

HHS Public Access

Author manuscript *Nature*. Author manuscript; available in PMC 2016 December 15.

Published in final edited form as: *Nature.*; 534(7608): 558–561. doi:10.1038/nature17978.

Rocaglates convert DEAD-box protein eIF4A into a sequenceselective translational repressor

Shintaro Iwasaki¹, Stephen N. Floor¹, and Nicholas T. Ingolia^{1,†}

¹Department of Molecular and Cell Biology, Center for RNA Systems Biology, University of California, Berkeley, CA 94720 USA

Abstract

Rocaglamide A (RocA) typifies a class of protein synthesis inhibitors that selectively kill aneuploid tumor cells and repress translation of specific mRNAs¹⁻⁴. RocA targets eukaryotic initiation factor 4A (eIF4A), an ATP-dependent DEAD-box RNA helicase; its mRNA selectivity is proposed to reflect highly structured 5' UTRs that depend strongly on eIF4A-mediated unwinding⁵. However, rocaglate treatment may not phenocopy the loss of eIF4A activity, as these drugs actually increase the affinity between eIF4A and RNA^{1,2,6}. Here, we show that secondary structure in 5' UTRs is only a minor determinant for RocA selectivity and RocA does not repress translation by reducing eIF4A availability. Rather, *in vitro* and in cells, RocA specifically clamps eIF4A onto polypurine sequences in an ATP-independent manner. This artificially clamped eIF4A blocks 43S scanning, leading to premature, upstream translation initiation and reducing protein expression from transcripts bearing the RocA-eIF4A target sequence. In elucidating the mechanism of selective translation repression by this lead anti-cancer compound, we provide an example of a drug stabilizing sequence-selective RNA-protein interactions.

We analyzed the global translational inhibition caused by RocA, as well as its marked mRNA selectivity, using ribosome profiling⁷. RocA treatment of HEK 293 cells caused a dose-dependent decrease in polysome formation and protein synthesis (Extended Data Figures 1a and 2a). Translation was inhibited without 4EBP dephosphorylation or eIF2a phosphorylation (Extended Data Figure 1b), but partially rescued by expression of RocA-resistant eIF4A proteins⁶ (Extended Data Figure 1c and d). We quantified the reduction in overall cytosolic ribosome footprints following normalization of our ribosome profiling data against footprints from the mitochondrial ribosome⁸, which employs molecular machinery distinct from the cytoplasmic translation apparatus (Figure 1a and Extended Data Figure 1e-h). We saw that RocA sensitivity varied widely across different transcripts (Figure 1a and b and Supplementary Tables 1a and b). This mRNA-specific translational repression occurred

[†]Correspondence and requests for materials should be addressed to N.T.I. (ingolia@berkeley.edu).

The authors declare no competing financial interests

Supplementary Information Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

Reprints and permissions information is available at www.nature.com/reprints.

Author Contributions S.I. performed all experiments and analyzed the data. Recombinant protein purification and fluorescence polarization assay were performed with the help of S.N.F. S.I. and N.T.I. designed the experiments and wrote the manuscript. Sequences generated in this study have been deposited in the Gene Expression Omnibus under accession numbers GSE70211 and GSE79392

even at a low, therapeutically relevant concentration of RocA (30 nM)¹⁻⁴, correlated well between different drug concentrations, and was not accompanied by significant changes in mRNA abundance (Extended Data Figure 2b-d and Supplementary Table 1c).

Given that eIF4A acts during the scanning of the pre-initiation 43S complex along the 5' UTR⁹, we reasoned that the varied RocA sensitivity of different mRNAs might be determined by their 5' UTR sequences. We confirmed that the 5' UTRs of selected mRNAs were sufficient to confer RocA sensitivity on a *Renilla* luciferase reporter, while the scanning-independent HCV IRES¹⁰ was totally resistant to the drug (Figure 1c and Extended Data Figure 2e). However, RocA sensitivity did not reflect either the calculated thermodynamic stability or experimentally derived DMS-Seq secondary structure measurement¹¹ of the 5' UTR, and the presence of predicted G-quadruplexes⁵ contributed only modestly (Extended Data Figure 3).

Because RocA enhances the RNA affinity of eIF4A^{1,2,6}, we suspected that it could induce effects beyond the simple loss of eIF4A activity. Indeed, we found that the eIF4A inhibitor Hippuristanol (Hipp), which decreases the affinity between eIF4A and RNA^{12,13}, yields a different spectrum of mRNA-specific repression (Extended Data Figure 4a-e). The mTOR inhibitor PP242, which inhibits formation of eIF4F (a complex of eIF4E/G/A)^{14,15}, represses a subset of these Hipp-sensitive mRNAs (Extended Data Figure 4f and g). Thus, RocA exerts effects beyond reduced eIF4A activity, particularly at low, therapeutic doses.

We next asked how RocA affected eIF4A occupancy across the transcriptome in cells by sequencing transcripts that co-purified with streptavidin binding peptide (SBP) tagged eIF4A (Extended Data Figure 5) (RIP-Seq). Increasing RocA doses elevated the overall amount of RNA that co-purified with SBP-tagged eIF4A, (Extended Data Figure 5d), and greatly changed the abundance of individual transcripts, leading to 15-fold or larger differences between mRNAs. Strikingly, enhanced eIF4A binding in the presence of RocA correlated strongly with translation inhibition by RocA (Figure 1d and Extended Data Figure 5f), suggesting that a selective increase of the eIF4A-RNA affinity underlies the specific translation inhibition caused by RocA.

This mRNA selectivity led us to explore the sequence preferences of eIF4A in the absence and presence of RocA. We measured the RNAs that bound to eIF4A out of a random pool of oligonucleotides using deep sequencing (RNA Bind-n-Seq)¹⁶ (Extended Data Figure 6a-c). We then calculated the enrichment of 4- to 6-nt motifs in RNAs retained on eIF4A, as DEAD-box RNA helicases typically contact 6 nt¹⁷. The motifs enriched from randomized synthetic RNA by Bind-n-Seq also predicted RIP-Seq enrichments of endogenous transcripts (Figure 2a and Extended Data Figure 6d). In both experiments, RocA greatly enhanced binding to short polypurine sequences (Figures 2b and c and Extended Data Figure 6e). Although drug-free eIF4A also had intrinsic RNA sequence preferences¹⁸ (Extended Data Figures 6g and 8b and c) and transcripts containing these preferred sequences were relatively resistant to Hipp treatment (Extended Data Figure 6h), RocA only selectively increases binding to a subset of sequences containing polypurine stretches (Extended Data Figure 6g).

Polypurine motifs were also enriched in the eIF4A binding sites detected by photocrosslinking and immunoprecipitation (iCLIP)¹⁹ following RocA treatment (Figure 2d and Extended Data Figure 7), and in the 5' UTRs of translationally RocA-sensitive mRNAs (Figure 2e). This striking correspondence among *in vitro* binding to recombinant protein, *ex vivo* co-purification, crosslinking in cells, and translational repression in cells led us to hypothesize that selective binding to polypurine motifs induced by RocA binding could explain mRNA-specific translational repression. We then directly confirmed that inserting the polypurine motif into an unstructured CAA repeat 5' UTR (Extended Data Figure 9a)²⁰ sensitized the reporter to RocA inhibiton (Figure 2f).

We found that RocA-induced, sequence-selective eIF4A binding occurs through ATPindependent clamping that suffices to repress translation of the clamped mRNA. The cycle of ATP-dependent RNA binding and subsequent release upon ATP hydrolysis is necessary for the efficient RNA remodeling activity of eIF4A as well as its role in translation²¹. Drugfree eIF4A bound RNA only in the presence of ATP (AMP-PNP and ADP-BeF_x) and transition state (ADP-AlF₄) analogues but not hydrolysis products (ADP + Pi). Remarkably, RocA clamped eIF4A on polypurine RNA, but not CAA-repeat RNA, in an ATPindependent manner (Figure 3a and b and Extended Data Figure 8a and b). Bind-n-Seq performed with ADP + Pi likewise recovered polypurine-enriched RNAs in the presence of RocA and no detectable RNA in the absence of RocA (Figure 3c and Extended Data Figure 6i). RocA provided polypurine-specific RNA binding activity to mutant eIF4A defective for ATP binding (VX₄GKT)²², which does not bind to RNA at all without RocA (Extended Data Figure 8d-f), and even to the truncated N-terminal domain of eIF4A, albeit with lower affinity (Extended Data Figure 8g). The eIF4A/RocA complex dissociated far more slowly from polypurine RNA than naïve eIF4A, even in the presence of ATP, whose hydrolysis ordinarily permits rapid dessociation (Figure 3d). High RNA affinity in the ADP-bound state can prolong RNA binding beyond the time required for adenosine nucleotide exchange to restore the high-affinity ATP-bound state and thus greatly reduce the effective dissociation rate. This effective dissociation rate from polypurine RNA measured in hydrolysable ATP (reflective of the intracellular environment) becomes much slower than the ~1 minute timescale of translation initiation²³, and could serve to directly block the ribosome.

In order to probe how clamped eIF4A repressed translation, we recapitulated RocA-induced, polypurine motif-specific translational repression in rabbit reticulocyte lysate (RRL) (Extended Data Figure 9b and c). In this system, RocA treatment represses the formation of 48S pre-initiation complexes on the start codon of sensitive mRNAs, which we assessed using a primer extension toeprinting assay^{20,24} (Figure 3e and Extended Data Figure 9d). Surprisingly, we observed additional RocA-dependent toeprints on the 5' UTR, corresponding to the position of polypurine motifs (Figure 3e), even without eIF4F recruitment (Extended Data Figure 9e). We recapitulated these toeprints using only purified eIF4A and drug, showing that they reflect eIF4A/RocA complexes clamped directly onto polypurine motifs, bypassing its canonical recruitment via cap and the eIF4F complex²¹ (Extended Data Figure 9f). RNase I footprinting revealed the full extent of the eIF4A protected region centered on the motif (Figure 3f and Extended Data Figure 9g).

