

Context and conformation dictate function of a transcription antitermination switch

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In bacteriophage λ , transcription elongation is regulated by the N protein, which binds a nascent mRNA hairpin (termed *boxB*) and enables RNA polymerase to read through distal terminators. We have examined the structure, energetics and *in vivo* function of a number of N-*boxB* complexes derived from *in vitro* protein selection. Trp18 fully stacks on the RNA loop in the wild-type structure, and can become partially or completely unstacked when the sequence context is changed three or four residues away, resulting in a recognition interface in which the best binding residues depend on the sequence context. Notably, *in vivo* antitermination activity correlates with the presence of a stacked aromatic residue at position 18, but not with N-*boxB* binding affinity. Our work demonstrates that RNA polymerase responds to subtle conformational changes in *cis*-acting regulatory complexes and that approximation of components is not sufficient to generate a fully functional transcription switch.

The regulation of transcription elongation is important in both prokaryotic and eukaryotic gene regulation^{1–5}. However, much less is known about the regulation of elongation than of initiation, even in classical systems such as bacteriophage λ , in which termination⁶ and elongation regulation were first described^{7,8}. The N protein is a key regulator of transcription elongation in phage λ ^{9–11}, interacting with (i) the nascent mRNA (ii) the transcription factor NusA and (iii) RNA polymerase, despite being only 107 amino acids long and totally unfolded in solution (Fig. 1a,b)¹². N increases the processivity of RNA polymerase, allowing it to read through both intrinsic and Rho-dependent termination signals^{9–11}, and initiates this antitermination activity by specifically binding the nascent mRNA at the *boxB* stem loop within the sequence known as *nut* (N-utilization)^{13,14} (Fig. 1b,c). Despite recent crystallographic work describing prokaryotic transcription machinery¹⁵, various features of this antitermination switch remain mysterious. For example, controversy exists over whether approximation of components in the antitermination machinery is sufficient for function¹⁶ or whether the detailed conformations of *cis*-acting elements¹⁷ have any role.

The RNA-binding domain of the λ N(1–22) peptide folds into a bent α -helix upon binding the *boxB* RNA and enforces formation of a GNRA tetraloop fold with one base extruded (Fig. 1c)^{17–21}. In the complex, Trp18 stacks on top of the tetraloop fold, effectively extending the RNA π -stack by one residue (Fig. 1c). Previous work *in vitro* and *in vivo* indicates that this tryptophan is important for both RNA binding and processive antitermination^{19,22}. We have previously used the RNA-binding domain of the λ N(1–22)-*boxB* complex as a model system to isolate high-affinity RNA-binding peptides by mRNA display^{23,24}. We were surprised when selections randomizing residues 13–22 of N showed no conservation of tryptophan at position 18 in

the selected peptides²³. Instead, a loose consensus containing Glu14 and Arg15 emerged, which also differed from the energetically important wild-type residues Lys14 and Gln15. Notably, selections in which residues 7, 14 and 15 were also randomized resulted in a sequence containing the Glu14Arg15 pair in the presence of Trp18. Here we address the structural, energetic and functional consequences of these mutations.

RESULTS

Context-dependent energetics

Our previous selections also hinted that certain residues in the RNA-binding domain of N might be energetically coupled²³. Subsequent sequence-covariation analysis of more than 60 different RNA-binding peptides isolated through *in vitro* selection experiments supported this idea, indicating a statistically significant linkage between (i) positions 14 and 15 and (ii) positions 15 and 18, and general linkage between (iii) adjacent residues and (iv) those positioned one helical turn away²⁵.

