-through containing RACK1-ybbR(Cy3/5) was further purified using size exclusion chromatography using a Superdex 75 column (23 mL).
equilibrated in SEC buffer (20 mM HEPES-KOH pH 7.5, 150 mM KOAc, 10% [v/v] glycerol, and 1 mM DTT). Fractions containing RACK1-ybbR(Cy3/5) were concentrated using a 10 kD MWCO Amicon Ultra centrifugal filter, aliquoted, flash frozen on liquid N₂, and stored at −80°C. The concentration was determined via absorption at 280 nm using a nanodrop for total protein, and at 548 or 646 nM for Cy3 or Cy5 labeled proteins, respectively. A concentration range of indicated recombinant proteins was analyzed by thermal melt assays in SEC buffer using SYPRO Orange (Molecular Probes, cat.# S6651) using a Bio-Rad CFX96 Touch Real-Time PCR C1000 System (ramping from 4°C to 100°C at 1°C per minute).

RNA transcription and fluorescent labeling

For single-molecule experiments, HCV IRES RNA was produced and labeled with cyanine dyes as described previously (Johnson et al. 2018). Briefly, an acceptor RNA was transcribed in vitro by T7 with GMP-priming for segmental labeling. Synthetic RNA donor oligos with cyanine dyes installed terminally at C44 or internally at U56 (TriLink) were ligated to the acceptor RNA using T4 RNA ligase 1 (NEB) (Fig. 3; Supplemental Fig. 4). For TIRFM experiments, RNAs were labeled at C44 with Cy5-hydrazide and for ZMW experiments, RNAs were labeled with cyanine dyes as described previously (Johnson et al. 2018). Briefly, 2.75% acrylamide (37.5:1) 0.5% Nusieve GTG agarose composite gels were prepared in the following buffer: 25 mM Tris-OAe pH 7.5, 4 mM KOAc, 2 mM Mg(OAc)₂, 0.5 mM DTT, 2.5% glycerol, 0.1% [v/v] TEMED, and 0.1% [v/v] fresh ammonium persulfate. The gels were cooled at 4°C for 20 min and allowed to further polymerize at room temperature for 2 h. Gels were prerun in ice-cold running buffer [25 mM Tris-OAe pH 7.5, 4 mM KOAc, and 2 mM Mg(OAc)₂] for 30–60 min at 4°C. All native gel electrophoresis assays were repeated at least three times, and respective ribosome, protein, and RNA samples were derived from a single purification. For gel shift assays with HCV IRES, the indicated amounts of purified ribosomal subunits were incubated for 10 min at 37°C with RNA in ribosome assay buffer [30 nM Cy5-labeled IRES in 30 mM HEPES-KOH pH 7.4, 100 mM KOAc, and 2 mM Mg(OAc)₂]. For analysis of recombinant protein incorporation into purified ribosomal subunits, 40 nM of purified recombinant proteins labeled with Cy5 were incubated for 10 min at 37°C with the indicated amount (titration experiment) or 75 nM (single concentration experiment) of 40S ribosomal subunits in ribosome assay buffer. For analysis of 80S ribosome complex formation, 60 nM of the indicated 40S and 60S ribosomal subunits, and 30 nM of Cy5-labeled IRES were incubated for 10 min at 37°C in ribosome assay buffer with the indicated amount of MgCl₂.

Native gel electrophoresis

Ribosomes and complexes of ribosomes and HCV IRES were analyzed using acrylamide/agarose composite gels as described previously (Johnson et al. 2018). Briefly, 2.75% acrylamide (37.5:1), 0.5% Nusieve GTG agarose composite gels were prepared in the following buffer: 25 mM Tris-OAe pH 7.5, 4 mM KOAc, 2 mM Mg(OAc)₂, 0.5 mM DTT, 2.5% glycerol, 0.1% [v/v] TEMED, and 0.1% [v/v] fresh ammonium persulfate. The gels were cooled at 4°C for 20 min and allowed to further polymerize at room temperature for 2 h. Gels were prerun in ice-cold running buffer [25 mM Tris-OAe pH 7.5, 4 mM KOAc, and 2 mM Mg(OAc)₂] for 30–60 min at 4°C. All native gel electrophoresis assays were repeated at least three times, and respective ribosome, protein, and RNA samples were derived from a single purification. For gel shift assays with HCV IRES, the indicated amounts of purified ribosomal subunits were incubated for 10 min at 37°C with RNA in ribosome assay buffer [30 nM Cy5-labeled IRES in 30 mM HEPES-KOH pH 7.4, 100 mM KOAc, and 2 mM Mg(OAc)₂]. For analysis of recombinant protein incorporation into purified ribosomal subunits, 40 nM of purified recombinant proteins labeled with Cy5 were incubated for 10 min at 37°C with the indicated amount (titration experiment) or 75 nM (single concentration experiment) of 40S ribosomal subunits in ribosome assay buffer. For analysis of 80S ribosome complex formation, 60 nM of the indicated 40S and 60S ribosomal subunits, and 30 nM of Cy5-labeled IRES were incubated for 10 min at 37°C in ribosome assay buffer with the indicated amount of MgCl₂.