Page 4

These eIF4F-independent eIF4A/RocA complexes directly repress translation. We preformed such stable complexes on an mRNA during a pre-incubation with recombinant eIF4A and RocA, and then showed that they repressed its subsequent translation in the absence of free RocA (Figure 4a). Recombinant forms of eIF4A bearing mutations that disrupt either ATP binding or eIF4G binding still retained the ability to clamp onto polypurine RNA in the presence of RocA (Extended Data Figures 8d-f,h-i, and 9h-i) and repress translation from the RNA as strongly as wild-type eIF4A/RocA complex (Extended Data Figure 9j). Furthermore, supplementation of recombinant eIF4A protein into an *in vitro* translation reaction actually strengthened the repressive effect of RocA (Figure 4b and Extended Data Figure 9k), confirming the dominant repressive effect of the eIF4A/RocA complex. In contrast, translation repression by Hipp, which decreases the affinity between eIF4A and RNA and thereby mimics a loss of its function, was relieved by the addition of recombinant eIF4A.

Assembly of an eIF4A/RocA complex could in principle repress 48S formation by blocking 40S attachment to the 5' end of an mRNA or subsequent 43S scanning along the 5' UTR. Because the impact of eIF4A/RocA bound to a single polypurine motif is unaffected by its distance from the 5' end (Extended Data Figure 9a and 1), we infer that eIF4A/RocA bound to these motifs blocks 43S scanning. We also found that eIF4A/RocA could inhibit translation from the polio virus (PV) internal ribosome entry site (IRES), which bypasses ordinary 40S recruitment but still depends on scanning (Extended Data Figure 9a)²⁵ when we inserted polypurine motifs in the scanned region (Figure 4c and Extended Data Figure 9m). Scanning inhibition suffices to explain repression by the eIF4A/RocA complex, although our data do not exclude an additional effect on 40S loading.

Impediments to 43S scanning by stable hairpins²⁶ or RNA-binding proteins²⁷ can enhance the translation from upstream open reading frames (uORFs) that otherwise would be skipped. We observed that RocA treatment, but not Hipp treatment, caused an analogous accumulation of translation on 5' UTRs despite the global reduction in footprints on proteincoding sequences (CDSes) (Extended Data Figure 10a and b). This enhancement occurred specifically on high-sensitivity transcripts (Figure 5d and Extended Data Figure 10c). The uORFs activated by RocA showed enrichment of a polypurine motif 20-30 nt downstream of the uORF initiation codon (Figure 5d, inset), reflecting the distance between the start site and the leading edge of the scanning complex²⁷. We tested directly whether eIF4A/RocA complexes on polypurine motifs can drive cryptic upstream initiation using a reporter mRNA with two alternative start sites that yield distinguishable protein isoforms. Insertion of a polypurine motif 30 nt downstream of the earlier AUG increased translation initiation from this codon upon RocA treatment (Figure 5e), confirming that clamped eIF4A/RocA complexes on polypurine motifs drive upstream translation initiation. We found evidence that that this enhanced upstream initiation could contribute to eIF4A/RocA-mediated repression of downstream CDSes²⁸, as RocA-sensitive transcripts showed more pre-existing uORF initiation²⁹ (Extended Data Figure 10d and e).

We have shown that RocA induces ATP-independent clamping of eIF4A onto polypurine sequences, creating an inhibitory roadblock for the scanning ribosome (Figure 5d). Our identification of the eIF4A/RocA binding motif provides the first observation of a drug that

stabilizes sequence-selective RNA-protein interactions³⁰. RocA may bind near the RNA interface on the N-terminal domain of eIF4A⁶, raising the possibility that the drug directly contacts purine bases of target RNAs. Alternatively, RocA might induce a conformational change leading to direct or indirect recognition of the polypurine motif by protein residues. Future structural insight into this polypurine selectivity may enable rocaglate derivatives with altered base selectivities that target different mRNA.

Methods

General methods

HEK 293 Flp-In T-Rex cells (Invitrogen) were cultured and recombined according to manufacturer's instructions. Stable integrants of SBP-tagged eIF4A were produced by co-transfection of these plasmids along with pOG44 by X-tremeGENE 9 (Roche) and selection using Hygromycin B. RocA, PP242, and Thapsigargin were purchased from Sigma. Proteins and DNAs/RNAs were stained with GelCode Blue Stain Reagent (Thermo Scientific) and SYBR Gold Nucleic Acid Gel Stain (Invitrogen), respectively.

Ribosome profiling

Library preparation and data analysis were performed according to the method previously described³², which monitors mitochondrial ribosomes as well^{8,32}. DMSO, RocA, Hipp, and PP242 were added to medium 30 min. prior to cell lysis. The libraries were sequenced on a HiSeq 2000/2500 (Illumina).

RIP-Seq

Cells with tetracycline-inducible, SBP-tagged eIF4A integrated stably were plated in a 10 cm dish and cultured for 3 days with 1 µg/ml tetracycline, incubated with DMSO, 0.03 µM, or 0.3 µM RocA for 30 min, washed once with 5 ml of ice-cold PBS, lysed with 600 µl of lysis buffer (20 mM Tris-HCl pH 7.4, 150 mM NaCl, 5 mM MgCl₂, and 1 mM DTT) containing 1% Triton X-100 and Turbo DNase I (Invitrogen) 25 U/ml, and then clarified by centrifugation for 10 min at 20,000 xg, 4 °C. The supernatant was incubated with 60 µl of Dynabeads M-270 Streptavidin (Invitrogen) equilibrated with lysis buffer containing 1% Triton X-100 and 7 mb beads were washed 5 times with lysis buffer containing 1% Triton X-100 and 1 M NaCl. SBP-eIF4A and bound RNAs were eluted with 25 µl of lysis buffer containing 5 mM biotin at 4 °C for 30 min. All buffers contained 0.001% DMSO with or without 0.03 or 0.3 µM RocA. RNAs were extracted with QIAzol (Qiagen) using the Direct-zol RNA miniprep (Zymo research). One-third of eluted RNA (~100 ng) was mixed with 1 ng of *in vitro* transcribed, spike-in *Renilla* luciferase RNA (*hRluc*) (see "DNA constructs") and sequencing libraries were prepared using Tru-seq Ribo-zero gold kit (Illumina). Libraries were sequenced on HiSeq2000/2500 (Illumina) sequencers.

iCLIP

Cells were cultured as described in "RIP-Seq". After medium was substituted with ice-cold PBS, the dishes on ice were irradiated with 150 mJ/cm² with UV-C (~254 nm) in UVP CL-1000 (UVP). Lysate was prepared as described in "RIP-Seq". The lysate from a 10 cm dish (600 μ l) was treated with 0.4 U of RNase I (Epicentre) at 37 °C for 3 min. Reaction was

quenched by the addition of 10 µl of SUPERase In RNase Inhibitor (Invitrogen), and then incubated with 60 µl of Dynabeads M-270 Streptavidin (Invitrogen) equilibrated with lysis buffer containing 1% Triton X-100 at 4 °C for 30 min. The beads were washed by CLIP wash buffer (20 mM Tris-Cl pH 7.4, 1 M NaCl, 2 mM EDTA, 1 mM DTT, and 1% Triton X-100) twice, by CLIP wash buffer containing 0.1% SDS and 0.05% sodium deocycholate twice, and then by lysis buffer containing 1% Triton X-100 twice. After discarding the supernatant, the beads were incubated with 10 U T4 PNK (NEB), 1x PNK buffer, and 0.33 μM ³²P-γ-ATP (3000 Ci/mmol, PerkinElmer) in 10 μl at 37 °C for 5 min and washed once with lysis buffer containing 1% Triton X-100. RNA-crosslinked proteins were eluted by 20 µl of lysis buffer containing 1% Triton X-100 and 5 mM biotin at 37 °C for 5 min, run onto NuPAGE (Invitrogen), and transferred to nitrocellulose membrane 0.45 µm (Biorad). The images of ³²P-labeled RNA-protein complex on the membrane were acquired by Typhoon TRIO (Amersham Biosciences). The membrane with the region containing SBPeIF4A/RNA complexes was excised and treated with 0.1 µg/µl Proteinase K (Thermo Scientific), 200 mM Tris-Cl, pH 7.4, 25 mM EDTA, pH 8.0, 300 mM NaCl, and 2% SDS in 200 µl at 55 °C for 20 min. RNAs were isolated by phenol/chloroform extraction and ethanol precipitation. Library preparation was performed according to the method previously described³² with following modifications. As linker DNA, 5'-

(Phos)NNNNNIIIIITGATCGGAAGAGAGCACACGTCTGAA(ddC)-3' where (Phos) indicated 5' phosphoryaltion and (ddC) indicates a terminal 2', 3'-dideoxycytidine, was used. The Ns indicate random barcode and the Is indicate a sample mulplexing barcode. For multiplexing, linker DNAs containing ATCGT for DMSO replicate #1, AGCTA for DMSO replicate #2, CGTAA for RocA 0.03 μ M, CTAGA for RocA 0.3 μ M, and GATCA for RocA 3 μ M in Is position were used, respectively. The linker DNAs were pre-adenylated by 5' DNA adenylation kit (NEB) before the ligation reaction. Instead of gel extraction, unreacted linkers were removed by the treatment of the ligation reaction with 5' deadenylase (NEB) and RecJ exonuclease (epicentre) at 30 °C for 45 min. Reverse transcription was performed with an oligo 5' -

(Phos)NNAGATCGGAAGAGCGTCGTGTAGGGAAAGAG(iSp18)GTGACTGGAGTTCA GACGTGTGCTC-3', where (Phos) indicates 5' phosphorylation and Ns indicate random barcode. PCR was performed with oligos, 5'-

AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTC-3' and 5'-CAAGCAGAAGACGGCATACGAGATCGTGATGTGACTGGAGTTCAGACGTGTG-3'. Libraries were sequenced on HiSeq4000 (Illumina) sequencers. Random barcode was used to eliminate PCR duplicates in the library.

