



**Randomized Codon Mutagenesis Reveals that the HIV Rev
Arginine-Rich Motif Is Robust to Substitutions and that
Double Substitution of Two Critical Residues Alters
Specificity**

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3 **1 Randomized Codon Mutagenesis Reveals that the HIV Rev Arginine-Rich Motif Is Robust to**
4 **2 Substitutions and that Double Substitution of Two Critical Residues Alters Specificity**
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1 ABSTRACT

2 The binding of the arginine-rich motif (ARM) of HIV Rev protein to its high-affinity site in stem
3 IIB in the Rev Response Element (RRE) initiates assembly of a ribonucleoprotein complex that
4 mediates the export of essential, incompletely spliced viral transcripts. Many biochemical, genetic,
5 and structural studies of Rev-RRE IIB have been published, yet the roles of many peptide residues
6 in Rev ARM are unconfirmed by mutagenesis. Rev aptamer I (RAI) is an optimized RRE IIB that
7 binds Rev with higher affinity and for which mutational data are sparse. Randomized-codon
8 libraries of Rev ARM were assayed for their ability to bind RRE IIB and RAI using a bacterial
9 reporter system based on bacteriophage λ N-nut antitermination. Most Rev ARM residues tolerated
10 substitutions without strong loss of binding to RRE IIB, and all except arginine 39 tolerated
11 substitution without strong loss of binding to RAI. The pattern of critical Rev residues is not the
12 same for RRE IIB and RAI, suggesting important differences between the interactions. The results
13 support and aid the interpretation of existing structural models. Observed clinical variation is
14 consistent with additional constraints on Rev mutation. By chance, we found double mutants of
15 two highly critical residues, arginine 35 (to glycine) and asparagine 40 (to valine or lysine), that
16 bind RRE IIB well, but not RAI. That an apparently distinct binding mode occurs with only two
17 mutations highlights the ability of arginine-rich motifs to evolve new recognition strategies and
18 supports the application of neutral theories of evolution to protein-RNA recognition.

19
20 **Keywords:** HIV Rev; Rev-Response Element; Rev Aptamer I; Protein-RNA recognition;
21 Arginine-rich motif; Neutral evolution.

22 INTRODUCTION

23 The human immunodeficiency virus (HIV) regulates the export of its singly and unspliced mRNAs
24 using the interaction between the virally encoded trans-activator protein, Rev, and a *cis*-acting,
25 ~350 nucleotide Rev Response Element (RRE), located in the *env* gene (Heaphy *et al.*, 1990;
26 Malim *et al.*, 1989; Zapp and Green, 1989). The 116 amino acid Rev protein includes an arginine-
27 rich motif (ARM) that binds to a high affinity site in stem IIB of RRE (Figure 1). A 17-mer Rev
28 peptide comprising residues 34-50 adopts an α -helical conformation and binds to RRE IIB with
29 high affinity and specificity *in vitro* (Battiste *et al.*, 1996; Kjems *et al.*, 1992; Tan *et al.*, 1993). The
30 Rev ARM binding site in RRE IIB includes an asymmetric internal loop with two purine:purine
31 base pairs (Bartel *et al.*, 1991; Hope *et al.*, 1990; Kjems *et al.*, 1992). Substitutions of Arg35,
32 Arg39, Asn40, and Arg44 to alanine and lysine (Tan *et al.*, 1993; Tan *et al.*, 1994) support NMR-
33 based structural models of Rev-RRE IIB (Battiste *et al.*, 1996) and Rev-RAI (Ye *et al.*, 1996), in
34 which these residues make base-specific contacts in the major groove (Figure 2). Though Trp45 is
35

1 not important for binding *in vitro*, it is required for essential Rev oligomerization *in vivo* (Kjems *et al.*, 1992; Zapp *et al.*, 1991). Despite intense investigation of the Rev-RRE IIB interaction, 2
3 mutational data are limited, and precise roles of specific ARM residues are uncertain. 4

5 Rev aptamer I (RAI) (Figure 1) was selected from a random RNA library based on a minimal RRE 6
7 IIB (Giver *et al.*, 1993). RAI shares substantial identity with RRE IIB and adopts a very similar 8
9 structure (Figure 2). The structural models show that the Rev peptide binds RAI similarly to RRE 10
11 IIB, though it has unique aspects including a U:A:U base triple in the Rev binding site (Ye *et al.*, 12
13 1996). RAI can replace RRE IIB functionally *in vivo* (Symensma *et al.*, 1996). The Rev-RAI 14
15 interaction has been examined far less than that of Rev-RRE IIB, and its tolerance to Rev-ARM 16
17 substitutions is unknown. 18
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23 Arginine-rich motifs binding small RNA structures occur in important biological systems, 24
25 including gene expression and its regulation (Chen and Varani, 2005). Because they are small, 26
27 structurally diverse, and occur in important systems, they serve as model systems to understand 28
29 RNA-protein recognition. (Draper, 1999). In addition to HIV Rev, examples of α -helical ARMs 30
31 include bacteriophage λ N (Legault *et al.*, 1998; Schärpf *et al.*, 2000) and its relatives P22 N (Cai *et al.*, 1998) and \square 21 N (Cilley and Williamson, 2003) that recognize small, boxB stem-loop RNAs 32
33 during the regulation of transcriptional antitermination. Important non- α -helical examples include 34
35 lentiviral Tat proteins that bind their small, stem-loop TAR RNAs (transactivating response 36
37 elements) during the regulation of transcriptional elongation (Ott *et al.*, 2011). HIV Tat binds its 38
39 target in an extended conformation (Long and Crothers, 1995), and its relative, bovine 40
41 immunodeficiency virus Tat (BIV Tat), binds its target in a β -turn conformation (Puglisi *et al.*, 42
43 1995). 44
45

46 In addition to RRE IIB and its variant RAI, unrelated Rev-binding RNAs (Xu and Ellington, 1996; 47
48 Ye *et al.*, 1999) and RRE IIB-binding peptides (Harada *et al.*, 1996; Harada *et al.*, 1997; Peled- 49
50 Zehavi *et al.*, 2003; Tan and Frankel, 1998) have been isolated by artificial selection. The 51
52 occurrence of diverse and specific recognition strategies raises questions as to how new recognition 53
54 strategies evolve: what mutations can occur without loss of function, how many different specific 55
56 interactions might exist, and what is the nature of evolutionary intermediates? Interestingly, the 57
58 conformational variability of proteins has been recently argued to facilitate evolvability (Tokuriki 59
60 and Tawfik, 2009).

1 Neutral theories of evolution (Kimura, 1991; Ohta, 2002) assert that most observed genetic change
2 is the result of the random fixation of selectively neutral mutations, rather than the result of
3 positive selection. When applied to molecular recognition, these theories predict that sequences are
4 sufficiently mutable that new recognition strategies can evolve by random genetic drift and
5 relaxed- and altered-specificity intermediates. Studies of arginine-rich peptide-RNA interactions,
6 including Rev-RRE relatives (Iwazaki *et al.*, 2005; Sugaya *et al.*, 2008a, Sugaya *et al.*, 2008b),
7 Tat-TAR (Smith *et al.*, 1998, Smith *et al.*, 2000), and P22N-boxB (Cocozaki *et al.*, 2008a;
8 Cocozaki *et al.*, 2008b), support these predictions by finding altered-specificity variants. Neutral
9 theories would predict that Rev ARM is sufficiently mutable such that distinct recognition
10 strategies are accessible by incremental changes without loss-of-fitness intermediates.

11
12 To improve understanding of the Rev-RRE IIB and Rev-RAI interactions, and to explore how well
13 neutral theories apply to RNA-protein interactions, we constructed randomized-codon libraries of
14 the Rev ARM and assayed them for their ability to bind RRE IIB and RAI using a bacterial
15 reporter system that recapitulates bacteriophage λ antitermination in *Escherichia coli* (Franklin,
16 1993). Arg39, Asn40, and Arg44 appear immutable with the RRE IIB reporter. Arg35 can be
17 substituted with the conservative, basic amino acid lysine with only moderate loss of activity. Most
18 residues are mutable, reflecting their less important role in recognition. In contrast to RRE IIB,
19 only Arg39 appears immutable with the RAI reporter, suggesting differences between RRE IIB and
20 RAI recognition strategies. The existing structural models are interpreted in light of results.
21 Clinical variation was compared to results and is consistent with the existence of additional
22 constraints on Rev mutation *in vivo*. Intriguingly, we found double mutants at two highly critical
23 residues, Arg35 (to glycine) and Asn40 (to valine or lysine), that bind RRE IIB well, but not RAI.
24 That an apparently distinct binding mode occurs with only two mutations highlights the ability of
25 arginine-rich motifs to evolve new recognition strategies and supports the application of neutral
26 theories of molecular evolution to small peptide-RNA recognition.

27 28 **MATERIALS AND METHODS**

29 **General**

30 Restriction enzymes and T4 DNA ligase were obtained from Roche (Germany). Bacterial media
31 components were obtained from Oxoid (UK). Fine chemicals were obtained from Amersham
32 (UK), Sigma (USA), and Amresco (USA). Laboratory chemicals were obtained from Acros
33 (Belgium). Disposable plasticware was obtained from Sarstedt (Germany).