We were curious to examine the structural and functional consequences of these changes and assess why these selected sequences are not found in the wild-type phage. We constructed 20 peptides with systematic variations at positions 14, 15 and 18 and measured the binding of each one to *boxB* RNA to investigate any energetic coupling experimentally. In wild-type λ N(1–22; K14Q15), the residues at position 18 with highest binding affinities were tryptophan and tyrosine, whereas arginine, alanine and glutamate variants showed similar stability (Fig. 2). In other sequence contexts, both the best residues at position 18 and the rank order shifted substantially. For example, peptides containing Trp18, Ala18 and Arg18 all showed similar affinity when residues 14 and 15 were glutamate and arginine (that is, in the E14R15 variant), respectively. However, the peptide with Arg18 was

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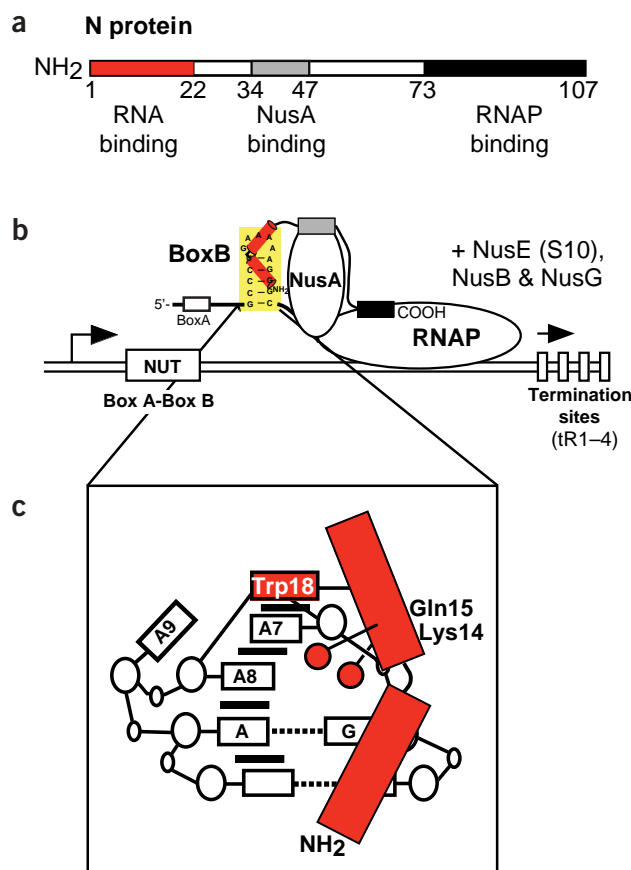


Figure 1 Structure and function of the N protein. (a) Domain structure of N. Regions for *boxB* RNA binding (residues 1–22), NusA binding (residues 34–47) and RNA polymerase binding (residues 73–107) are indicated³⁷. (b) Formation of the processive antitermination complex. The minimal antitermination complex⁴⁸ forms when N binds the *boxB* in the nascent mRNA and is stabilized by binding of NusA. Association of the other Nus proteins results in a processive antitermination complex. (c) Schematic of N peptide–*boxB* RNA complex indicating the peptide and loop stacking geometry.

peptide positions 14, 15 and 18 that were unique to each library. For the K14Q15-X18 and L14Q15-X18 libraries, we observed a strong preference for tryptophan and tyrosine at position 18. Fluorescence measurements of these variants showed quenching of 2-aminopurine (2AP)-7-substituted *boxB* similar to that of the wild-type peptide, consistent with π -stacking of the aromatic side chain on the RNA loop^{19,22} (Fig. 3b). Our selection results correlated well with previous genetic work demonstrating that the L14Q15 variant is highly functional and that tryptophan and tyrosine are the only functional substitutions at position 18 of N protein²². In contrast, our selections showed that aromatic and planar residues rarely occur at position 18 in the K14R15 and E14R15 variants. The selections therefore provided an experimental demonstration that the fitness landscape for *boxB* binding is rugged for the interfacial residues, and were suggestive of structural differences that would explain the observed sequence preferences.