Bind-n-Seq

SBP-tagged eIF4A was purified as described in "RIP-Seq", without DMSO or RocA treatment. The beads tethering SBP-eIF4A were treated with 1x Micrococcal Nuclease Buffer (NEB), 0.5x lysis buffer, 0.5% Triton X-100, and 200 U/µl Micrococcal Nuclease (NEB) in 30 µl at 25 °C for 30 min, washed 5 times with lysis buffer containing 1% Triton X-100, 1M NaCl, and 5 mM EGTA pH 7.4, and rinsed twice with lysis buffer containing 0.1% Triton X-100. The beads were incubated in lysis buffer containing 0.1% Triton X-100, 2 mM AMP-PNP, 0.33 U/µl SUPERase In RNase Inhibitor (Invitrogen), 1 µM N₃₀ RNA [(N)₃₀CTGTAGGCACCATCAAT, bold characters represent DNA sequence for reverse

transcription primer hybridization] in 30 μ l at 37 °C for 30 min, and washed 5 times with lysis buffer containing 0.1% Triton X-100, 2 mM AMP-PNP, and 0.1% DMSO. SBPeIF4A/RNA complex was eluted with 30 μ l of lysis buffer containing 0.1% Triton X-100, 2 mM AMP-PNP, and 5 mM biotin. 0.1% DMSO with or without 30 or 300 nM RocA was present in all buffers during the RNA binding reaction, wash, and elution. RNAs were extracted with QIAzol (Qiagen) using the Direct-zol RNA miniprep (Zymo Research) and converted into DNA library as the same method of ribosome profiling³². For Bind-n-Seq with ADP + Pi, 2 mM ADP, 2 mM Na₂HPO₄, 50 μ M N₃₀ RNA, and 3 μ M RocA were used.

The 30-nt randomized RNA followed by 3' DNA sequence for reverse transcription priming was designed to avoid ligation biases and sequencing of contaminating RNA fragments from cells during SBP-eIF4A purification, and to cover the entire sequence with a single 50-bp mode of HiSeq (Illumina) sequencers.

Our read depth (~10⁸ reads) is less than the theoretical complexity ($4^{30} \approx 10^{18}$), so that the probability that the same sequence appears multiple times in the library is quite low. Therefore, we assumed that reads with exactly the same sequence and length in the library reflect PCR duplicates and counted them only once. Motif enrichment in the range of interest (4-6 nt) was calculated as the ratio of the motif frequency between libraries¹⁶.

Spearman's correlation of motif number in 5' UTR versus RIP-Seq -fold change caused by RocA treatment was used as motif prediction in RIP-Seq. High scoring motifs were defined as those with enrichment of the prediction or the enrichment is >1.5 standard deviation from the mean in RIP-Seq and Bind-n-Seq, respectively.

Data analysis

The reads were aligned to the hg19 human genome reference and the resulting aligned reads were mapped to UCSC known reference genes, downloaded from UCSC genome browser on July 2013. A UCSC bed file of known genes was used for the 5' UTR analysis. For mitochondria footprints alignments, we used the RefSeq genes track corresponding to the mitochondrial chromosome (chrM), downloaded from UCSC genome browser. Specific Asite nucleotides were empirically estimated based on the length of each footprint. The offsets were 14 for 26-29 nt and 15 for 30-31 nt. For mitochondria footprints, 9 for 26-27 nt, 11 for 28-29 nt, 12 for 30 nt, 13 for 31 nt, and 18 for 32-34 nt. For mRNA fragments, we used offset 14. For measuring footprint density and mRNA fragments of RIP-Seq between samples, we restricted our analysis to genes, which have at least 40 and 100 summed counts in each sample, respectively. For CDSs, the analysis only included the transcript positions beginning 15 codons following the start codon and stopping 5 codons preceding the stop codon. For 5' UTRs, we included the transcript positions from the 5' end of the mRNA until 5 codons preceding the start codon. DESeq³³ was used to calculate relative enrichment of genes in the library, including the mitochondrial footprints and spike-in hRluc mRNA counts. The calculated -fold change was re-normalized to the value of the summed mitochondria footprints or the spike-in hRluc mRNA fragments.

High-sensitivity messages were defined as transcripts with reduction more than 2-fold from the median, and with q value <0.01, between 3 μ M RocA-treated and untreated cells. Low-

sensitivity transcripts are defined as same as high-sensitivity but with accumulation over 2-fold.

For calculation of G, RNALfold (ViennaRNA Package)³⁴ was run with -L30 - g options on 5' UTRs sequences from UCSC foldUtr5 table. The minimum G along each 5' UTR was used as a representative free energy value for each gene.

The presence of G-quadruplexes was predicted with RNAfold (ViennaRNA Package). The Gini differences across 5' UTRs were calculated using published data¹¹. Analysis was restricted to the mRNAs bearing 5' UTRs which have one or more reads on A/C positions on average.

"uORF translation intensity" was calculated using published data²⁹. To incorporate the number and intensity of each upstream initiation site in the 5' UTR, we calculated the density of 5' UTR footprints for each transcript as mentioned above, as the great majority of these footprints derive from ribosomes trapped on first codons (Extended Data Figure 10c). To normalize mRNA abundance in cells, we normalized the density by footprint counts from the CDS start codon region using the genomic position between start codon and 6 nt downstream. We restricted the analysis to transcripts with at least 10 counts from CDS start codons and re-normalized the value to the median as 1.

Fluorescence polarization assay

Proteins (0-50 μ M) were incubated in 14.4 mM HEPES-NaOH, 108 mM NaCl, 1 mM MgCl₂, 0.36 mM TCEP, 14.4% glycerol, 0.1% DMSO, and 10 nM 5'FAM-labeled RNA with or without 50 μ M RocA in 10 μ l reaction for 30 min at 25 °C. The experiments were performed with 1 mM AMP-PNP (for AMP-PNP), 1 mM ADP, 5 mM BeCl₂, and 25 mM NaF (for ADP-BeF_x), 1 mM ADP, 5 mM AlCl₃, and 25 mM NaF (for ADP-AlF₄), or 1 mM ADP and 1 mM Na₂HPO₄ (for ADP + Pi). For the condition without ATP analog, MgCl₂ was omitted from the reaction.

For competition assay, the complexes were preformed with 1 mM ATP or AMP-PNP, 1 μ M eIF4A, 10 nM FAM-labeled RNA, and 50 μ M RocA and chased with 100 μ M non-labeled RNA. Because of the low affinity, 50 μ M eIF4A was used with ATP and DMSO.

Fluorescence polarization was measured using an Infinite F-200 PRO (TECAN). K_d and $t_{1/2}$ was calculated with fitting to Hill equation and one-phase exponential decay equation, respectively, by Igor Pro software (WaveMatrics).

In vitro translation and toeprinting assay

In vitro translation was performed with nuclease-treated RRL system (Promega), according to manufacture's instruction. Fifty nM reporter mRNAs (see "DNA constructs") was incubated in 50% RRL with RocA (concentration shown in the figure legends) or 1% DMSO in 10 μ l at 30 °C for 1 hr. For the detection of SBP, 20 μ l of the reaction was used with uORF + CAACAA or uORF + AGAGAG mRNAs and concentrated with 10 μ l of Dynabeads M-270 Streptavidin (Invitrogen).

Toeprinting assay was performed as previously described³⁵. Reaction was pre-incubated with RRL in the presence of 2 mM GMP-PNP or m⁷GTP and 3 μ M RocA or 1% DMSO at 30 °C for 5 min, and then incubated with 50 nM mRNAs at 30 °C for 5 min, followed by reverse transcription with 10 U/ μ l ProtoScript II (NEB) with 250 nM 5′ 6-FAM labeled primer (5′-6-FAM-ATGCAGAAAAATCACGGC-3′) at 30 °C for 15 min. Ten μ M of recombinant eIF4A was used instead of RRL in 30 mM HEPES, pH 7.3, 100 mM KOAc, 1 mM Mg(OAc)₂, and 1 mM DTT in the presence or absence of 10 μ M RocA. cDNA was purified by phenol extraction, concentratated using Oligo clean & concentrator (Zymo Research), and run with GeneScan 600 LIZ size standard v2.0 (Life Technologies) on 3730 DNA Analyzer (Life Technologies). Data were analyzed by GeneMapper software (Life Technologies). For pre-formation of eIF4A/RocA complex on mRNA, 30 μ l of the reaction was loaded on G-25 column equilibrated with 30 mM HEPES, pH 7.3, 100 mM KOAc, 1 mM Mg(OAc)₂, and 1 mM DTT to remove free RocA. The flow through mRNA was used for *in vitro* translation at 20 nM.

Dideoxy-terminated sequencing of RNA was performed by reverse transcriptions using 0.125 mM individual dideoxy-NTP and 0.5 mM each deoxy-NTP with the same 5' 6- FAM labeled primer and ProtoScript II, according to manufacture's instructions.

RNase I footprinting assay

Reporter RNA was incubated with recombinant eIF4A and RocA in 12 µl as described in toeprinting assay. The reaction was treated with 1 µl of 0.001 U/µl RNase I (Epicentere) at room temperature for 5 min. After quenching the digestion by the addition of 1 µl of SUPERase In RNase Inhibitor (Invitrogen), RNA was extracted by Oligo clean & concentrator (Zymo Research) and reverse transcribed by ProtoScript II (NEB) with 5' 6-FAM labeled primer (5'-6-FAM-ATGCAGAAAAATCACGGC-3') according to manufacturer's intruction. The cDNA was run on 3730 DNA Analyzer (Life Technologies) as described in toeprinting assay. Data were analyzed by GeneMapper software (Life Technologies).

Polysome profiling

Cells lysate was prepared as described previously³². Lysate containing 15 μ g total RNA was loaded on to 10-50% linear sucrose gradients containing 20 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM MgCl₂, 1 mM DTT, 100 μ g/ml cycloheximide, and 2 U/ml SUPERase In RNase Inhibitor and sedimented by ultracentrifugation at 36,000 rpm for 2 hr at 4 °C with SW41 rotor (Beckman Coulter). Gradients were fractionated using Gradient station (Biocomp). UV absorbance was detected by ECONO UV monitor (Biorad).