1 Strains and plasmids

2 *Escherichia coli* supporting antitermination, N567 (Franklin and Doelling, 1989), pACnutTAT13,
3 and pBRptac-N* (Franklin, 1993), were obtained from Naomi Franklin (University of Utah).
4 Plasmids expressing the Rev14 N fusion, (pBRN-HIVRev14), the BIV Tat N fusion (pBRN-
5 BIVTat), and the NMR RRE reporter (pAC-HIV-RRE) (Harada *et al.*, 1996) were obtained from
6 Kazuo Harada (Tokyo Gakugei University).
7

8 Construction of libraries and mutants

9 Plasmids expressing the Rev17 N fusion and reporting on RRE IIB binding, RAI binding, and BIV
10 TAR binding were constructed in house. Rev17 and its libraries were cloned into pBRptac-N* as
11 NcoI-BsmI fragments replacing the amino terminus of λ N (residues 1–19). Rev17 and library
12 fragments were generated by mutually priming oligonucleotides followed by NcoI-BsmI digestion
13 using oligonucleotide pairs based on the Rev17 coding strand 5'-CC ATG GCA ACC CGC CAG
14 GCC CGT CGT AAC CGT AGA CGT CGT TGG CGT GAG CGT CAG CGT GCA GCT GCG
15 GCG AAT GCA-3'. RRE IIB, RAI, and BIV TAR reporters were cloned into pACnutTAT13
16 replacing the λ left *nut* site oligonucleotides based on the coding strand sequences flanked by PstI
17 and BamHI sites: 5'-CTG CAG TCG ACG CTC TTA AAA ATT AAG GTC TGG GCG CAG
18 CGT CAA TGA CGC TGA CGG TAC AGG CCA GCA TTC AAA GCA GGG ATC C-3' for
19 RRE IIB, 5'-CTG CAG TCG ACG CTC TTA AAA ATT AAG CTC GTG TAG CTC ATT AGC
20 TCC GAG CAG CAT TCA AAG CAG GGA TCC-3' for RAI, and 5'-CTG CAG TCG ACG CTC
21 TTA AAA ATT AAG CTC GTG TAG CTC ATT AGC TCC GAG CAG CAT TCA AAG CAG
22 GGA TCC-3' for BIV TAR. Standard molecular biology procedures were used, and all clones were
23 tested by function. Entire synthetic inserts were confirmed by sequencing using PBRNR2 (5'-GGC
24 TTG CTG TAC CAT GTG-3') for N-fusion constructs and PACF2 (5'-AAT CAC TGC ATA ATT
25 CGT GTC-3') for RNA-reporter constructs.
26

27 Library screening with X-gal

28 Competent N567 host cells carrying reporter plasmids were transformed with the library or clones
29 of interest and control plasmids. Approximately 10-100 ng of plasmid per 100 μ l of competent
30 cells were transformed by heat shock and plated on tryptone plates containing 50 μ g/ml ampicillin
31 and 15 μ g/ml chloramphenicol as antibiotics, 80 μ g/ml X-gal (5-bromo-4-chloro-3-indolyl- β -D-
32 galactoside) as the chromogenic substrate of the β -galactosidase reporter protein, and 0.05 mM
33 IPTG (isopropyl β -D-thiogalactoside) to induce the tac promoters expressing N protein and the

1 reporter transcript. The plates were scored and photographed after one day at 34°C and after a
2 second day at 24°C.

4 **ARM-RNA recognition assays with ONPG**

5 For each interaction, from 3 to 6 representative colonies were picked from X-gal plates for solution
6 assays. For measurement of ARM-RNA recognition via N-mediated antitermination, cultures were
7 grown overnight at 30°C with aeration in tryptone with 50 µg/ml ampicillin and 15 µg/ml
8 chloramphenicol as antibiotics, and with 0.05 mM IPTG. The cells were then permeabilized,
9 assayed for β-galactosidase activity using ONPG (*ortho*-nitrophenol-β-D-galactoside), and units of
10 β-galactosidase were calculated following Miller (1992). Percent activities are reported normalized
11 to the activity of Rev17 for RRE and RAI reporters and BIV Tat for the BIV TAR reporter of that
12 day's experiment.

14 **Structure visualization**

15 Jmol (<http://www.jmol.org/>), an open-source Java viewer for chemical structures in 3D, was used
16 to view NMR structures of Rev-RRE (Protein Data Bank 1ETF and 1ETG; Battiste *et al.*, 1996)
17 and Rev-RAI (Protein Data Bank 1ULL; Ye *et al.*, 1996).

19 **RESULTS**

20 **Anti-termination assay reflects Rev-RRE IIB and Rev-RAI binding**

21 A protein-RNA reporter system (Franklin, 1993) based on bacteriophage λ N-nut antitermination
22 has been previously exploited for screening and assay of heterologous interactions, including Rev-
23 RRE IIB and relatives (Harada *et al.*, 1996; Harada *et al.*, 1997; Iwazaki *et al.*, 2005; Peled-Zehavi
24 *et al.*, 2003; Sugaya *et al.*, 2008a. Sugaya *et al.*, 2008b). The system consists of two plasmids
25 transformed into *Escherichia coli*. One plasmid is an N-fusion supplier plasmid in which the RNA-
26 binding peptide of interest is fused to amino terminus of λ N protein, replacing that of N. The other
27 plasmid is a boxB reporter plasmid, in which the RNA of interest replaces the boxB hairpin in the λ
28 *nut* (N-utilization) site upstream from transcriptional terminators. Expression of the downstream β-
29 galactosidase gene reflects binding of the peptide-RNA interaction by N-nut mediated
30 antitermination. To choose the context for library screening, we first compared the activity of two
31 different Rev sequences, Rev14 and Rev17 (Table 1), expressed as fusions to the activation domain
32 of N, and three reporters, in which the Rev-binding RNAs replaces boxB in the reporter transcript
33 (Figure 1A). Rev14 is Rev 34-47, used previously (Harada *et al.*, 1996; Peled-Zehavi *et al.*, 2003).
34 Rev17 is Rev34-50, also used previously (Battiste *et al.*, 1996; Tan *et al.*, 1993; Tan and Frankel,

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3 1 1994). NMR RRE is the high affinity site in RRE IIB (C44-G53:C65-G76) truncated and capped
4 2 by a tetraloop, previously used for structural studies and in previous screens (Harada *et al.*, 1996,
5 3 Harada *et al.* 1997). RRE IIB is RRE C44- G76 with the apical 9 nt of the wild-type stem loop
6 4 intact. RAI is Rev Aptamer I, used for NMR studies (Ye *et al.*, 1996). An N supplier and a nut
7 5 reporter for the heterologous BIV Tat-TAR interaction served as a specificity control (Harada *et*
8 6 *al.*, 1996). Based on solution assays indicating increased activity with Rev17 and RRE IIB
9 7 constructs without loss of specificity, they were chosen as the basis for library construction and
10 8 screening, and they were used for all further experiments (Table 1).
11 9

10 **Rev Arginine-Rich Domain Is Robust to Substitutions**

11 A set of 15 plasmid libraries, spanning Rev Thr34 to Arg48, each expressing a Rev17-N fusion
12 12 with one Rev ARM codon randomized, was constructed via cassette mutagenesis, and transformed
13 13 into reporter cells on plates containing the chromogenic substrate X-gal. The proportion of each
14 14 library transformation appearing to have wild-type activity by deposition of blue pigment was
15 15 estimated by eye (Figure 4). Computer-assisted quantification of RRE IIB plate images produced
16 16 similar results to those estimated by eye (data not shown). The proportion of positives ranged from
17 17 less than 1% for Asn40 to 50% for Arg36, Arg 41, and Glu47. The existing structural model of
18 18 Rev-RRE posits essential roles for base-specific contacts by four residues: Arg35, Arg39, Asn40,
19 19 and Arg44 (Battiste *et al.*, 1996), and the results of Figure 2A accord with these roles, though the
20 20 Arg35 library screen has more positives than the 6/64 (~9%) that would be expected for only
21 21 arginine codons. Some libraries had more positives than was expected from published mutational
22 22 analysis, including as Thr34, Arg38, and Arg46 (Tan *et al.*, 1993; Tan and Frankel, 1994).
23 23 Mutation of Ala37 had been neglected previously, yet its library contained relatively few positives.
24 24
25 25 Screening the libraries with the RAI reporter revealed a different pattern, in which all libraries
26 26 contain more positives than with the RRE IIB reporter, and only Arg39 appeared to be poorly
27 27 tolerant of substitution. Notably, the two most restrictive Rev positions when screened against RRE
28 28 IIB, Asn40 and Arg44, have far higher proportions of positives when screened against RAI,
29 29 suggesting differences between Rev-RRE IIB and Rev-RAI interactions. No library had positives
30 30 when screened against BIV TAR reporter cells.
31 31

32 **Activities of Single Mutations Support and Refine Existing Structural Models**

33 To determine the effect of specific mutations, colonies displaying wild-type activity were selected
34 34 by eye from each screen, grown individually, and pooled for plasmid preparations. The N-supplier

1 plasmid was separated from the reporter plasmid by gel electrophoresis and transformed into *E.*
2 *coli* cells. Colonies were selected randomly and restreaked, and plasmids were prepared and
3 individually tested against reporters. Clones were sequenced and measured for activity by solution
4 assay against Rev, RAI, and BIV TAR reporters (Table 2).