An adaptive RNA-peptide interface

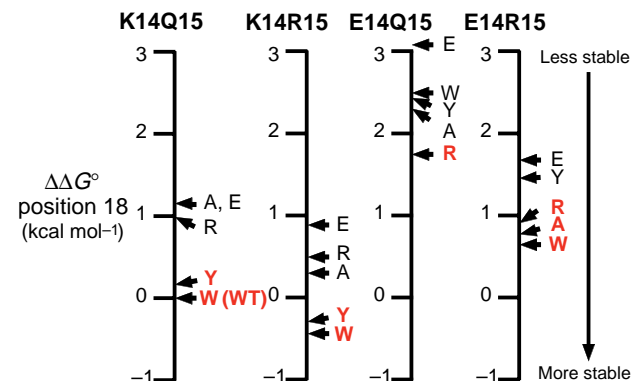
NMR provided a means to examine the structural origins of the context dependences we had observed. The indole NH proton of Trp18 and the RNA imino protons provide well-resolved spectroscopic handles in our RNA–peptide complexes. We reasoned that structural variations in the complexes should be revealed by intensity or chemical-shift changes of some of these resonances. All of the free peptides examined were unfolded in solution²³ and showed induced α -helix formation upon binding the *boxB* RNA (see Supplementary Fig. 1 online). The free *boxB* RNA showed three imino resonances; the rest were lost to exchange with solvent (Fig. 3a). In the wild-type λ N(1–22)–*boxB* complex (K14Q15, Fig. 3a), the RNA showed two more resonances from imino protons (five total), four corresponding to stem base pairs (U5, G14, G13 and G12) and one upfield at ~11 p.p.m. for the G6–A10 sheared pair. The most notable change in the peptide spectra occurred at Trp18, which shifted upfield by ~0.8 p.p.m. as a result of the ring-current effect of the adjacent, π -stacked RNA base (A7), consistent with previous structural work^{17–19}.

the best binder in a designed sequence containing E14Q15. Thus, the energetic landscape at position 18 depends on the sequence context one helical turn away at positions 14 and 15.

Probing sequence preferences with mRNA display

We next investigated the context-dependent energetics of RNA binding using *in vitro* selection experiments to rank the amino acids at position 18 in relation to different sequence contexts at positions 14 and 15. We inserted a saturation cassette (NNG/C codon) into the position corresponding to residue 18 of the protein. We then carried out mRNA display–based peptide selections²³ for five libraries with variants differing at positions 14 and 15: (i) K14Q15 (ii) L14Q15 (iii) K14H15 (iv) K14R15 and (v) E14R15. After two rounds of selection against *boxB* RNA^{23,24}, ~20 sequences were cloned from each library and assessed (Table 1). We observed good correlation between the number of times a peptide sequence occurred and its binding affinity for the *boxB* RNA, confirming our approach. The selected sequences showed strong context effects for *boxB* RNA binding and

Figure 2 Relative stability of N peptide–*boxB* RNA complexes that differ at positions 14, 15 and 18. The dissociation constant (K_d) for the wild-type (WT) complex was determined to be 1.2 nM for 2AP-7–labeled *boxB* RNA. Uncertainties in K_d values were <10%. For complexes with small fluorescence change at 2AP-7 (such as E14Q15), 2AP-8 and 2AP-9 *boxB* RNA were used. Complexes are ranked by their free energy change relative to the wild-type N–*boxB* complex ($\Delta\Delta G^\circ$; kcal mol⁻¹), with the most stable shown in red. Complexes are indicated by the peptide residue identity at positions 14 and 15 (top) and residue 18 (adjacent to arrows).