Metabolic labeling of nascent peptide by OP-puro

Nascent peptides in HEK 293 cells were labeled by 40 μ M OP-puro (Jena Bioscience) in 24 well dishes with 0-3 μ M RocA for 30 min. Cells were washed with PBS and lysed with 50 μ l of lysis buffer (20 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM MgCl₂, and 1 mM DTT) containing 1% Triton X-100, and then clarified by centrifugation with 20,000 xg at 4 °C for 10 min. Nascent peptides were labeled with 5 μ M Alexa Fluor 488 Azide (ThermoFisher Scientific) by Click-it cell reaction kit (ThermoFisher Scientific) according to

manufacturer's instruction and run on SDS-PAGE. Images were acquired by FluorChem R imaging sysmtem (ProteinSimple) and quatified by AlphaView (ProteinSimple).

DNA constructs

DNA fragments containing 5' UTRs sequences, listed below, were inserted between T7 promoter and ORF of *Renilla* luciferase (hRluc) in psiCHECK2 (Promega). We cloned the exactly same sequence of G-quadruplex and its control sequence used in⁵. These plasmids were digested by *Not*I and used as *in vitro* transcription template.

PTGES3 (uc001slu.4);

EIF2S3 (uc004dbc.3);

TTTCCTTCCTCTTTTGGCAAC

HNRNPC (uc001vzy.3);

AGGAATGGGGCGGGGACTAGGCCTTCGCCTCGGCGGCAGAGGAGACTCGGGG GCCATTTTGTGAAGAGACGAAGACTGAGCGGTTGTGGCCGCGTTGCCGACCTCC AGCAGCAGTCGGCTTCTCTACGCAGAACCCGGGAGTAGGAGACTCAGAATCGAA TCTCTTCTCCCTCCCCTTCTTGTGAGATTTTTTTGATCTTCAGCTACATTTCGGCT TTGTGAGAAACCTTACCATCAAACACG

GPX1 (uc021wxw.1);

CAGTTAAAAGGAGGCGCCTGCTGGCCTCCCCTTACAGTGCTTGTTCGGGGGCGCTC CGCTGGCTTCTTGGACAATTGCGCC

TMA7 (uc003cte.1);

GGGGAAGCGGCGGCAGGCGCC

KMT2A (uc001pta.3);

CTGCTTCACTTCACGGGGGCGAAC

HCV IRES;

CCGCTCAATGCCTGGAGATTTGGGCGTGCCCCCGCAAGACTGCTAGCCGAGT AGTGTTGGGTCGCGAAAGGCCTTGTGGTACTGCCTGATAGGGTGCTTGCGAG TGCCCCGGGAGGTCTCGTAGACCGTGCACCATGAGCACGAATCCTAAACCTC AAAGAAAAACCAAACGTAAC

G-quadruplex;

CTAGGTTGAAAGTACTTTGACGGCGGCGGCGGCGGTCAATCTTACGGCGGCGG CGGACATAGATACGGCGGCGGCGGCGGTAGAAACTACGGCGGCGGCGGATTA GAATAGTAAA

(bold characters represent G quadruplex forming sequences)

Randomized control for G-quadruplex;

CTAGGGCGCACGTACTTCGACAACGTCAGCGTTCAGCGTTCCAACGTCAGCG TACAGCGATCCAACGTCAGCGTTCTGCGCTACAACGTCAGCGTATCCGCGTA GCACA

CAA repeat;

7x AGAGAG motifs;

1x AGAGAG left;

1x AGAGAG middle;

1x AGAGAG right;

PV IRES;

TTAAAACAGCTCTGGGGTTGTACCCACCCAGAGGCCCACGTGGCGGCTAGT ACTCCGGTATTGCGGTACCCTTGTACGCCTGTTTTATACTCCCTTCCCGTAACT TAGACGCACAAAACCAAGTTCAATAGAAGGGGGGTACAAACCAGTACCACCA CGAACAAGCACTTCTGTTTCCCCGGTGATGTCGTATAGACTGCTTGCGTGGTT

PV IRES 3x AGAGAG;

The following DNA fragments, coding *Drosophila msl-25'* UTR and SBP, were amplified by PCR and used for *in vitro* transcription template.

uORF + CAACAA;

uORF + AGAGAG;

TAATACGACTCACTATAGGGCAGCATAACCATTGTTGATGACTCGAGACCTCT CAAACGTAAACCAAAGAGAGGCACGTGACACCATGGACGAGAAAACCACC

Reporter RNAs were *in vitro* transcribed, capped, and polyadenylated using T7-Scribe Standard RNA IVT Kit, ScriptCap m⁷G Capping System, ScriptCap 2'-O-Methyltransferase Kit, and A-Plus Poly(A) Polymerase Tailing Kit (CELLSCRIPT). Capping reaction was skipped for PV IRES and PV IRES 3x AGAGAG reporters.

For the generation of stable cell-lines, PCR products containing CDS region of *EIF4AI* mRNA and SBP amplified from cDNA from Human adult normal brain (Invitrogen) and from pASW³⁶ (a kind gift from Y. Tomari), respectively were inserted into *Hin*dIII site in pcDNA5/FRT/TO (Invitrogen) by Gibson assembly (NEB). P159Q, F163L, and Q194E mutations were introduced by site-directed mutagenesis.

For recombinant eIF4A protein expression, PCR products containing CDS region of *EIF4AI* mRNA were inserted into pHM-GWA³⁷ to construct pHisMBP-eIF4A. VX₄GKT (A82V) and D296A-T298K mutations were introduced by site-directed mutagenesis. His-tag, MBP-tag, Tobacco Etch Virus protease cleavage site, and N-terminal region of eIF4A (1-237) were cloned into pET-28a, to construct pHisMBP-eIF4A (1-237).

Reporter assay in HEK 293 cells

Transfections were performed in 24 well dishes with TransIT-mRNA Transfection Kit (Mirus) according to manufacturer's instruction, at half scale. Three hours after transfection, RocA was added to the medium and 9 hr after transfection, cells were washed with PBS and lysed with Passive lysis buffer (Promega). Luciferase assay was performed with *Renilla*-Glo Luciferase Assay System (Promega) according to manufacturer's instruction. Luminescence was detected with GloMax-Multi Jr system (Progema).

For stable cell line with SBP-tagged eIF4A and its mutants, HEK 293 Flp-In T-Rex cells were cultured for 4 days with 1 μ g/ml tetracycline prior to the experiments. Tetracycline was included in media during experiments.

qPCR

Cell lysate or *in vitro* translation reaction for luciferase assay was treated with 40 U/ml TurboDNase for 10 min on ice, and then RNA was extracted by TRI Reagent (Sigma) and Direct-zol RNA miniprep (Zymo Research). Reverse transcriptions were performed with ProtoScript II (NEB) and random primer mix (NEB) according to manufacturer's instruction. qPCR was performed with Fast EvaGreen qPCR Mix (BIOTIUM) in BioRad CFX96 Touch Real Time PCR Detection System (Bio-rad) with oligos, 5'-TCGTCCATGCTGAGAGTGTC-3' and 5'-CTAACCTCGCCCTTCTCCTT-3'. RNA from non-transfected cells or *in vitro* translation reaction without the addition of mRNAs was used as qPCR background.

Purification of recombinant eIF4A proteins

Typically, BL21 Star (DE3) E. coli cells (Invitrogen) transformed with pHisMBP-eIF4A, pHisMBP-eIF4A (VX₄GKT), pHisMBP-eIF4A (D296A-T298K), or pHisMBP-eIF4A (1-237) in 1.5 L culture were cultivated to OD₆₀₀ of 0.5 at 37 °C with 50 µg/ml kanamycin and then grown at 16 °C overnight with 1 mM IPTG. The cell pellets were resuspended in His buffer (20 mM HEPES-NaOH, pH 7.5, 500 mM NaCl, 10 mM imidazole, 10 mM β mercaptoethanol) with 0.5% NP-40, sonicated, and centrifuged at $35,000 \times g$ for 20 min. The supernatant was incubated with 1.5 ml bed volume of Ni-NTA Superflow (Qiagen) for 1 hr. The beads were loaded on a gravity column and washed with His buffer containing 1 M NaCl. The proteins were eluted with 50 mM Na-phosphate buffer, pH 7.5, 500 mM NaCl, 100 mM Na₂SO₄, 250 mM imidazole, 10 mM β -mercaptoethanol, treated with Tobacco Etch Virus protease overnight, dialyzed to 20 mM HEPES-NaOH, pH 7.0, 150 mM NaCl, 0.5 mM TCEP, and 10% glycerol, and loaded on MBPTrap HP 5 ml (GE Healthcare). The flowthrough fractions were collected, concentrated with Amicon Ultra 10kDa (Millipore), and loaded onto a HiLoad 16/600 Superdex 75 pg column (GE Healthcare) equilibrated with 20 mM HEPES-NaOH, pH 7.5, 150 mM NaCl, 0.5 mM TCEP. The peak fractions were collected, concentrated with Amicon Ultra 10kDa (Millipore), mixed with 0.25 volume of 80% glycerol, shock-frozen in liquid nitrogen, and stored at -80 °C. All purification steps were performed at 4 °C. Column chromatography was performed using an AKTA purifier (GE Healthcare).

Pulldown assay

The lysate of *E. coli* cells expressing eIF4A WT or eIF4A D296A-T298K proteins from 1 ml culture was prepared as described in "Purification of recombinant eIF4A proteins" and incubated with 10 μ l of HisPur Ni-NTA Magnetic Beads (Thermo Scientific) at 4 °C for 30 min. The beads were washed 5 times with His buffer containing 1 M NaCl, rinsed once with 20 mM HEPES-NaOH, pH 7.5, 10 mM NaCl, 10 mM imidazole, 10 mM β -mercaptoethanol, and incubated with RRL (Promega) at 25 °C for 30 min. After 5 washes with His buffer, the proteins were eluted from the beads by SDS sample buffer.