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6 As expected from the base-specific contacts observed in the Rev-RRE model (Battiste *et al.*, 1996),
7 published data on mutants (Tan *et al.*, 1993; Tan and Frankel, 1994), and the proportion of active
8 library members (Figure 4), no functional mutants of Arg39, Asn40, and Arg44 were found, and
9 only lysine could replace Arg35, with moderately reduced activity (Table 2). The average Rev-RAI
10 structural model has a similar core structure to Rev-RRE. Using mutational data of Rev-RRE to
11 support the Rev-RAI structural model, Ye *et al.* (1996) posit base-specific contacts by Arg35,
12 Arg39, Asn40, and Arg44. Screening the Rev libraries against RAI, a markedly higher proportion
13 of active members was noted. The pattern of restrictive residues appears distinct: Arg39 is
14 immutable with RAI, Arg44 intermediate, and Asn40 tolerant to substitution.

15
16 In the average Rev-RRE model, Arg39 is deep in the major groove within hydrogen-bonding
17 distance of G70 (Figure 2). The Rev-RAI model shows hydrogen bonds between the Arg39 epsilon
18 and η -nitrogens to the equivalent of G70 O6 and G70 N7, respectively, and structure 3 of the Rev-
19 RRE PDB file shows a similar orientation. We recovered only arginine from RRE IIB and RAI
20 screens. Though the proportion of positives in the Arg39 library was consistent with no mutation
21 being functional, we prepared the R39K construct to determine to whether the positive charge or
22 the guanidium group of Arg39 was important. The poor activity of R39K with both RRE II B and
23 RAI reporters (Table 2) strongly supports the proposed archetypal arginine-guanine interaction
24 between Arg39 and G70 being responsible for the critical role of Arg39.

25
26 In the Rev-RRE model, Asn40 is within hydrogen-bonding distance and nearly co-planar with the
27 G47:A73 base pair (Figure2). The Rev-RAI average structure shows Asn40's carboxamide
28 donating a hydrogen bond from its nitrogen to the O6 of the G47 equivalent and accepting a
29 hydrogen bond from its oxygen to N6 of the A73 equivalent. The Asn40 library had the lowest
30 proportion of active mutants, and no active single substitution was found in screens against the
31 RRE IIB reporter. Substitutions with arginine, leucine, valine, and glycine had only background
32 activity. Glutamine was not recovered, despite sequencing 16 clones, strongly supporting the
33 critical role of Asn40 hydrogen bonds to G47:A73 in the Rev-RRE interaction.

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3 1 The Rev-RRE model positions Arg44 in the major groove close to U45 and G46, though the side
4 2 chain is not well resolved. The Rev-RAI model is similarly resolved and suggests hydrogen bonds
5 3 to the equivalent U45 O4 and G46, though the published analysis only discusses the interaction
6 4 with the U45 (Ye *et al.*, 1996). Individual structures of both interactions suggest hydrogen bonding
7 5 to either or both U45 O4 and G46 O6, though the contact to U45 O4 is most frequent. We observed
8 6 a low proportion of active library members recovered only arginine from RRE IIB screens, and
9 7 found R44K to have moderate activity with RRE IIB. The results support the critical role of Arg44
10 8 as donating hydrogen bonds to U45.
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13 10 The Rev-RRE model has Arg35 within hydrogen bonding distance of U66 and G67. Individual
14 11 Rev-RRE structures and the average Rev-RAI structure suggest η -nitrogens of Arg35 donate
15 12 hydrogen bonds to O6 and N7 of the equivalent to G67. Individual Rev-RRE structures
16 13 infrequently suggest an Arg35 hydrogen bond to O4 of U66. The RAI equivalent of U66 is a
17 14 cytosine lacking an O4, preventing the Rev-RAI interaction from supporting the Arg35-U66
18 15 interaction. The observation that more than 10% of library members were active in RRE IIB
19 16 screens suggested functional substitutions existed, and indeed, R35K was recovered from screens
20 17 with RRE IIB, though with moderately reduced activity. Interestingly, R35S showed activity well
21 18 above background with the RRE IIB reporter, consistent with a critical role for the positive charge
22 19 of Arg35 and the donation of at least one hydrogen bond to G67.
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26 21 In contrast to literature reports that Thr34 was important (Tan *et al.*, 1993), and the Rev-RRE and
27 22 Rev-RAI models suggesting important contact between the Thr34 hydroxyl to the 5'-phosphate of
28 23 G47, we observed a proportion of the library active with RRE IIB and moderate to high activity of
29 24 smaller amino acids, including glycine and proline (Table 2). Interestingly, most individual Rev-
30 25 RAI structures and some Rev-RRE structures suggest the backbone nitrogen of the terminal Thr34
31 26 making the phosphate contact. The recovery of T34N and T34G from screening and the relatively
32 27 high activity of these mutants, including T34P, conflicts with published gel shift data showing
33 28 T34A had low affinity (Tan *et al.*, 1993). The T34A mutant was constructed in hopes of resolving
34 29 this discrepancy, but it too showed more than 50% activity. Our data suggest that the hydroxyl
35 30 contact is not important, though the Thr34 amide could be, and that steric occlusion of the side
36 31 chain limits Thr34 mutation to smaller residues. Why the activity of the T34A mutant might differ
37 32 between our system and a gel shift assay is not clear. The effect of chemical modification of the
38 33 G47 phosphate (Kjems *et al.*, 1992) could be a reflection of steric crowding rather than a hydrogen
39 34 bond with Thr34. The activity of several small residues unable to make productive contacts with
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1 phosphate, especially T34G, suggest that the threonine side chain is either not making a hydrogen
2 bond to G47 phosphate, or that it contributes little to binding.

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4 Arg46 is suggested to stabilize the concentration of three phosphates near RRE U72 (Battiste *et al.*,
5 1996). As a result of the alterations to the equivalent region of RAI caused by the formation of the
6 A:U:A base triple, Arg46 is near only two phosphates. That our library had approximately a quarter
7 of its members active with RRE IIB suggested that Arg46 may not be absolutely required, but we
8 found no strongly active substitutions. R46K, which would be capable of favorable electrostatic
9 interactions with phosphates, had only a quarter of the activity of wild-type Rev with RRE IIB.
10 Arg46 appears relatively important, supporting its role in stabilizing the charge of three phosphates
11 in the Rev-RRE IIB interaction, though it appears to contribute to recognition less than that of the
12 residues contacting bases.

13
14 The Rev-RRE model shows Arg38 close to the phosphates 5' to U66 and G67 and proposed to
15 donate hydrogen bonds to them. From RRE IIB screens, we recovered R38K and R38L with only
16 moderately reduced activity (Table 2). Why R38A and R38K severely decreased gel shift affinity
17 (Tan *et al.*, 1993) and reporter activation (Tan and Frankel, 1994), respectively, is unclear. Similar
18 to Thr34, the difference in context may be responsible, and the varying activity of multiple mutants
19 is consistent with it having a relatively minor role in RRE recognition

20
21 In the Rev-RRE model, Arg 41, Arg42, Arg43, and Arg48 are positioned to interact with backbone
22 phosphates. Arg48 is seen with Gln49 contacting the non-ARM Asn26 in a Rev crystal structure
23 (Daugherty *et al.*, 2010). Screening Arg41, Arg42, Arg43, and Arg48 libraries with RRE IIB and
24 RAI reporters indicated active mutants, and a variety of moderately active substitutions were
25 recovered. Our data suggest that these arginines make minor contributions to affinity consistent
26 with roles contacting backbone phosphates, and they are individually dispensable for Rev-ARM
27 binding to RRE IIB. The moderate to high activities of R38L, R41L, and R42L and their adjacent
28 locations in the complex suggest favorable van der Waals interactions between the arginine
29 aliphatic chains.

30
31 Overlooked in previous mutational studies, Ala37 appears to be restricted in the Rev-RRE
32 interaction to smaller amino acids. The models suggest it is sterically confined between the peptide
33 and RNA, making van der Waals contacts to a backbone ribose. Consistent with steric
34 confinement, the proportion of positives in the Ala37 library was low, and only A37S had even

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3 1 moderate activity with RRE IIB. Though Ala37 may not be making significant enthalpic
4 2 contributions, it could have a role in stabilizing the bound structure conformationally, and it is
5 3 likely few substitutions would be allowed.
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10 5 Gln36 is in the major groove, and the Rev-RRE model suggests it could make van der Waals
11 6 contacts to G47 and G48 (Battiste *et al.*, 1996). Gel shift data indicate that it can be substituted
12 7 with alanine (Tan *et al.*, 1993). As expected, the Gln36 library had a large proportion of positives,
13 8 and active substitutions were recovered. Its position in the major groove could allow its
14 9 carboxamide nitrogen to donate a hydrogen bond to the N7 of G48. Interestingly, only one mutant
15 10 in all libraries was found to have significantly higher than wild type activity with RRE IIB, Q36S.
16 11 It is conceivable that the high activity of Q36S with RRE IIB and RAI is the result of a new,
17 12 favorable contact between the serine hydroxyl and N7 of G48 and its equivalent in RAI.
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24 14 As expected from the distance between Glu47 and the RNA, its negative charge, and gel shift
25 15 affinity unaffected by substitution (Tan *et al.*, 1993), the Glu47 library had a large proportion of
26 16 positives. Many different substitutions had activities similar to or higher than the wild type
27 17 sequence. Glu47 does not appear to have a role in recognition. Though Trp45 has an essential role
28 18 in Rev function *in vivo* (Zapp *et al.*, 1991), mutational data *in vitro* and the Rev-RRE model
29 19 indicate it has little role in recognition, though it could to make van der Waals contact to A68. As
30 20 expected, a high proportion of its library was active in screens with both RRE IIB and RAI
31 21 reporters, and a wide variety of substitutions were found to have near wild-type activity.
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23 **Compensatory mutations of critical residues suggest a distinct recognition strategy nearby**