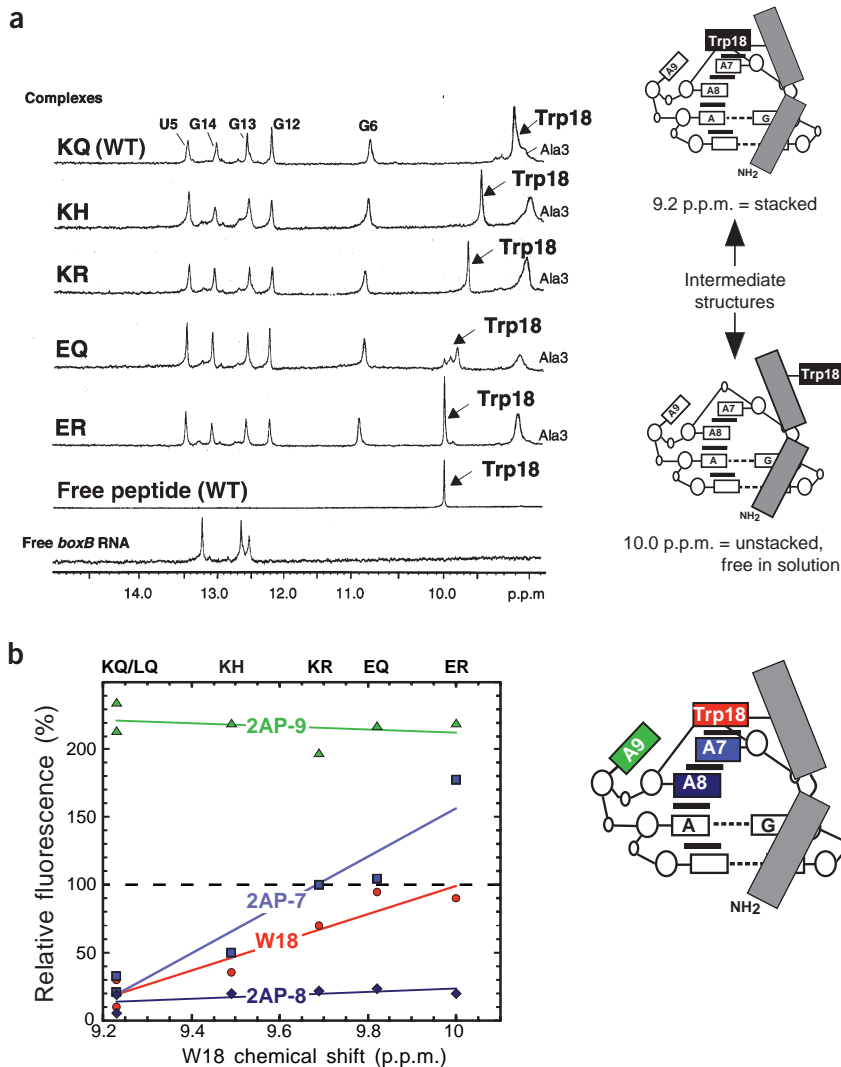


Figure 3 Structure and folding characteristics of *N*-*boxB* complexes as detected by NMR and fluorescence spectroscopy. **(a)** One-dimensional NMR spectra of peptide-RNA complexes containing Trp18 and residues K14Q14 (KQ; wild type, WT), K14H15 (KH), K14R15 (KR), E14Q15 (EQ) or E14R15 (ER) at positions 14 and 15. The positions of the imino protons and the tryptophan indole proton are indicated. One-dimensional spectra are also shown for the free peptide (wild type) and free *boxB* RNA. **(b)** Correlation of *N*-*boxB* fluorescence with Trp18 chemical shift. The relative fluorescence of Trp18 (unlabeled complex) or 2-aminopurine at RNA position 7, 8 or 9 (2AP-7, 2AP-8 and 2AP-9) is plotted versus the ^1H chemical shift of Trp18 (p.p.m.). The identities of peptide residues 14 and 15 are indicated at the top of the figure. Linear fits to the data and the stacking schematic of the wild-type complex are also shown.

cates increased stacking of the fluorophore, whereas enhancement indicates increased exposure to aqueous solvent. The stacked structure seen in the wild-type complex showed quenching of both position 7 (2AP-7) of the RNA and Trp18 of the peptide, whereas fluorescence enhancement for 2AP-7 was seen for the unstacked E14R15 variant. The relative fluorescence at 2AP-7 and for Trp18 showed linear correlations with each other and also with the Trp18 chemical shift we observed by NMR (Fig. 3b). RNA positions adjacent to this interface (2AP-8 and 2AP-9) showed a consistent amount of fluorescence quenching (22–28%) or enhancement (200–235%) irrespective of the Trp18 chemical shift for different complexes (Fig. 3b).

Taken together, the NMR, CD and fluorescence data indicate that the complexes have similarity in the RNA stem, the sheared GA pair, the flipped base (position A9) in the loop and the induced helical structure of the peptides. However, the critical Trp18–A7 interaction showed structural and energetic variability depending on the sequence context at positions 14 and 15, one helical turn away. Structurally, variable amounts of stacking between Trp18