ATP crosslinking assay

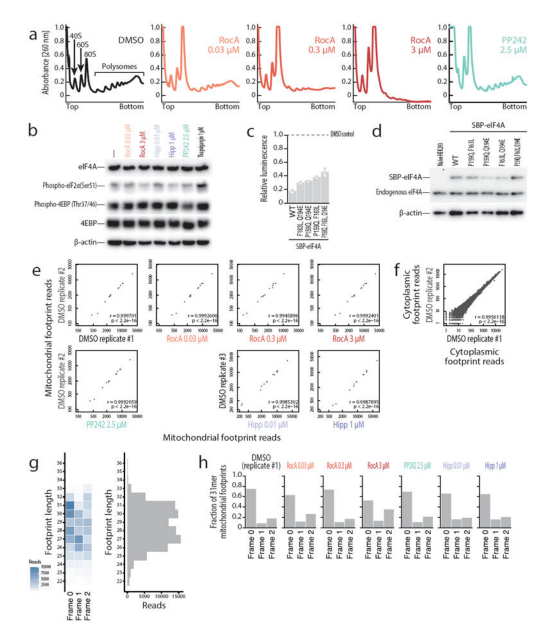
Ten μ M recombinant eIF4A WT and VX₄GKT was incubated with 1 μ M [γ -³²P]-ATP (3000 Ci/mmol, Perkin Elmer) in 30 mM Hepes-KOH, pH 7.3 (Fisher Scientific), 100 mM KOAc, 5 mM Mg(OAc)₂, and 1 mM DTT in 20 μ l reaction at 37 °C for 15 min. The reactions were exposed to 1500 mJ/cm² using UV₂₅₄ (CL1000, UVP) at a distance of 2 cm from the lamp on ice and run on SDS-PAGE. The images were acquired by Typhoon TRIO (Amersham Biosciences).

Western blotting

Anti-eIF4AI (#2490, Cell signaling) (1:1000), anti-phospho-eIF2 α (Ser51) (D9G8 #3398, Cell Signaling) (1:1000), anti-4E-BP1 (#9452, Cell Signaling) (1:2000), anti-phospho-4EBP (Thr37/46) (236B4 #2855, Cell Signaling) (1:2000), anti- β -actin (ab20272, Abcam) (1:1000), anti-eIF4E (#9742, Cell Signaling) (1:1000), anti-eIF4G (#2498, Cell Signaling) (1:1000), and anti-SBP-tag (SB19-C4 sc-101595, Santa Cruz Biotechnology) (1:1000) were

used as primary antibodies. Chemiluminescence was induced by Pierce ECL Western Blotting Substrate (Thermo Scientific) and images were acquired by FluorChem R imaging sysmtem (ProteinSimple).

Extended Data



Extended Data Figure 1. RocA represses translation, targeting to eIF4A

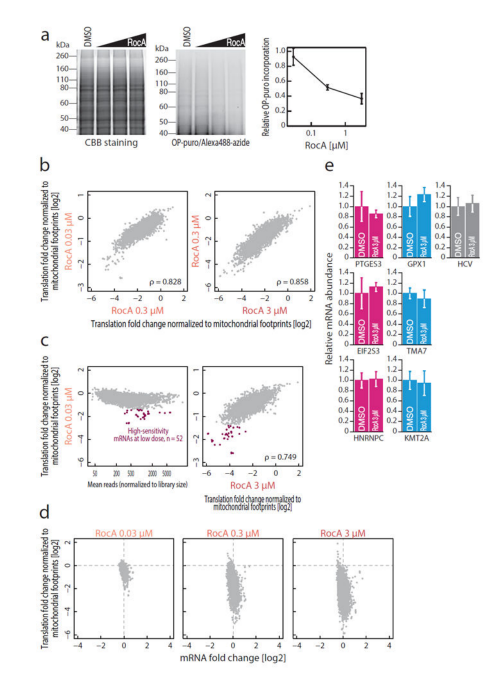
(a) Polysome profiling experiments with RocA and PP242 treatments. RocA disrupts polysomes dose-dependently.

(b) Western blot of phospho-eIF2 α and phospho-4EBP shows that effect of RocA is independent of known translation control targeting to eIFs. Phosphorylation of eIF2 α and dephosphorylation of 4EBP were induced by Thapsigargin and PP242, respectively.

(c and d) Luciferase reporter assay possessing *PTGES35'* UTR (Figure 1c) with exogenous expression of WT or RocA resistant eIF4A mutants (c) and western blot of endogenous and exogenous eIF4A (d). eIF4A is the main molecular target of RocA. Data represent mean and S.D. (n = 3).

(e and f) Correlation of sum of the footprint reads to 13 mitochondrial mRNAs among different conditions (e) and correlation of sum of the footprint reads from cytoplasmic ribosomes to each transcript between biological replicates (f). r is Pearson's correlation. P value is calculated by Student's t-test.

(g and h) Tile plot of codon periodicity along length of mitochondria footprints (g, left) and mitochondria footprint length distribution (g, right) and codon periodicities of 31 nt mitochondrial footprints among different conditions (h). Footprints with 31-nt length showed most homogenous codon periodicity and this periodicity was retained with RocA treatment, showing that mitochondrial ribosome translates even in high doses of RocA.



Extended Data Figure 2. Extended Data Figure 2. RocA represses translation without mRNA degradation

(a) Metabolic labeling of nascent peptides with OP-puro. The OP-puro incorporated nascent peptides were visualized by Click reaction with Alexa488-azide (middle) and quantified (right). Data represent mean and S.D. (n = 3).

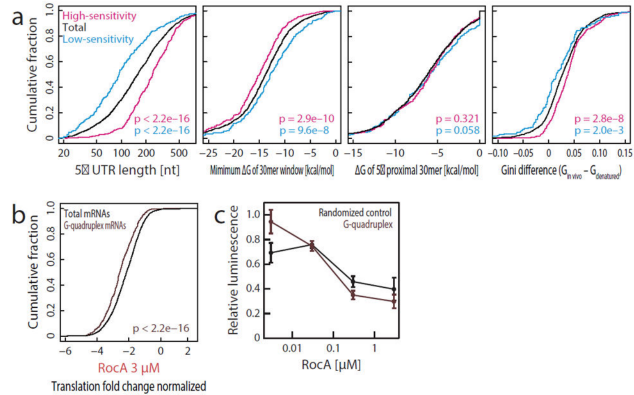
(b) Correlation of translation -fold change among different concentrations of RocA treatments. ρ: Spearman's rank correlation coefficient.

(c) MA plot of mean footprint reads between 0.03 μ M RocA treatment and non-treatment normalized to library sizes to footprints -fold change by 0.03 μ M RocA treatment (left) and the correlation of translation -fold change between 0.03 and 3 μ M of RocA treatments

(right), highlighting high-sensitivity mRNAs at 0.03 μ M RocA treatment. ρ : Spearman's rank correlation coefficient.

(d) Scatter plots of footprints -fold change normalized to mitochondrial footprints and mRNA -fold change by RocA treatments. RocA represses translation without significant mRNA change.

(e) qPCR from the samples of Figure 1c. Data represent mean and S.D. (n = 3).



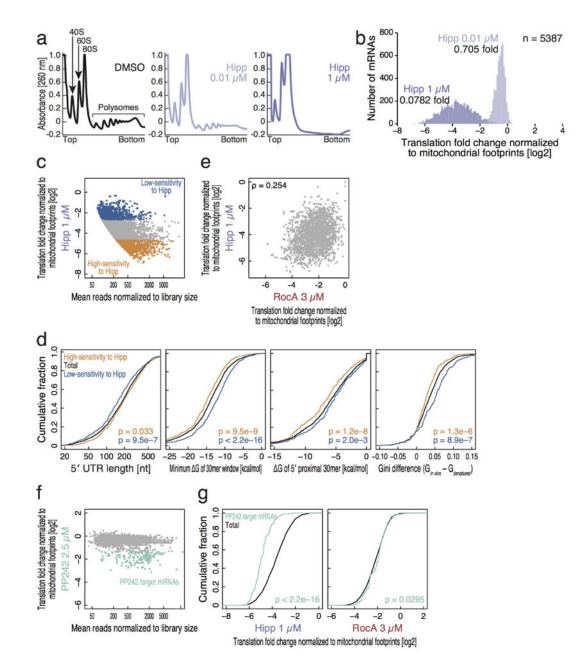
to mitochondrial footprints [log2]

Extended Data Figure 3. Secondary structure in 5' UTR is not strong determinant of RocA sensitivity

(a) Cumulative fractions along length of 5' UTR, minimum G among all 30-mer windows along a 5' UTR, G in cap-proximal region (30 nt) of 5' UTR, and Gini difference are plotted to total, RocA high-sensitivity, and RocA low-sensitivity mRNAs. Significance is calculated by Mann-Whitney U test.

(b) Cumulative fractions along translation -fold change by RocA are plotted to total mRNAs and mRNAs with predicted G-quadruplexes in 5' UTRs. Significance is calculated by Mann-Whitney U test. The impact of presence of G-quadruplex in 5' UTR is modest in RocA sensitivity.

(c) The 5' UTRs with G-quadruplexes and randomized control sequence were fused to *Renilla* luciferase and these reporter mRNAs were transfected prior to treatment with RocA as indicated. Data represent mean and S.D. (n = 3). G-quadruplex does not show the prominent RocA sensitivity.



Extended Data Figure 4. Characterization of translational inhibition by Hippuristanol and PP242

(a) Polysome profiling experiments with Hipp treatments. Hipp disrupts polysomes dose-dependently.

(b) Histograms of number of transcripts along footprints -fold change with 0.01 and 1 μ M Hipp treatment compared to non-treatment, normalized to mitochondrial footprints. Median -fold change is shown. Bin width is 0.1.