24 24 By chance, the first eight active clones isolated from the Asn40 library screen with RRE IIB
25 25 included four clones representing double substitutions, three of R35G-N40V and one of R35G-
26 26 N40K. The Arg35 codon in the Asn40 library should be CGC, and the double mutants had glycine
27 27 GGC codons. Presumably, active single substitutions in the Asn40 library were so rare that that
28 28 active double substitutions arising from single nucleotide errors in the Asn40 library
29 29 oligonucleotide synthesis were relatively common. Both double mutants displayed moderate
30 30 activity with RRE IIB (Table 3). We prepared the single substitution N40V construct, and assayed
31 31 it for activity, which was as expected minimal, as were single substitutions R35G and N40L (Table
32 32 2).
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1 To pursue these observations, a library randomized at both Arg35 and Asn40 was constructed and
2 screened with RRE IIB. The double substitution library displayed a much higher proportion that
3 did the Asn40 library (~4% vs 0.4%, see online supporting material Figures S1 and S2). Nine
4 sequences were isolated, of which five were R35G-N40V, two were R35G-N40L, and two were
5 R35G-N40A. No wild type sequences were recovered. The double substitutions were assayed for
6 activity (Table 3). That functional double mutants of two critical residues making base-specific
7 contacts restore activity when replaced by the absence of a side chain and aliphatic side chains
8 argues that the double mutant binds RRE using an alternate recognition strategy dependent on a
9 glycine at position 35 and independent of hydrogen bonding side chains at position 40.
10 Interestingly, R35G-N40L maintained a much greater activity on RRE IIB compared to wild type
11 compared to RAI, which was not observed of any single substitution, demonstrating significant
12 differences between RRE IIB and RAI specificities.

13 14 **Clinical Variation Suggests Constraints on Rev Genetic Drift Other Than Binding**

15 Extraction of HIV Rev basic domain (Thr34-Arg48) sequences from the 2009 Los Alamos HIV
16 Compendium (Kuiken *et al.*, 2009) followed by filtering of incomplete and stop-codon-containing
17 variants yield 1379 entries comprising 129 distinct sequences. With regard to only the composition
18 of sequences, the number of substitutions relative to our basis sequence, HXB2, indicates several
19 interesting features (Table 4). Very few substitutions are seen at most positions. Trp45, required for
20 multimerization, appears least frequently mutated. Critical residues that make base-specific
21 contacts, Arg35, Asn40, and Arg44, have the least number of substitutions. The critical residue
22 Arg39 has a large number of R39K substitutions, but otherwise has as few substitutions as other
23 critical, base-contacting residues. Slightly more substitutions are seen at arginines that do not
24 contact bases: Arg28, Arg41, Arg42, Arg43, Arg46, Arg48. Relatively more substitutions are seen
25 at non-arginines that we found mutable: Thr34, Gln36, Ala37, Glu47. Many frequent substitutions
26 appear quite conservative, such as T34S, A37T, R38Q, R39K, R41Q, R43K, R44K, R46K, and
27 R48K. Our basis sequence for Rev-ARM, HXB2, is not the most common: E47A constitutes more
28 than half of residue 47 entries.

29
30 The frequency of R39K was surprising, because our data shows this mutant has very low activity.
31 Further analysis showed that of the 515 entries with R39K, none were single mutations. Indeed,
32 excluding those of Glu47, there were only 18 single substitutions in the 34-48 region: the
33 remainder were multiple substitutions. Of the 515 R39K entries, 498 are also E47A. This
34 frequency suggested covariation, in which the replacement of the negatively charged Glu37 with an

1 uncharged residue compensated for mutation of Arg39 to Lys. We prepared and assayed the R39K-
2 E47A construct and found it weakly active, yet well above the background activity of R39K (Table
3 3).

4 5 **DISCUSSION**

6 Noting incomplete mutational data in support of the Rev-RRE and Rev-RAI structural models, we
7 constructed 15 libraries containing all single substitutions in Rev35-48 and screened them for
8 activity using a well-established protein-RNA reporter system (Cocozaki *et al.*, 2008b; Franklin,
9 1993; Harada *et al.*, 1996; Iwazaki *et al.*, 2005). Our results show that Rev basic domain residues
10 range from invariant to highly mutable when binding RRE IIB. The data support the Rev-RRE
11 structural model with minor refinements, consistent with critical, base-specific contacts by Arg35,
12 Arg39, Asn40, and Arg44. The Rev-RAI interaction appears far more mutable than that of Rev-
13 RRE and is absolutely dependent on only one critical residue, Arg39, suggesting differences
14 between Rev-RRE IIB and Rev-RAI recognition. New findings include the activity of R35K with
15 RRE IIB, the unexpected mutability of Thr34, the restricted mutability of Ala37, and the fortuitous
16 discovery of active double mutants of Arg35-Asn40. Comparison to variation in clinical sequences
17 is consistent with Rev mutation being limited by factors other than binding RRE IIB. The
18 mutability of Rev with different partners and the activity of the double mutant support the
19 application of neutral theories of evolution to small peptide-RNA recognition.

20 21 **Rev-RRE and Rev-RAI structural models**

22 Supported by published gel shifts of alanine substitutions (Tan *et al.*, 1993) and heterologous
23 reporter assays of arginine-to-lysine substitutions (Tan and Frankel, 1994), the NMR-based
24 structural models of Rev-RRE (Battiste *et al.*, 1996) and Rev-RAI (Ye *et al.*, 1996) present four
25 residues making important base-specific contacts, Arg35, Arg39, Asn40, and Arg44, while the
26 backbone amide or hydroxyl of Thr34 and remaining arginines contact phosphates. Ala36 and
27 Trp45 are shown making van der Waals contacts, and Glu47 does not contribute to binding. It is
28 tempting to imagine well-defined interactions in which each of the critical residues makes ideal
29 hydrogen bonds in plane to their base pair partners. The mutability of Rev seen here, the
30 conformational flexibility of Rev arginines observed in Rev-RRE IIB and Rev-RAI complexes
31 (Wilkinson *et al.*, 2004), and adaptive binding of partners in these complexes (Battiste *et al.*, 1994,
32 Ye *et al.*, 1996), suggest that less perfect interactions and even an ensemble of minor variations is
33 likely.

1 The similarity of the Rev-RRE and Rev-RAI structural models promotes the assumption they are
2 identical in all important aspects. How similar are the two interactions, and how much can we infer
3 about one interaction from the other? We found the Rev ARM to be highly mutable with the RAI
4 reporter, consistent with Rev-RAI having more intermolecular contacts, perhaps unsurprisingly as
5 it is the product of artificial selection for affinity, whereas the natural Rev-RRE interaction may
6 have evolved under selection for biologically relevant rates of binding and unbinding in addition to
7 specificity. The structures have important differences. The U:A:U triple in RAI introduces a large
8 aberration in the backbone that repositions phosphates and more encloses the peptide, possibly
9 facilitating more or better contacts. RAI, lacking the equivalent of RRE U66, cannot form a
10 hydrogen bond between U66 O4 and Arg35, which was found to be important for Rev binding by
11 conservation in selection experiments (Bartel *et al.*, 1991) and footprinting and modification-
12 interference experiments (Kjems *et al.*, 1992). Counter-intuitively, arginines in Rev ARM are more
13 dynamic in the higher affinity Rev-RAI interaction than in Rev-RRE IIB (Wilkinson *et al.*, 2004).

14
15 The two interactions' determinants of specificity, while overlapping, are not identical. Only Arg39
16 of the critical Rev-RRE residues Arg35, Arg39, Asn40, and Arg44, is absolutely required for RAI
17 binding, obscuring the defining pattern of the Rev-RRE recognition strategy. Rev substitutions
18 throughout the peptide decreased binding to RRE IIB at least 10-fold more than RAI: A37L,
19 R38G, N40L, R42S, R43L, R44K, and R46S (Table 2). Though no Rev single substitution was
20 found to increase binding to RRE IIB markedly more than RAI, the Rev double mutant R35G-
21 N40L maintained 10-fold more activity to RRE than to RAI. Thus, our results do not cast doubt on
22 Arg35, Asn40, and Arg44 making base-specific contacts to RAI, but they do show that the
23 recognition strategy of Rev-RAI, though structurally similar, has a distinct specificity.