Single-molecule spectroscopy

For equilibrium experiments, 100 nM refolded HCV IRES annealed to a biotinylated DNA oligo was incubated with 200 nM of 40S(RACK1-ybbR-Cy3) ribosomal subunits for 10 min at 37°C in ribosome assay buffer. Conditions were identical for experiments with 80S ribosomes except that 400 nM 60S ribosomal subunit was included and ribosome assay buffer was supplemented with 3 mM Mg(OAc)₂ (5 mM total). For experiments with in vitro reconstituted ribosomes, 1 µM of ΔRACK1 40S ribosomal subunits were first incubated with 2 µM of Cy3-labeled recombinant protein for 10 min at 37°C in ribosome assay buffer, and subsequently, 200 nM of in vitro reconstituted 40S ribosomal subunits were incubated with refolded HCV IRES for 10 min at 37°C in ribosome assay buffer. Ribosome-IRES complexes were immobilized in a channel of neutraVin-coated quartz slides by addition of 40 µL 400 pM complex (based on the IRES concentration) and incubating for 5 min. Unbound complexes were washed from the channel using ribosome assay buffer supplemented with 2 mM Trollox (TYS), 2 mM protocatechueic acid (PCA), and 0.06 U/µL protocatechuate-3,4-dioxygenase (PCD). Complexes were imaged using a prism-based total internal reflection fluorescence (TIRF) microscope as described previously (Atkén et al. 2008), using 532 and 647 nm excitation lasers with movies collected at five frames per second and EM gain set to 650. Short movies (50 frames) were collected using either (single) or both (dual) excitation lasers, whereas long movies (1500 frames) were collected using excitation with the 532 nm laser. Colocalized molecules were identified and analyzed using custom MATLAB (R2017a, MathWorks) scripts (available upon request).
For real-time experiments in ZMWs, all imaging was conducted using a modified Pacific Biosystems RS II DNA sequencer as described previously (Chen et al. 2014; Johnson et al. 2018). ZMW chips purchased from Pacific Biosciences (PacBio) were initially washed with TP50 buffer (50 mM Tris-OAc pH 7.5, 100 mM KCl), coated with neutravidin by incubating 20 µL of 75 nM neutravidin in TP50 supplemented with blocking oligo for 5 min, and washed 3–6× with ribosome assay buffer. 40S-IRES complexes were formed by incubating 1 µM ΔRACK1 40S ribosomal sub-units with 200 nM refolded HCV IRES labeled at C44 with Cy3.5 annealed to a biotinylated DNA oligo. Complexes were immobilized on the neutravidin-coated ZMW surface by incubating 40 µL of 1 nM of complex (via the IRES) for 5 min. Unbound complex was removed via 3–5 washes with ribosome assay buffer. Immediately before imaging, buffer was removed from the surface of the ZMW chip and replaced with 20 µL ribosome assay buffer supplemented with 2 mM TSY, 2 mM PCA, and 0.06 U/µL PCD. Upon start of image acquisition, 20 µL of Cy5-labeled recombinant protein in the same buffer was delivered to the chip surface. For experiments with delivery of 40 nM 40S-RACK1-ybbR-Cy5, all conditions were the same except 40 µL of 0.5 nM HCV IRES labeled with either Cy3 or Cy3.5 at C44 was immobilized on the ZMW surface. All movies were collected at five frames per second for 20 min to maximize the capture of single-molecule photobleaching events of the Cy3.5 dye. Unless noted, the 532 nm laser was used at 0.8 W/µm² illumination and the temperature of the imaging chamber was 20°C. The “low laser power” experiment was conducted using the 532 nm laser at 0.32 W/µm² illumination at 20°C. All movies were analyzed using custom MATLAB scripts (available upon request) to extract fluorescence intensity and kinetic parameters. The respective ribosome, protein, and RNA samples were derived from a single purification for all equilibrium and real-time single-molecule analyses.

For the single-molecule analyses, n represents the number of molecules that were analyzed, unless otherwise noted. For robust kinetic fits, we typically analyzed >175 molecules per condition, and only wells with a single IRES molecule, identified by a single-step photobleaching event of the donor fluorophore. To overcome nonuniform mixing on the surface of the ZMW chip, which occurs within 1 sec at the center and is complete at ∼7 sec, we concentrated our analyses on molecules present in the center of the chip (Chen et al. 2014). At the lowest concentration (5 nM), the slow rate was dominant, and we were unable to confidently fit the association times with single or double exponential functions. For all analyses that compare smFRET efficiencies, smFRET events were assigned to a single state and single molecules were identified by a single-step photobleaching event of the donor fluorophore, based on relative fluorescence intensities. We then corrected the background fluorescence present in each optical channel and calculated distributions of observed smFRET efficiencies for the respective dye pairs.

The 30°C and 37°C real-time association experiments were conducted with the PacBio RS II imaging chamber set to 28°C and 35°C, respectively, to account for the increased temperature due to laser excitation. We delivered RACK1 at 75 nM for several reasons. First, the fast association rate was dominant, which enabled robust fitting of the data. Second, evaporation of the sample is a concern at the highest temperature in our microscope. We thus focused our measurements on the impact of temperature on fast binding events, which predominately occurred within 30 sec.

Third, this concentration was within a window that would allow observation of faster or slower binding events.

DATA DEPOSITION

The single-molecule data sets generated during and/or analyzed during the current study are available from the corresponding author upon request. All custom MATLAB analysis scripts are available upon request or are available on GitHub (https://github.com/corsepius/FRETAutomation).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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RACK1 on and off the ribosome

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