All the complexes gave the same number and pattern of imino proton peaks at 10–14 p.p.m. (including the G6–A10 sheared pair imino), indicating that the mutant peptides induce similar RNA stem structure in all cases. Consistent with this idea, the N-terminal peptide (N(1–11)) present in all our mutants also induced the same RNA spectra with *boxB* (data not shown), indicating that stem recognition occurs predominantly with the N-terminal 11 residues. However, the Trp18 indole NH resonance in five of the RNA-peptide complexes showed considerable variability (Fig. 3a). Whereas both the wild-type complex and L14Q15 complex (data not shown) showed almost complete stacking on the loop as indicated by the large upfield Trp18 chemical shift, in the E14R15 complex, Trp18 appeared almost completely unstacked as indicated by the same chemical shift as in the free peptide. The other complexes (K14H15, K14R15, and E14Q15) showed intermediate levels of stacking, as gauged by their Trp18 chemical shifts between the wild type and E14R15.

The NMR observations correlated well with fluorescence experiments used to determine the K_d values for RNA-peptide complex formation. In these experiments, we measured the fluorescence ratio of the complex to free peptide or free 2AP-labeled RNA. Quenching indi-

Table 1 Amino acid composition at position 18 after two rounds of *in vitro* selection for binding *boxB* RNA

Library	Trp	Tyr	Phe	His	Total aromatic	Total other residues (%)
K14Q15-X18	19.2	19.2	7.7	3.8	50	50
L14Q15-X18	30.0	35.0	0	5.0	70	30
K14H15-X18	0	17.9	17.9	3.6	39.4	61.6
K14R15-X18	4.2	4.2	4.2	12.4	25	75
E14R15-X18	0	0	5.3	15.8	21.1	78.9

For each library, fractions (%) of sequences with tryptophan, tyrosine, phenylalanine and histidine at position 18 are listed separately, and their sum is listed as Total aromatic; the rest of the residues are listed as Total others. The numbers of sequences for each library are as follows: 26 for the K14Q15-X18 library, 20 for L14Q15-X18, 28 for K14H15-X18, 24 for K14R15-X18 and 19 for E14R15-X18.