24 25 **Clinical sequence variation**

26 To what extent does the mutability of the Rev basic domain seen in our binding assay reflect its
27 potential mutability in the wild? Though our reporter system has been found to accord with affinity
28 measurements *in vitro* (Harada *et al.*, 1996; Harada *et al.*, 1997; Iwazaki *et al.*, 2005; Sugaya *et al.*,
29 2008b), N-mediated antitermination relies on bacterial host factors unrelated to the host factors
30 involved in Rev-mediated nuclear export in human cells. Compared to our results, the clinical data
31 have very little variability, though the profile is similar in that critical residues have the lowest
32 frequency of mutations, and unimportant residues have the highest, with the exception of Arg39
33 (Table 4). The very low levels of substitution and our isolation of few mutations conferring higher-
34 than-wild-type activity are consistent with purifying selection operating against weaker binders. In

1 agreement with this, Glu47 is the most variable residue in the clinic, it has no clear RNA-binding
2 role being negatively charged and projecting away from the RNA, and we isolated many functional
3 mutants. In the clinic, the conservative mutation R39K is very common (Table 4), though we found
4 the R39K mutant had only background activity with both RRE IIB and RAI reporters (Table 2).
5 This specific discrepancy between clinical sequences and our assays is ameliorated by the
6 observation that the vast majority of R39K occur in conjunction with E47A and the moderately
7 restored activity of our R39K-E47A construct (Table 3). Interestingly, neither R35G nor N40V
8 were found in the database (Table 4). Thus, Rev ARM does display some diversity of recognition
9 strategy *in vivo*, though much less than found in our reporter system. In addition maintaining
10 affinity to RRE IIB, there are likely strong constraints on Rev ARM evolution *in vivo*, including
11 the overlapping open reading frames of *env* and *tat* and its function as a nuclear localization signal.
12

13 Evolution of recognition

14 How different is the recognition strategy of R35G-N40V? How are the loss of critical, base-
15 specific contacts made by Arg35 and Asn40 compensated? Energetically, there should be new
16 contacts to compensate for the loss of hydrogen bonds. The replacement of the charged
17 guanidinium group of Arg35 with a proton suggests deeper penetration of the peptide and
18 conformational flexibility. The replacement of the carboxamide of Asn40 with an aliphatic group
19 suggests burial of hydrophobic surface by extensive van der Waals interactions. We argue that
20 specificity to RRE in preference to RAI upon mutation of two critical residues is most consistent
21 with the double mutant binding using a distinct recognition strategy, though more physical data or
22 computational modeling will be needed understand how structurally similar the interaction is to
23 Rev-RRE.

24
25 Proteins are observed to be quite tolerate of random substitution (Guo *et al.*, 2004; Bloom *et al.*,
26 2005). Neutral theories of evolution (Kimura, 1991; Ohta, 2002;) assert that most observed genetic
27 change is neutral or nearly neutral, such that genetic drift can lead to new phenotypes without
28 positive selection. RNA has been used as a model to study neutral theories, its sequence being its
29 genotype and its folding being its phenotype (Fontana, 2002). Intersection sequences (Schultes and
30 Bartel, 2000), RNAs able to fold into different structures, would allow transitions to new
31 phenotypes. Similar to intersection sequences, chameleon peptides (Minor and Kim, 1996) are able
32 to adopt different conformations in different contexts, possibly facilitating evolutionary transitions
33 and modeling how alternate structural conformers facilitate protein evolution (Tokuriki and
34 Tawfik, 2009). When applied to small peptide-RNA recognition such as the interaction between

1 Rev ARM and RRE IIB, neutral theories would explain the evolution of new recognition strategies
2 by incremental genetic drift along paths of active mutations of one strategy until paths originating
3 from different strategy are encountered, at which point transition can occur. Thus, neutral theories
4 predict peptide and RNA partners will be robust (Wilke and Adami, 2003), robustness being the
5 ability to incur mutations without loss of activity, and that there are other recognition strategies
6 nearby.

7
8 We see a range of alterations to the Rev-RRE recognition strategy: the minor variation of R39K-
9 E47A, a somewhat different strategy employed by Rev-RAI, and an apparently distinct strategy
10 employed by the Rev R35G-N40V double mutant when binding RRE IIB. Many ARM-RNA
11 variations have been isolated from random libraries (Bayer *et al.*, 2005; Das and Frankel, 2003;
12 Harada *et al.*, 1996; Harada *et al.*, 1997; Peled-Zehavi *et al.*, 2003; Tan and Frankel, 1998; Xu and
13 Ellington, 1996; Ye *et al.*, 1999). These variations illustrate the potential evolution of recognition
14 strategies. The R39K-E47A variation may represent a simple energetic compensation in which the
15 decreased affinity caused by the disruption of a guanidinium hydrogen bond is restored by the
16 removal of a distal, destabilizing negative charge. Rev-RAI may represent a variation in which Rev
17 makes sufficient additional contacts that the otherwise critical residues Arg35, Asn40, and Arg44
18 are individually dispensable. The robustness of Rev-RAI suggest that the recognition strategy of
19 Rev-RAI, possibly very similar to Rev-RRE in structure, is poised between being largely identical
20 and being able to access new recognition strategies with very few mutations. One can imagine new
21 recognition strategies easily arising by incremental mutations that repeatedly acquire new contacts
22 and dispense of old contacts, thus maintaining binding while specificity alters.

23
24 Our findings support neutral theories, both by the robustness of Rev ARM to substitutions, and by
25 the discovery of a distinct recognition strategy with only two Rev mutations, R35G and N40V. Are
26 there examples from related systems supporting the possibility that Rev R35G-N40V binds in a
27 distinct conformation? In an example of chameleonism, JDV Tat adopts distinct strategies to bind
28 both BIV TAR and HIV-1 TAR, in β -turn and extended conformations, respectively (Smith *et al.*,
29 2000). Hybrid boxB RNAs are able to bind two normally type-specific N proteins in
30 bacteriophages λ and P22 (Cocozaki *et al.*, 2008a), and a single substitution relaxed P22N ARM
31 specificity such that it complemented both N-deficient λ and P22 viruses *in vivo* (Cocozaki *et al.*,
32 2008b).

33 34 CONCLUSION

1 Our findings that Rev can access increasingly variant recognition strategies resonates with other
2 studies on the diversity and adaptability of the Rev and RRE and viral ARM-RNA interactions
3 (Williamson, 2000). Importantly, Rev binds other sites on RRE than stem IIB (Daugherty *et al.*,
4 2008), and a Rev binds a non-RAI aptamer (Xu and Ellington, 1996) in an extended, non-helical
5 conformation (Ye *et al.*, 1999). Using the N-boxB antitermination system applied here, structurally
6 diverse peptides have been isolated that bind RRE IIB (Harada *et al.*, 1996; Peled-Zehavi *et al.*,
7 2003). Mutational analyses and further selections have shown how few mutations in ARMs or RRE
8 can increase affinity and alter specificity (Harada *et al.*, 1997, Iwazaki *et al.*, 2005; Sugaya *et al.*,
9 2008a; Sugaya *et al.*, 2008b). Thus, there are many different specificities accessible, often with few
10 structural changes. The adaptive binding (Battiste *et al.*, 1994; Patel, 1999; Ye *et al.*, 1996) and the
11 conformational flexibility of Rev arginines (Wilkinson *et al.*, 2004) observed in Rev-RRE and
12 Rev-RAI complexes may reflect requirements of specificity (Leulliot and Varani, 2001), but they
13 may also cooperate with the mutability of Rev reported herein to allow multiple partners and
14 binding modes that facilitate evolutionary transitions.

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6

7 **FIGURE LEGENDS**

8 **Figure 1.** Rev-binding RNAs and Rev peptide. (A) Sequence and secondary structures of the high-
9 affinity site in RRE used in structural studies (NMR RRE), the longer site used in this study (RRE
10 IIB), and the Rev aptamer I used in structural studies (RAI). Wild-type nucleotides are bold, and
11 the added GNRA tetraloop in NMR RRE is lower case. Boxed residues in RRE IIB have been
12 identified as important by biochemical and mutational studies. Numbered RRE IIB bases contact
13 Rev. Numbers are standard to RRE studies and refer to placement in the entire RRE element
14 (Malim *et al.*, 1989). (B) The sequence of the Rev peptide fusion to the N activation domain is
15 shown with HIV Rev residues numbered. Randomized codon libraries were made of residues 34-
16 48, shown bold.

17
18 **Figure 2.** Structural Models of Rev-RRE IIB and Rev-RAI. NMR models of (A) HIV Rev-RRE
19 IIB (1ETF) and (B) Rev-RAI (1ULL) are shown to the right with RNA white cartoon, peptide
20 backbone gray wireframe, and side chains of critical residues Arg35, Arg39, Asn40, and Arg44
21 colored by atom type. Views and renderings to the right are identical, except RNAs are rendered at
22 van der Waals radii. Critical side-chain labels are placed level to their locations.

23
24 **Figure 3.** Experiment Approach. The arginine-rich motif of HIV Rev (residues 34-50) replaces the
25 ARM of λ N such that Rev-ARM is fused to the activation domain of λ N and flanked by
26 methionine-alanine at the amino terminus and 4 alanines at the fusion junction. Oligonucleotide
27 libraries, coding for Rev-ARM sequences in which codons of residues 34-48 are individually
28 randomized, were inserted into N-fusion supplier plasmids and screened for functional members by
29 by transformation into *E. coli* hosting RNA reporter plasmids. When the N-fusion protein
30 containing a Rev-ARM mutant binds RRE IIB or RAI in the *nut* site of the reporter transcript, the
31 N activation domain recruits host factors to assemble an anti-termination complex causing
32 downstream transcriptional terminators to be ignored and expression of β -galactosidase from the
33 LacZ reporter gene. β -galactosidase activity can be monitored colorimetrically in colonies with
34 Xgal plates and quantified with ONPG in extracts of cultures. The BIV Tat-TAR interaction placed
35 in this reporter system serves as a heterologous control for specificity.