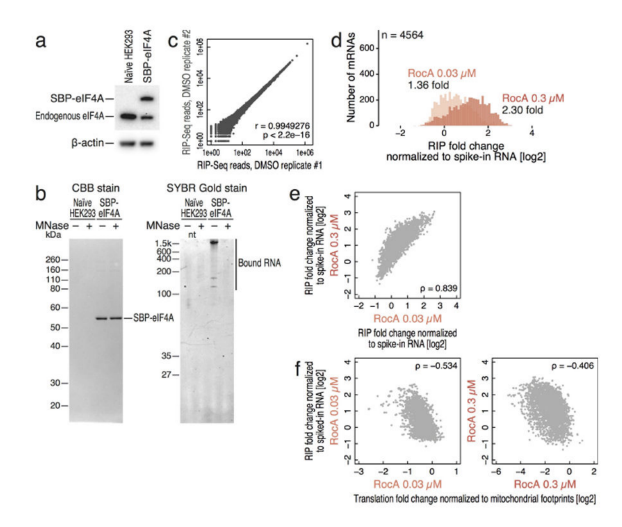
(c) MA plot of mean footprint reads between 1 μ M Hipp treatment and non-treatment normalized to library sizes to translation -fold change by 1 μ M Hipp treatment, highlighting high-sensitivity and low-sensitivity mRNAs.

(d) Cumulative fractions along length of 5' UTR, minimum G among all 30-mer windows along a 5' UTR, G in cap-proximal region (30 nt) of 5' UTR, and Gini difference are plotted to total, Hipp high-sensitivity, and Hipp low-sensitivity mRNAs. Significance is calculated by Mann-Whitney U test.

(e) Translation -fold changes by RocA and Hipp are modestly correlated. ρ : Spearman's rank correlation coefficient.

(f) MA plot of mean footprint reads between 2.5 μ M PP242 treatment and non-treatment normalized to library sizes to translation -fold change by PP242 treatment, highlighting PP242 target mRNAs.

(g) Cumulative distributions of translation -fold change caused by RocA and Hipp treatment are plotted for total and PP242-target mRNAs. Significance is calculated by Mann-Whitney U test.



Extended Data Figure 5. Purification of SBP-tagged eIF4A and co-purified RNA from HEK 293 cells

(a) Western blot of exogenous SBP-eIF4A and endogenous eIF4A in tetracycline-inducible stable cell line. Expression of physiological levels of the tagged allele attenuated

endogenous eIF4A expression but preserved overall eIF4A levels, likely reflecting the same feedback loop previously reported between eIF4AI and eIF4AII³¹.

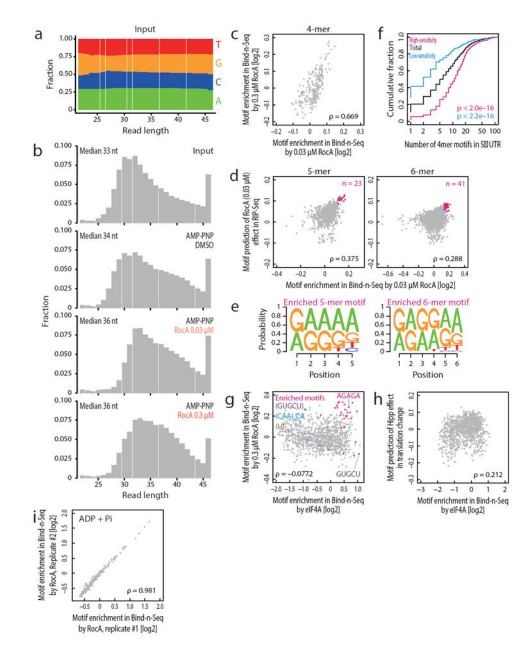
(b) CBB staining of purified SBP-eIF4A and SYBR Gold staining of purified RNA bound to SBP-eIF4A with or without Micrococcal Nuclease (MNase).

(c) Correlation of sum of the mRNA fragment reads of each transcript between biological replicates of RIP-seq. r is Pearson's correlation coefficient. P value is calculated by Student's t-test.

(d) Histogram of the number of transcripts along RNA/eIF4A interaction -fold change by RIP-Seq when cells are treated with 0.03 or 0.3 μ M RocA normalized to spiked-in RNA. Data present the same mRNAs analyzed in Figure 1a. Median -fold change is shown. Bin width is 0.1.

(e) Correlation of RIP -fold change between different concentration of RocA treatments. ρ : Spearman's rank correlation coefficient.

(f) Correlation of translation -fold change to RIP -fold change with the same concentration of RocA treatment. ρ : Spearman's rank correlation.



Extended Data Figure 6. Motif enrichment by Bind-n-Seq

(a) Nucleotide composition in each length of reads in input RNAs for Bind-n-Seq. Input RNAs are random in entire read length.

(b) Length distribution of reads from Bind-n-Seq. RNAs bound to eIF4A showed longer length distribution, indicating that eIF4A has preference for longer RNAs.

(c) Correlations of 4-mer motif enrichment in Bind-n-Seq by 0.03 μ M RocA treatment to that by 0.3 μ M RocA treatment.

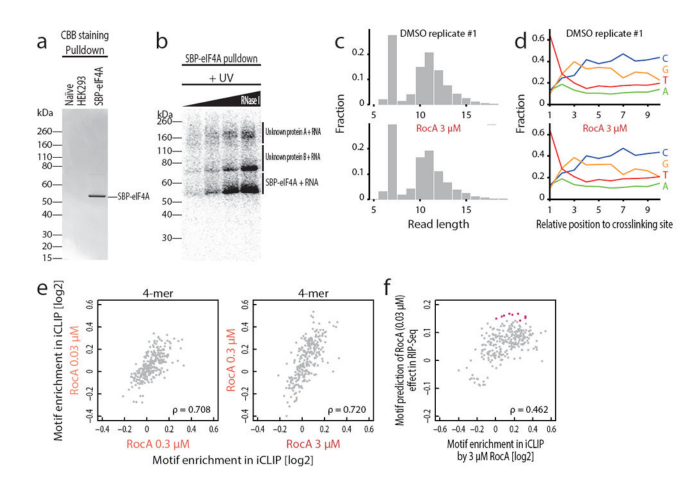
(d) Correlations between 5-mer and 6-mer motif enrichment in Bind-n-Seq by 0.03 μ M RocA treatment and motif prediction of 0.03 μ M RocA effect in RIP-Seq. ρ : Spearman's rank correlation.

(e) Highest-scoring 5-mer and 6-mer motifs in Bind-n-Seq and RIP-Seq.

(f) Cumulative fractions along number of 4-mer motifs (Figure 2b) in 5' UTR are plotted to total, RocA high-sensitivity, and RocA low-sensitivity mRNAs. Significance is calculated by Mann-Whitney U test.

(g) Correlations of Bind-n-Seq motif enrichment (5-mer) by eIF4A to that by 0.03 μ M RocA treatment. The motifs appeared in RNAs used in Extended Data figure 8 are highlighted. (h) Correlation of Bind-n-Seq motif enrichment (5-mer) by eIF4A to motif prediction of Hipp effect in translation change, which is define as Spearman's correlation of motif number in 5' UTR to translation -fold change by Hipp. mRNAs with high affinity motif to eIF4A in 5' UTR are resistant to Hipp treatment.

(i) The correlation between enriched motifs of replicates in Bind-n-Seq with ADP + Pi. ρ : Spearman's rank correlation.



Extended Data Figure 7. Characterization of iCLIP data

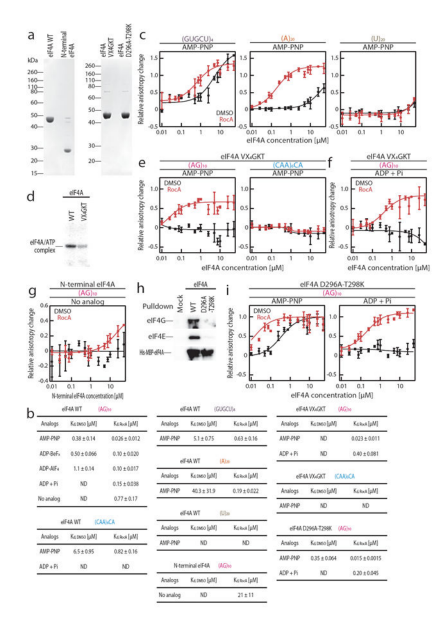
(a) CBB staining of purified SBP-eIF4A protein in iCLIP procedure.

(b) Visualization of RNA-crosslinked with SBP-eIF4A and unknown proteins by ³²P labeling of RNA. We avoided the contamination of RNAs cross-linked to the additional, co-purifying, unknown proteins.

(c) Distribution of read length in iCLIP libraries. Avoidance of contaminating RNAs restricted us to short RNAs, which likely correspond to the region of RNA physically protected by eIF4A binding, or footprint

(d) Nucleotide bias along the reads in iCLIP libraries. The crosslinking bias for U may underestimate the preference for polypurine motifs.

(e) Correlations of iCLIP motif enrichment (4-mer) by different RocA concentrations. (f) Correlations of iCLIP motif enrichment (4-mer) by 3 μ M RocA and motif prediction of 0.03 μ M RocA effect in RIP-Seq. The motifs shown in Figure 3b are highlighted. ρ : Spearman's rank correlation.



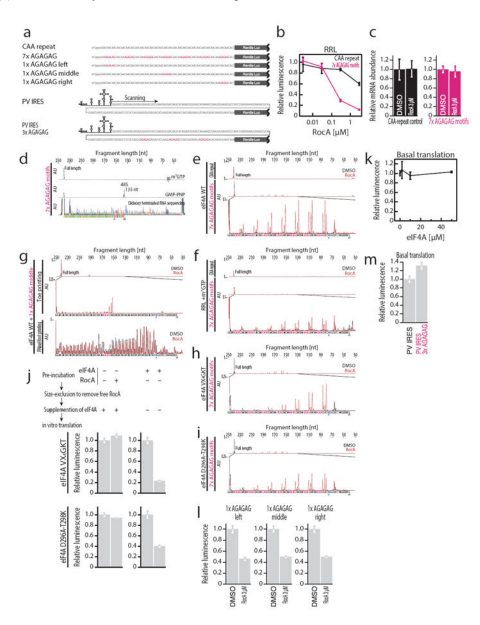
Extended Data Figure 8. eIF4A/RNA affinity measured by fluorescence polarization

(a) CBB staining of recombinant proteins used in this study.