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4 **Figure 4.** Rev Library Activities Against RRE IIB and RAI. Estimated proportions of Rev library
5 members displaying wild-type activity in X-gal plate screens with RRE IIB (A) and RAI (B)
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8 reporters are shown as histograms, by Rev residue.
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2 **Table 1.** Antitermination activities of ARM-RNA interactions.

| N-fusion ^a | Sequences ^b | β-gal units ^c | | | | percent activation ^d | | | |
|-----------------------|------------------------|--------------------------|----------------------|------------------|-------------------|---------------------------------|----------------------|------------------|-------------------|
| | | NMR RRE ^e | RRE IIB ^f | RAI ^g | BTAR ^h | NMR RRE ^e | RRE IIB ^f | RAI ^g | BTAR ^h |
| Rev14 ⁱ | -TRQARRNRRRRWRR- | 46±8 | 130±30 | 470±90 | 2±0.4 | 46±8 | 57±13 | 101±20 | 4.2±0.7 |
| Rev17 ^j | -TRQARRNRRRRWRERQR- | 100±15 | 230±30 | 460±70 | 2±0.6 | 100±15 | 100±14 | 100±16 | 4±1.1 |
| BTat ^k | MGRPRGTRGKRRIRRGGGNA | 1±0.07 | 2±0.06 | 2±0.4 | 60±10 | 1.37±0.07 | 0.7±0.03 | 0.51±0.09 | 100±18 |

3 ^a Laboratory names of N-fusions expressed by pBRN plasmids.4 ^b The sequences of ARM fused to N. See footnotes h, i, j.5 ^c Plasmids were transformed into reporter cells with the named RNA in place of λ *nut* site boxB. At least four replicates of each clone were
6 grown at 30°C overnight in tryptone medium supplemented with 50 μM IPTG and assayed for β-galactosidase activity with ONPG.7 ^d Percent activation represents antitermination activities of N-fusions normalized to Rev17-NMR RRE, Rev17-RRE IIB Rev17-RAI, or BIV Tat-
8 TAR.9 ^e NMR RRE = 5'-GGUCUGGGCGCAGCGCAAGCUGACGGUACAGGCC-3'10 ^f RRE IIB = 5'-GGUCUGGGCGCAGCGUCA AUGACGCUGACGGUACAGGCC-3'11 ^g RAI = 5'-GGCUGGACUCGGACUUCGGUACUGGAGAAACAGGCC-3'12 ^h BTAR = 5'-GCUCGUGUAGCUCAUUAAGCUCCGAGC13 ⁱ Rev14 = MA TRQARRNRRRRWRR AAAA14 ^j Rev17 = MA TRQARRNRRRRWRERQR AAAA15 ^k BTat = MG RPRGTRGKRRIRR GGGNAAN

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Table 2. Activities of single Rev substitutions.

| Clone ^a | Rev17 N-fusion ^b | percent activation ^c | | |
|----------------------|-----------------------------|---------------------------------|-----------|----------|
| | | RRE IIB | RAI | BTAR |
| II820-2 | T34S | 114±15 | 148±26 | 2.9±0.4 |
| V635-1 | T34N | 68±7 | 129±28 | 2.5±0.4 |
| V658-2 | T34G | 93±9 | 158±39 | 1.8±0.4 |
| II752-1 | T34P | 66±7 | 134±30 | 3.1±0.4 |
| V635-2 | T34L | 38±2 | 117±25 | 2.2±0.5 |
| II922-3 ^d | T34A | 66±9 | 141±16 | 2±0.5 |
| V635-3 | R35K | 49±10 | 33±6 | 1.8±0.3 |
| V715-1 | R35S | 17±5 | 70±13 | 5.4±0.8 |
| V696-1 | R35G | 5.5±0.5 | 16±4 | 5±0.4 |
| V635-5 | Q36S | 308±42 | 172±21 | 1.7±0.11 |
| II750-2 | Q36L | 70±4 | 124±23 | 3.9±0.4 |
| V658-6 | A37S | 48±7 | 163±15 | 1.8±0.5 |
| II778-2 | A37C | 2.3±0.1 | 74±6 | 2.0±0.5 |
| II778-1 | A37L | 2.0±0.2 | 64±9 | 2.8±0.4 |
| II778-5 | A37V | 5.3±0.4 | 43±8 | 2.2±0.6 |
| II752-10 | R38K | 74±15 | 208±35 | 2.7±0.5 |
| V658-10 | R38L | 52±20 | 157±24 | 1.0±0.2 |
| II752-9 | R38E | 29±2 | 127±27 | 2.5±0.4 |
| II751-2 | R38G | 4.1±0.3 | 90±8 | 2.0±0.2 |
| II922-1 ^d | R39K | 2.2±0.4 | 0.73±0.25 | 2±0.2 |
| V701-1 | N40L | 0.8±0.3 | 119±21 | 2.2±0.2 |
| V701-7 | N40R | 1.2±0.2 | 58±13 | 6.5±0.6 |
| V696-10 | N40G | 0.68±0.2 | 41±6 | 5±1.0 |
| II945-1 ^d | N40V | 0.72±0.14 | 12±3 | 1.9±0.2 |
| II751-4 | R41L | 46±5 | 146±40 | 1.5±0.3 |
| 752-14 | R41I | 24±4 | 147±25 | 1.4±0.4 |
| V635-15 | R41S | 28±2 | 205±68 | 1.6±0.3 |
| II909-2 ^d | R41K | 82±6 | 135±36 | 2±0.2 |
| V635-18 | R42L | 80±13 | 189±29 | 1.6±0.2 |
| II747-18 | R42M | 64±6 | 187±12 | 2.3±0.06 |
| V635-17 | R42A | 60±7 | 199±55 | 1.6±0.11 |
| V658-17 | R42F | 40±6 | 128±36 | 1.7±0.11 |
| II752-16 | R42S | 11±6 | 150±22 | 1.7±0.3 |
| II747-20 | R43K | 55±10 | 92±25 | 2.7±0.9 |
| V635-20 | R43S | 63±5 | 137±25 | 2.0±0.1 |
| V696-13 | R43T | 14±2 | 92±12 | 3.0±2 |

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|----------|------|-----------|--------|----------|
| V715-5 | R43L | 3±0.7 | 42±4 | 3.0±0.9 |
| II755-5 | R44K | 0.88±0.11 | 44±13 | 1.5±0.4 |
| V635-23 | W45Q | 117±22 | 134±26 | 2.9±0.6 |
| II755-6 | W45L | 103±20 | 108±28 | 2.3±0.6 |
| II747-24 | W45N | 127±31 | 118±9 | 1.6±0.14 |
| II752-20 | W45H | 80±6 | 115±27 | 2.3±1 |
| V658-23 | W45A | 101±11 | 117±17 | 1.6±0.2 |
| II752-19 | W45T | 81±13 | 103±13 | 2.6±0.5 |
| V635-24 | W45S | 71±7 | 107±15 | 2.3±0.6 |
| V635-25 | R46K | 24±2 | 50±9 | 3.2±0.9 |
| V635-26 | R46L | 10±1 | 117±20 | 2.5±0.9 |
| II751-6 | R46A | 5±1 | 109±9 | 2.9±0.8 |
| II755-7 | R46S | 4±1 | 91±22 | 1.9±0.1 |
| V696-15 | E47V | 136±15 | 101±17 | 1±0.1 |
| V635-28 | E47M | 142±37 | 96±24 | 2.8±0.5 |
| V696-14 | E47R | 150±22 | 107±16 | 4.0±1.5 |
| V696-19 | E47C | 134±10 | 95±21 | 2.1±0.2 |
| V635-27 | E47N | 63±13 | 102±26 | 2.5±0.7 |
| II752-24 | E47S | 120±27 | 117±14 | 3.1±1.1 |
| II752-23 | E47I | 78±7 | 58±7 | 2.7±0.6 |
| V715-12 | E47A | 121±31 | 56±5 | 3.3±1.5 |
| V635-29 | R48K | 75±12 | 77±11 | 2.5±1.3 |
| V751-1 | R48Q | 21±4 | 39±12 | 1.9±0.2 |
| II755-9 | R48H | 50±11 | 98±16 | 2.1±0.3 |
| II747-30 | R48A | 35±5 | 87±9 | 2.1±0.4 |

^a Laboratory stock numbers of single mutants clones in Rev17 N-fusion.

^b Rev mutants are indicated by single-letter codes for wild type and mutant residues separated by the numerical position in Rev.

^c Library plasmids were transformed into RRE IIB and BTAR reporter cells. At least four replicates of each clone were grown at 30°C overnight in tryptone medium supplemented with IPTG and assayed for β-galactosidase activity with ONPG. Percent activation represents antitermination activities of Rev mutants normalized to Rev17-RRE, Rev17-RAI, or BIV Tat-TAR.

^d Synthetic constructs.

1 **Table 3.** Activity of Rev double mutants.

| Clone ^a | Mutants ^b | Sequences ^c | percent activation ^d | | |
|-----------------------|----------------------|--------------------------|---------------------------------|----------|---------|
| | | | RRE IIB | RAI | BTAR |
| V795-4 | R35G; N40V | TGQARR <u>V</u> RRRRWRER | 46±1.0 | 10±0.95 | 1.9±0.2 |
| V696-11 | R35G; N40K | TGQARR <u>K</u> RRRRWRER | 40±4.0 | 8.9±2.1 | 1.9±0.2 |
| V795-1 | R35G; N40L | TGQARR <u>L</u> RRRRWRER | 31±1.4 | 3.1±0.35 | 2.0±0.5 |
| II1026-1 ^e | R39K; E47A | TRQARK <u>N</u> RRRRWRAR | 9.3±1.4 | 11±3.4 | 2.2±0.4 |

2 ^a Laboratory stock numbers of Rev17 N-fusions.

3 ^b Rev mutants are indicated by single-letter codes for wild type and mutant residues separated by the numbered position.

4 ^c The sequences of Rev mutants with mutations indicated by underlining.