(b) Summary of K_d between RNA and eIF4A among the conditions assayed. (c, e-g, i) Direct measurement of the eIF4A/RNA affinity by fluorescence polarization for eIF4A WT, eIF4A (VX₄GKT), or eIF4A (D296A-T298K) and 5' FAM-labeled RNAs in the presence or absence of RocA. Data represent mean and S.D. (n = 3).



(h) Pulldown assay with His-MBP-eIF4A expressed in E. coli and eIF4E/G in RRL.



Extended Data Figure 9. Characterization of toeprinting assay

(a) Diagram of the reporters used in this study.

(b, and c) In vitro translation in RRL with mRNAs containing seven polypurine motif

(AGAGAG) insertions (b) and qPCR from the samples (c).

(d) Dideoxy terminated sequencing of RNA by reverse transcription verified the toeprinting product length terminated by 48S ribosomes.

(e) Ribosome to eprinting assay performed in RRL in the presence of m7-GTP in the presence or absence of 3 μ M RocA treatment.

(f) Toeprinting assay using 10 μM recombinant eIF4A in the presence or absence of 10 μM RocA treatment.

(g) Toeprinting assay (top) and RNase I footprinting assay (bottom) using 10 μ M recombinant eIF4A with mRNA containing one AGAGAG motif at the middle in the presence or absence of 10 μ M RocA treatment.

(h and i) Toeprinting assay using 10 μ M recombinant eIF4A (VX₄GKT) or (D296A-T298K) with mRNA containing seven AGAGAG motifs in the presence or absence of 10 μ M RocA treatment.

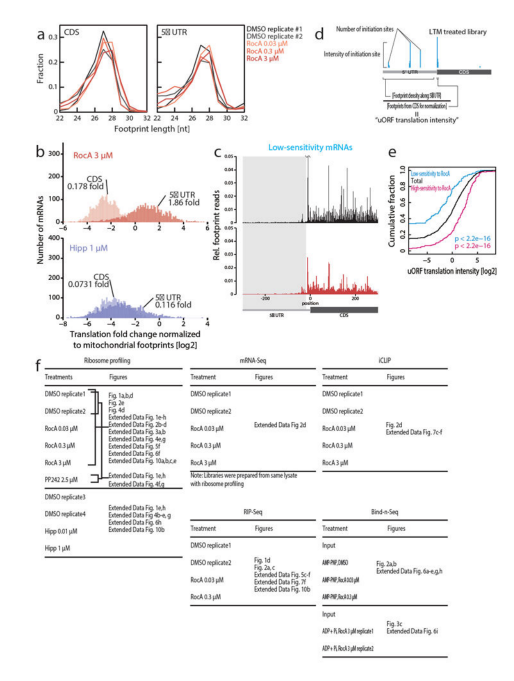
(j) Pre-formation of the complex with RocA and eIF4A (VX₄GKT) or (D296A-T298K) on the mRNA bearing seven polypurine motifs represses the translation from the mRNA in RRL.

(k) Basal translation level from mRNA containing seven AGAGAG with the supplementation of recombinant eIF4A.

(l) *In vitro* translation in RRL with mRNAs with single polypurine motif (AGAGAG) insertion at the different positions in 5' UTR

(m) Basal translation level from mRNAs bearing PV IRES and PV IRES with three AGAGAG.

In b-c and h-j, data represent mean and S.D. (n = 3).



Extended Data Figure 10. 5' UTR footprints accumulated in RocA treatments are come from uORFs $% \mathcal{A}$

(a) The distributions of specific footprint length, which is a hallmark of 80S ribosomes⁸, from CDS and 5' UTR are indistinguishable.

(b) The change in ribosome footprint counts for 5' UTRs and CDSs when cells are treated with 3 μ M RocA or 1 μ M Hipp compared to non-treatment, normalized to mitochondrial footprints. Median -fold change is shown. Bin width is 0.1. Analysis is restricted to mRNAs bearing footprints in the 5' UTR in the non-treatment condition.

(c) Meta-gene analysis of low-sensitivity transcripts to RocA. Reads are normalized to the sum of mitochondrial footprints reads.

(d) The illustration of the definition of uORF translation intensity.

(e) Transcripts sensitive to RocA contain more active uORFs, as measured by cumulative distributions of the uORF translation intensity (c). Significance is calculated by Mann-Whitney U test.

(f) The summary of deep sequencing based approaches used in this study and corresponding figures.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We are grateful to J. Tanaka for providing hippuristanol, to Y. Tomari for sharing DNA constructs, to H. Asahara and UC Berkeley DNA sequencing facility for the help of toeprinting assay, and to A. Pinder and F. Tan for support of deep sequencing analysis. We also thank the members of Ingolia, Lareau, and Tomari laboratories for faithful discussion and technical support. N.T.I. is a Damon-Runyon-Rachleff Innovator supported in part by the Damon Runyon Cancer Research Foundation (DRR-37-15), the Searle Scholars Program (11-SSP-229), and the National Institute of General Medical Sciences of the National Institutes of Health (P50GM102706). This work used the Vincent J. Coates Genomics Sequencing Laboratory at UC Berkeley, supported by NIH S10 Instrumentation Grants S10RR029668, S10RR027303, and OD018174. S.I. is a recipient of Human Frontier Science Program long-term fellowship. S.N.F. is a HHMI Fellow of the Helen Hay Whitney Foundation.

References

- Cencic R, et al. Antitumor activity and mechanism of action of the cyclopenta[b]benzofuran, silvestrol. PLoS One. 2009; 4:e5223. [PubMed: 19401772]
- 2. Bordeleau ME, et al. Therapeutic suppression of translation initiation modulates chemosensitivity in a mouse lymphoma model. J Clin Invest. 2008; 118:2651–2660. [PubMed: 18551192]
- Liu T, et al. Synthetic silvestrol analogues as potent and selective protein synthesis inhibitors. J Med Chem. 2012; 55:8859–8878. [PubMed: 23025805]
- 4. Santagata S, et al. Tight coordination of protein translation and HSF1 activation supports the anabolic malignant state. Science. 2013; 341:1238303. [PubMed: 23869022]
- Wolfe AL, et al. RNA G-quadruplexes cause eIF4A-dependent oncogene translation in cancer. Nature. 2014; 513:65–70. [PubMed: 25079319]
- Sadlish H, et al. Evidence for a functionally relevant rocaglamide binding site on the eIF4A-RNA complex. ACS Chem Biol. 2013
- Ingolia NT, Ghaemmaghami S, Newman JR, Weissman JS. Genome-wide analysis in vivo of translation with nucleotide resolution using ribosome profiling. Science. 2009; 324:218–223. [PubMed: 19213877]
- Ingolia NT, et al. Ribosome profiling reveals pervasive translation outside of annotated proteincoding genes. Cell Rep. 2014; 8:1365–1379. [PubMed: 25159147]
- Sonenberg N, Hinnebusch AG. Regulation of translation initiation in eukaryotes: mechanisms and biological targets. Cell. 2009; 136:731–745. [PubMed: 19239892]
- Pestova TV, Shatsky IN, Fletcher SP, Jackson RJ, Hellen CU. A prokaryotic-like mode of cytoplasmic eukaryotic ribosome binding to the initiation codon during internal translation initiation of hepatitis C and classical swine fever virus RNAs. Genes Dev. 1998; 12:67–83. [PubMed: 9420332]
- Rouskin S, Zubradt M, Washietl S, Kellis M, Weissman JS. Genome-wide probing of RNA structure reveals active unfolding of mRNA structures in vivo. Nature. 2014; 505:701–705. [PubMed: 24336214]
- 12. Bordeleau ME, et al. Functional characterization of IRESes by an inhibitor of the RNA helicase eIF4A. Nat Chem Biol. 2006; 2:213–220. [PubMed: 16532013]

- Lindqvist L, et al. Selective pharmacological targeting of a DEAD box RNA helicase. PLoS One. 2008; 3:e1583. [PubMed: 18270573]
- Hsieh AC, et al. The translational landscape of mTOR signalling steers cancer initiation and metastasis. Nature. 2012; 485:55–61. [PubMed: 22367541]
- Thoreen CC, et al. A unifying model for mTORC1-mediated regulation of mRNA translation. Nature. 2012; 485:109–113. [PubMed: 22552098]
- Lambert N, et al. RNA Bind-n-Seq: quantitative assessment of the sequence and structural binding specificity of RNA binding proteins. Mol Cell. 2014; 54:887–900. [PubMed: 24837674]
- 17. Linder P, Jankowsky E. From unwinding to clamping the DEAD box RNA helicase family. Nat Rev Mol Cell Biol. 2011; 12:505–516. [PubMed: 21779027]
- Abramson RD, et al. The ATP-dependent interaction of eukaryotic initiation factors with mRNA. J Biol Chem. 1987; 262:3826–3832. [PubMed: 2950099]
- König J, et al. iCLIP reveals the function of hnRNP particles in splicing at individual nucleotide resolution. Nat Struct Mol Biol. 2010; 17:909–915. [PubMed: 20601959]
- Pestova TV, Kolupaeva VG. The roles of individual eukaryotic translation initiation factors in ribosomal scanning and initiation codon selection. Genes Dev. 2002; 16:2906–2922. [PubMed: 12435632]
- Parsyan A, et al. mRNA helicases: the tacticians of translational control. Nat Rev Mol Cell Biol. 2011; 12:235–245. [PubMed: 21427765]
- 22. Pause A, Sonenberg N. Mutational analysis of a DEAD box RNA helicase: the mammalian translation initiation factor eIF-4A. EMBO J. 1992; 11:2643–2654. [PubMed: 1378397]
- Shah P, Ding Y, Niemczyk M, Kudla G, Plotkin JB. Rate-limiting steps in yeast protein translation. Cell. 2013; 153:1589–1601. [PubMed: 23791185]
- Shirokikh NE, et al. Quantitative analysis of ribosome-mRNA complexes at different translation stages. Nucleic Acids Res. 2010; 38:e15. [PubMed: 19910372]
- Balvay L, Soto Rifo R, Ricci EP, Decimo D, Ohlmann T. Structural and functional diversity of viral IRESes. Biochim Biophys Acta. 2009; 1789:542–557. [PubMed: 19632368]
- Kozak M. Downstream secondary structure facilitates recognition of initiator codons by eukaryotic ribosomes. Proc Natl Acad Sci U S A. 1990; 87:8301–8305. [PubMed: 2236042]
- 27. Medenbach J, Seiler M, Hentze MW. Translational control via protein-regulated upstream open reading frames. Cell. 2011; 145:902–913. [PubMed: 21663794]
- Arribere JA, Gilbert WV. Roles for transcript leaders in translation and mRNA decay revealed by transcript leader sequencing. Genome Res. 2013; 23:977–987. [PubMed: 23580730]
- 29. Lee S, et al. Global mapping of translation initiation sites in mammalian cells at single-nucleotide resolution. Proc Natl Acad Sci U S A. 2012; 109:E2424–32. [PubMed: 22927429]
- Pommier Y, Marchand C. Interfacial inhibitors: targeting macromolecular complexes. Nat Rev Drug Discov. 2012; 11:25–36. [PubMed: 22173432]
- Galicia-Vazquez G, Cencic R, Robert F, Agenor AQ, Pelletier J. A cellular response linking eIF4AI activity to eIF4AII transcription. RNA. 2012; 18:1373–1384. [PubMed: 22589333]
- Ingolia NT, Brar GA, Rouskin S, McGeachy AM, Weissman JS. The ribosome profiling strategy for monitoring translation in vivo by deep sequencing of ribosome-protected mRNA fragments. Nat Protoc. 2012; 7:1534–1550. [PubMed: 22836135]
- Anders S, Huber W. Differential expression analysis for sequence count data. Genome Biol. 2010; 11:R106. [PubMed: 20979621]
- 34. Lorenz R, et al. ViennaRNA Package 2.0. Algorithms Mol Biol. 2011; 6:26. [PubMed: 22115189]
- Dmitriev SE, Pisarev AV, Rubtsova MP, Dunaevsky YE, Shatsky IN. Conversion of 48S translation preinitiation complexes into 80S initiation complexes as revealed by toeprinting. FEBS Lett. 2003; 533:99–104. [PubMed: 12505166]
- Iwasaki S, et al. Hsc70/Hsp90 chaperone machinery mediates ATP-dependent RISC loading of small RNA duplexes. Mol Cell. 2010; 39:292–299. [PubMed: 20605501]
- Busso D, Delagoutte-Busso B, Moras D. Construction of a set Gateway-based destination vectors for high-throughput cloning and expression screening in *Escherichia coli*. Anal Biochem. 2005; 343:313–321. [PubMed: 15993367]

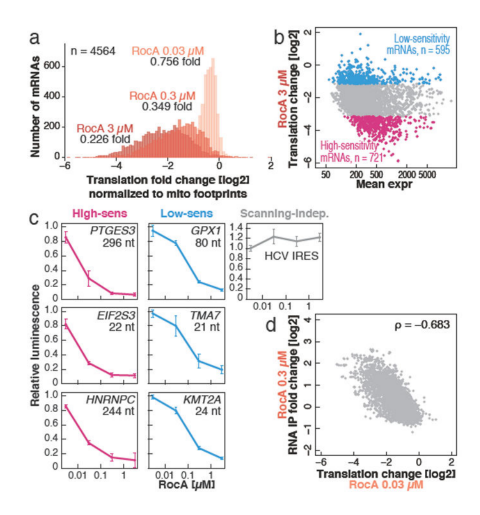


Figure 1. RNA sequence selectivity is imparted upon eIF4A by RocA causing selective translation repression

(a) Histogram of the number of transcripts along translation -fold change by ribosome profiling when cells are treated with 0.03, 0.3, or 3 μ M RocA, normalized to the number of mitochondrial footprints. Median -fold change is shown. Bin width is 0.1.

(b) MA plot of mean footprint reads between 3 μ M RocA treatment and non-treatment normalized to library sizes versus translation -fold change by 3 μ M RocA treatment, highlighting high-sensitivity and low-sensitivity mRNAs.

(c) The 5' UTRs of indicated genes were fused to *Renilla* luciferase and these reporter mRNAs were transfected prior to treatment with RocA as indicated. Data represent mean and standard deviation (S.D.) (n = 3).

(d) Correlation of translation -fold change to RIP -fold change with RocA treatment. ρ : Spearman's rank correlation.

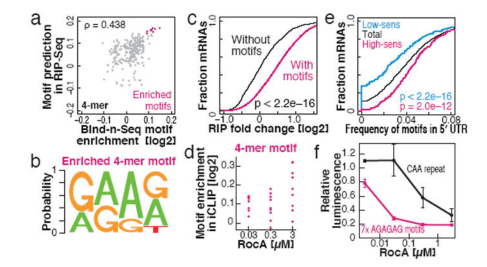


Figure 2. RNA Bind-n-Seq and iCLIP reveal that RocA preferentially increases the affinity between eIF4A and polypurine motif

(a) Correlations between 4-mer motif enrichment in Bind-n-Seq by 0.03 μ M RocA treatment and motif prediction of 0.03 μ M RocA effect in RIP-Seq. ρ : Spearman's rank correlation.

(b) Highest-scoring elements in Bind-n-Seq and RIP-Seq.

(c) The change in mRNA binding for mRNAs with or without the enriched 4-mer motif (b) in their 5' UTRs is shown as the RIP -fold change by RocA normalized to spike-in RNA. Significance is calculated by Mann-Whitney U test.

(d) Enrichment of 4-mer motifs (b) in iCLIP by RocA treatment relative to control DMSO treatment.

(e) The frequency of the 4-mer motif (b) in the 5' UTR predicts whether a mRNA is high- or low-sensitivity, based on the difference in cumulative distributions of motifs in the 5' UTR. Significance is calculated by Mann-Whitney U test.

(f) Reporter assay in HEK 293 cells with a CAA-repeat 5' UTR containing seven polypurine motif (AGAGAG) insertions (Extended Data Figure 9a). Data represent mean and S.D. (n = 3).

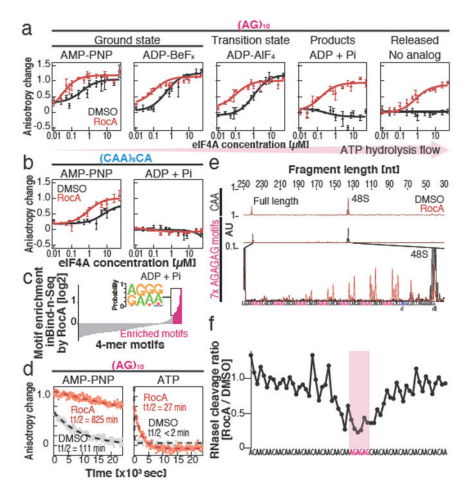


Figure 3. RocA clamps eIF4A on polypurine motif even after ATP hydrolysis (a, b) Direct measurement of the eIF4A/RNA affinity by fluorescence polarization for eIF4A and 5' FAM-labeled RNAs in the presence or absence of RocA. Data represent mean and S.D. (n = 3).

(c) Motif enrichments along entire 4-mer motifs in Bind-n-Seq with ADP + Pi and highestscoring elements (inset).

(d) Competition assay with unlabeled RNA. Data represent mean (n = 3).

(e) Ribosome to eprinting assay performed in RRL in the presence of GMP-PNP in the presence or absence of 3 μ M RocA treatment.

(f) Relative RNase I cleavage protected by eIF4A/RocA complex on mRNA containg one AGAGAG at the middle in footprinting assay. See the original data in Extended Data Figure 9f.

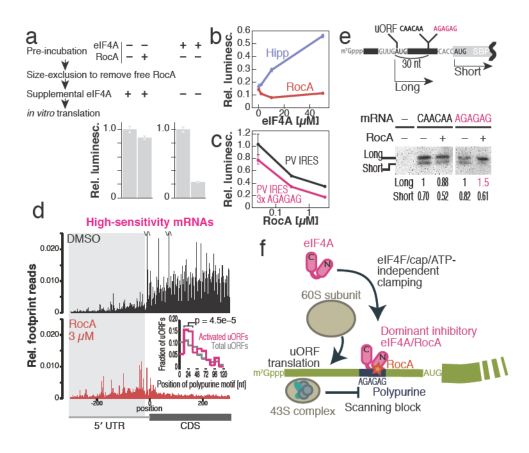


Figure 4. eIF4A/RocA complexes on polypurine motifs block scanning of pre-initiation complex, inducing uORF translation

(a) Pre-formation of the complex with RocA and eIF4A on the mRNA bearing seven polypurine motifs represses the translation from the mRNA in RRL.

(b) The supplementation of recombinant eIF4A protein to RRL *in vitro* transaltion reaction with 10 μ M Hipp or 3 μ M RocA.

(c) *In vitro* translation in RRL with mRNAs with native PV IRES and that with three polypurine motifs (Extended Data Figure 9a).

(d) Meta-gene analysis of high-sensitivity transcripts to RocA. Reads are normalized to the sum of mitochondrial footprints reads. Histogram of the position of the first polypurine motif (6-mer) after uORF initiation codon (inset). P value is calculated by Fisher's exact test. Bin width is 12 nt.

(e) Western blot of SBP translated from uORF and downstream major ORF in RRL with 0.03 μ M RocA treatment. Quantification of bands normalized to long form with DMSO treatment is shown. For gel source data, see Supplementary Fig. 1.

(f) Schematic representation of RocA-mediated translation control. RocA clamps eIF4A onto mRNA by selective affinity enhancement for a polypurine motif in eIF4F-, cap-, and ATP-independent manners, which then blocks scanning of pre-initiation complex, introducing premature translation from uORF and inhibiting downstream ORF translation. In b and c, data represent mean and S.D. (n = 3).