5 ^d Plasmid clones were transformed into RRE IIB, RAI, and BTAR reporter cells. At least four replicates of each clone were grown at 30°C
6 overnight in tryptone medium supplemented with IPTG and assayed for β-galactosidase activity with ONPG. Percent activation represents
7 antitermination activities of Rev mutants normalized to Rev17-RRE, Rev17-RAI, or BIV Tat-TAR.

8 ^e Synthetic construct.

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1 **Table 4.** Composition of Rev 34-48 in clinical HIV sequences.

| Codon ^b | Clinical variation of HIV-1 compared to HXB2 Rev 35-48 ^a | | | | | | | | | | | | | | |
|--------------------|---|-------------|-------------|-------------|-------------|------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|------------------|-------------|
| | T34 | R35 | Q36 | A37 | R38 | R39 | N40 | R41 | R42 | R43 | R44 | W45 | R46 | E47 | R48 |
| Ala | 2 | | 2 | 1308 | | | | 1 | 1 | | | | | 987 ^d | |
| Arg | | 1369 | 10 | | 1313 | 856 | 1 | 1357 | 1369 | 1362 | 1373 | 2 | 1364 | 31 | 1350 |
| Asn | 2 | | 14 | | | | 1370 | | | | | | 1 | | |
| Asp | | | | | | | 1 | | | | | | | 1 | |
| Cys | | | | | | | | | | | | | | | |
| Gln | | 3 | 1328 | | 56 | 1 | 1 | 14 | | | | | | 41 | |
| Glu | | | | | | | | | | | 1 | | | 285 | |
| Gly | 1 | | | | 1 | | | 2 | 1 | 3 | 1 | | 3 | 2 | 5 |
| His | | | | | | | 1 | | | | | | | | |
| Ile | | | | | | | 1 | | | | | | 1 | | |
| Leu | | | | | 1 | | | | | | | 1 | | | |
| Lys | | 2 | 12 | | 1 | 515 | 2 | 2 | 4 | 13 | 4 | | 9 | 19 | 23 |
| Met | | | | | | | | | | | | | | | |
| Phe | | | | | | | | | | | | | | | |
| Pro | 4 | 4 | 1 | | 1 | 1 | 1 | 4 | | | | | | | |
| Ser | 142 | | 10 | 8 | 1 | 1 | 1 | | | 1 | | | 1 | 1 | 1 |
| Thr | 1228 | 1 | 2 | 63 | | 5 | | | 4 | | | | | 12 | |
| Trp | | | | | 5 | | | | | | | 1376 | | | |
| Tyr | | | | | | | 1 | | | | | | | | |
| Val | | | | | | | | | | | | | | | |
| Total ^c | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 | 1379 |

^a The Los Alamos HIV sequence database (Kuiken *et al.*, 2009) was queried for Rev34-48 as nucleotides aligning to 8402-8447 of the reference genome HXB2. Incomplete, artifact, and stop-codon containing sequences were removed, and the 1379 remaining sequences were analyzed. The total number of each amino acid at each position, indicated by single-letter code for amino acid and location in Rev protein, was counted, and the number referring to the identity of HXB2 at that position is in bold face type.

^b The 20 possible amino acids are represented by standard three-letter codes.

^c The total number of sequences analyzed is shown as a checksum.

^d E47A is more common than the reference HXB2 sequence.

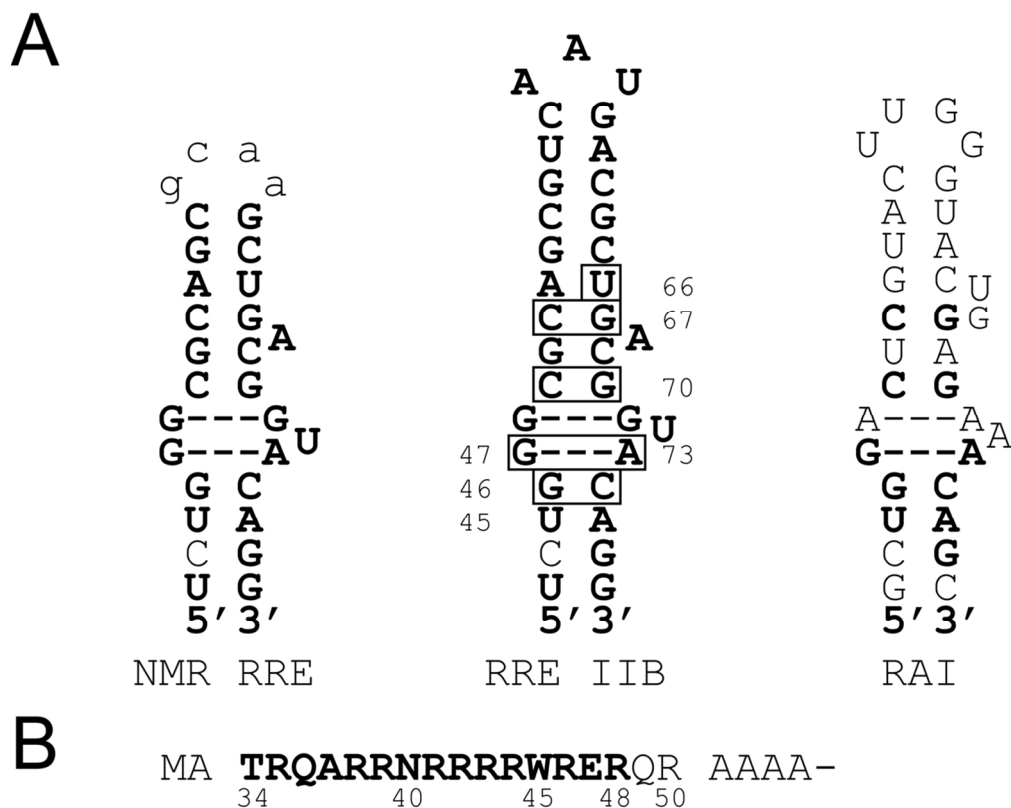


Figure 1. Rev-binding RNAs and Rev peptide. (A) Sequence and secondary structures of the high-affinity site in RRE used in structural studies (NMR RRE), the longer site used in this study (RRE IIB), and the Rev aptamer I used in structural studies (RAI). Wild-type nucleotides are bold, and the added GNRA tetraloop in NMR RRE is lower case. Boxed residues in RRE IIB have been identified as important by biochemical and mutational studies. Numbered RRE IIB bases contact Rev. Numbers are standard to RRE studies and refer to placement in the entire RRE element (Malim et al., 1989). (B) The sequence of the Rev peptide fusion to the N activation domain is shown with HIV Rev residues numbered. Randomized codon libraries were made of residues 34-48, shown bold.

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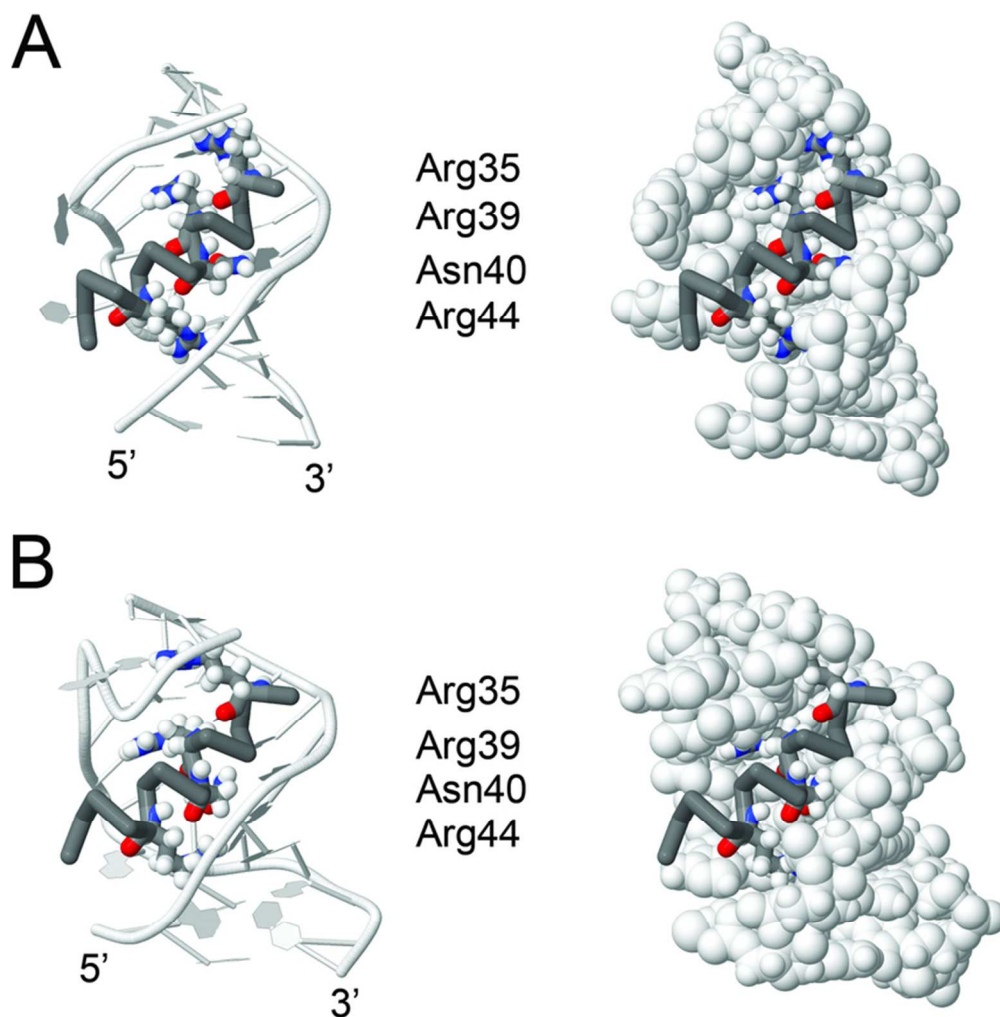


Figure 2. Structural Models of Rev-RRE IIB and Rev-RAI. NMR models of (A) HIV Rev-RRE IIB (1ETF) and (B) Rev-RAI (1ULL) are shown to the right with RNA white cartoon, peptide backbone gray wireframe, and side chains of critical residues Arg35, Arg39, Asn40, and Arg44 colored by atom type. Views and renderings to the right are identical, except RNAs are rendered at van der Waals radii. Critical side-chain labels are placed level to their locations.
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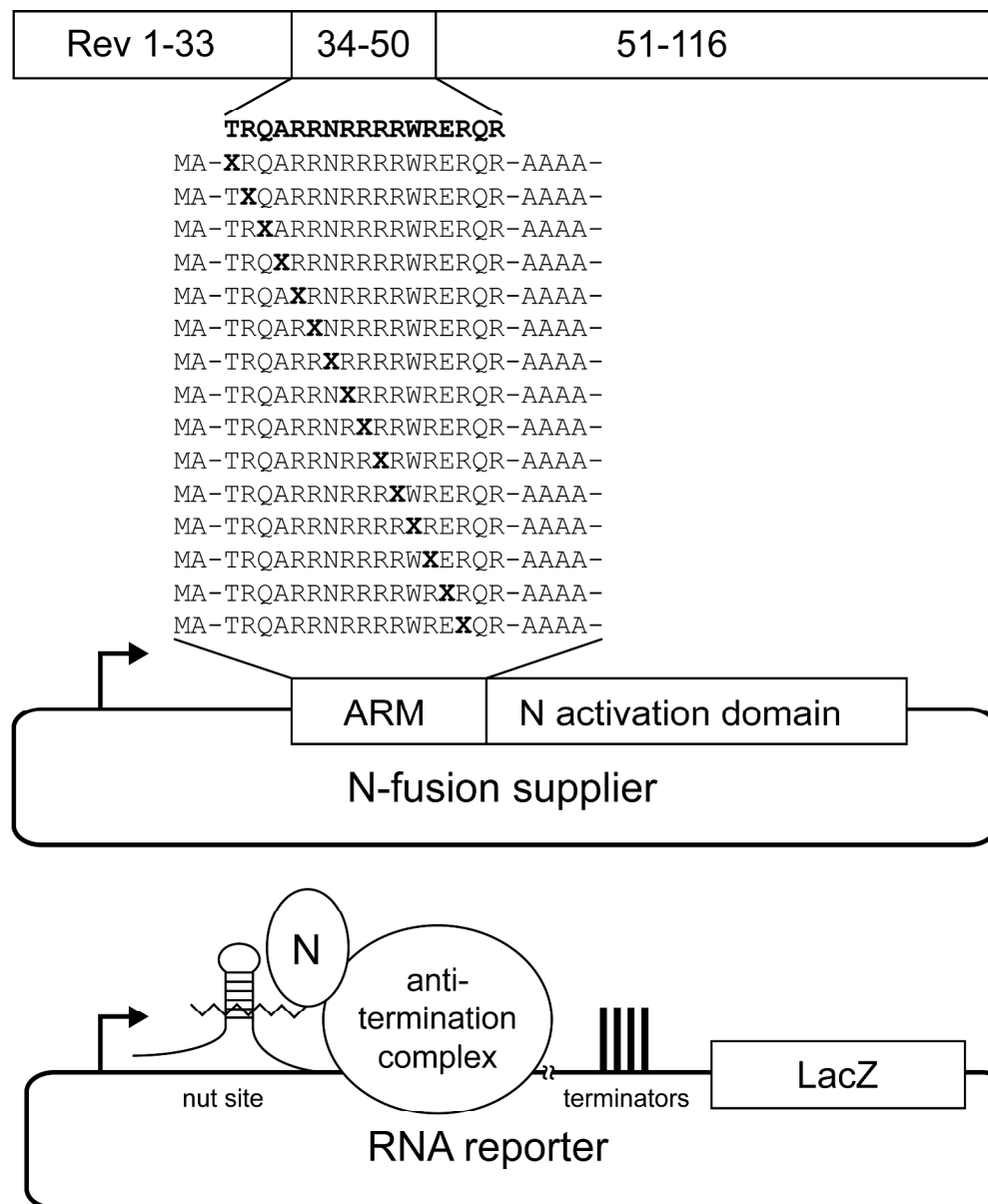


Figure 3. Experiment Approach. The arginine-rich motif of HIV Rev (residues 34-50) replaces the ARM of λ N such that Rev-ARM is fused to the activation domain of λ N and flanked by methionine-alanine at the amino terminus and 4 alanines at the fusion junction. Oligonucleotide libraries, coding for Rev-ARM sequences in which codons of residues 34-48 are individually randomized, were inserted into N-fusion supplier plasmids and screened for functional members by transformation into *E. coli* hosting RNA reporter plasmids. When the N-fusion protein containing a Rev-ARM mutant binds RRE IIB or RAI in the nut site of the reporter transcript, the N activation domain recruits host factors to assemble an anti-termination complex causing downstream transcriptional terminators to be ignored and expression of β -galactosidase from the LacZ reporter gene. β -galactosidase activity can be monitored colorimetrically in colonies with Xgal plates and quantified with ONPG in extracts of cultures. The BIV Tat-TAR interaction placed in this reporter system serves as a heterologous control for specificity.

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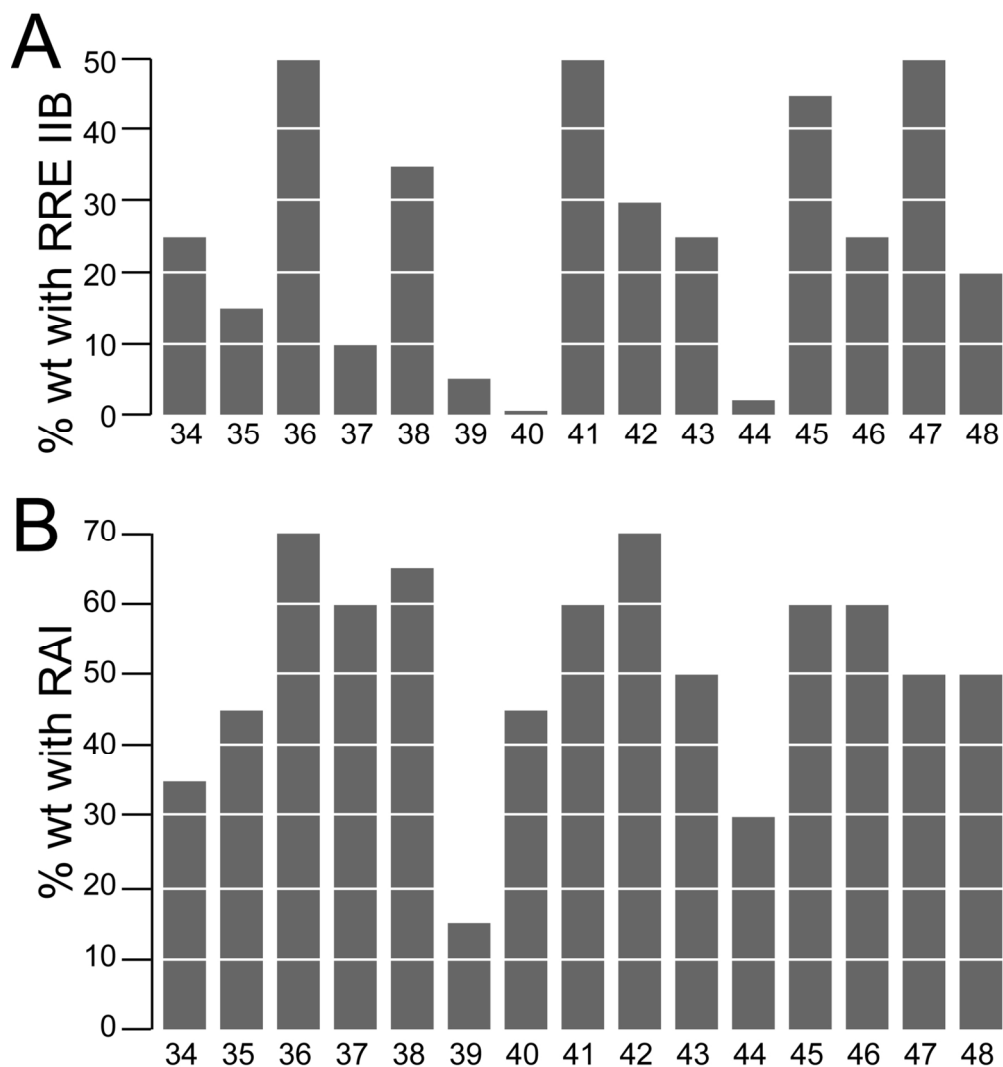


Figure 4. Rev Library Activities Against RRE IIB and RAI. Estimated proportions of Rev library members displaying wild-type activity in X-gal plate screens with RRE IIB (A) and RAI (B) reporters are shown as histograms, by Rev residue.
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