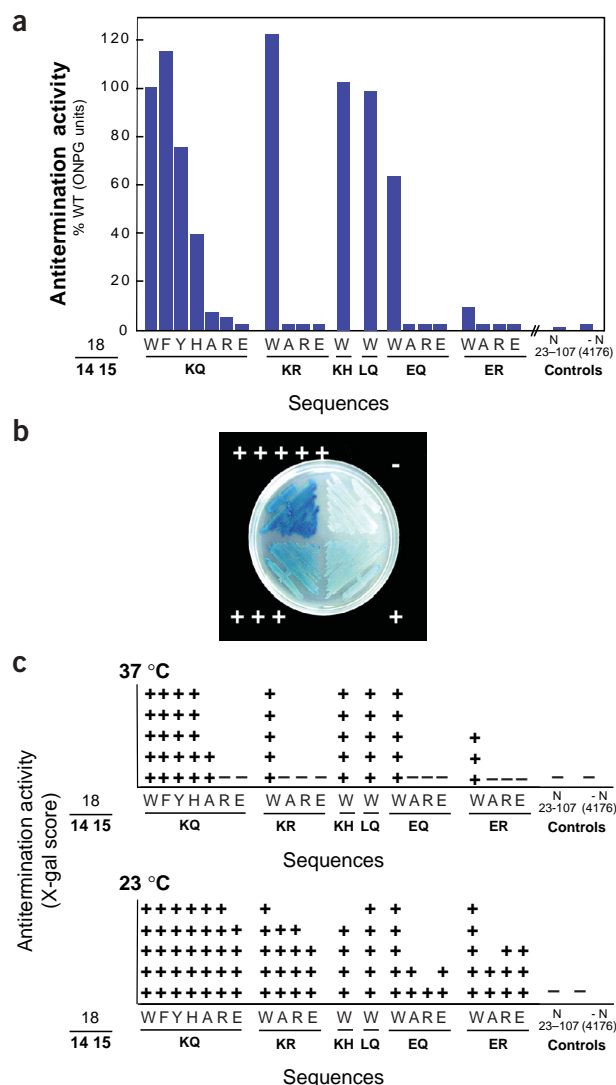


Figure 4 Antitermination activities of N mutants, as determined using *E. coli* strains carrying a two-plasmid reporter system²². Functional antitermination through four terminators (tR1–4) results in expression of β -galactosidase. (a) Antitermination activity in solution, determined by quantifying β -galactosidase using a colorimetric assay based on *o*-nitrophenyl- β -D-galactopyranoside. β -galactosidase activity is expressed as a percentage of that of the wild-type (WT) N reporter construct⁴⁹. The residue identities at positions 14, 15 and 18 are indicated. Control constructs in which N lacks an RNA-binding domain (N(23–107)) or the N expressor plasmid is absent (N 4176)²² are indicated. (b) Plate-scoring assay of mutants quantified on the basis of X-gal signal. (c) Plate-scoring assay of temperature sensitivity in the antitermination activity of N mutants grown at 37 °C or 23 °C.

Additionally, NMR and alanine scanning mutagenesis has also indicated that positions 3, 14 and 19 are important in recognition by phage N protein^{17–19,27}. Finally, one group of researchers had reported an *in vivo* screen based solely on approximation between a nascent transcript and N mutants containing arginine-rich RNA-binding domains at their amino terminus²⁸. However, this system alone generated only 2% as much readthrough as the wild-type N-*boxB* system, suggesting that the distinct conformation of the components has an important role in generating a fully functional antitermination switch. Furthermore, some RNA mutations that support N binding do not support the formation of higher-order complexes with the transcription elongation-termination factor NusA^{13,17}. The identities of RNA bases in the pentaloop of *boxB* affect N binding¹⁹ and antitermination²⁹ in somewhat different ways. G6 and A10 are absolutely required to maintain the GA sheared base pair, and purine bases are highly preferred for positions 7–9, with most pyrimidine substitutions resulting in <10% of wild-type antitermination activities except the A9U mutation (the flipped-out base), which has 35% of wild-type activity²⁹. Work in our laboratory had previously shown that wild-type N peptide discriminated between *boxB* RNA and a GAAA tetraloop with approximately ten-fold specificity²³.

These previous studies therefore motivated us to examine the *in vivo* activity of the N mutants we had studied. Our selected proteins are one-, two- and three-residue N mutants that preserve the known essential sequence elements that had been studied previously. Our measurements here of these mutants showed a wide range of stabilities (~3.5 kcal mol⁻¹) and marked structural differences at the Trp18–A7 peptide-RNA interface. We were interested to see if differences in the conformation or stability of these complexes would affect *in vivo* antitermination activity. We inserted these sequences into a β -galactosidase reporter construct and measured transcription antitermination efficiency²² in solution (Fig. 4a) and on agar plates (Fig. 4b,c). The activities of the complexes showed a full dynamic range, with some having activity equal to or better than that of the wild-type complex, whereas others show no activity, comparable to strains lacking N altogether (Fig. 4a). In the wild-type sequence context (K14Q15), tryptophan and other aromatic residues were necessary at position 18 for efficient antitermination, consistent with previous *in vivo* work²². However, assessment of the unstacked E14R15 mutant indicated that Trp18 alone is not sufficient for efficient antitermination, resulting in only about 9% the activity of the wild type. This loss of activity did not seem to be caused by lack of expression, as both the E14R15 mutant and wild-type N proteins were expressed *in vivo* as judged by northwestern analysis (data not shown). Thus, variations at positions 14 and 15 that tend to unstack Trp18 resulted in little or no antitermination activity. Finally, constructs lacking the RNA-binding domain of N(23–107) showed no

and A7 were observed, with the wild-type and L14Q15 complexes being fully stacked and the E14R15 complex completely unstacked. It is likely that different complexes have different population distributions of stacked and unstacked conformations in dynamic exchange²⁶. In terms of the energetics and selection, the wild-type and L14Q15 sequence contexts showed that Trp18 and Tyr18 are the clear winners, whereas the best residues in other sequence backgrounds varied somewhat, and in the E14R15 context, Arg18, Ala18 and Trp18 appeared energetically similar.

Stacking, not energetics, dictates *in vivo* function

Previous work from a number of groups has supported the idea that binding alone between N and *boxB* is sufficient to construct a functional antitermination switch. *In vitro* and *in vivo* studies have indicated that arginines 6, 7, 8, 10 and 11, as well as Trp18 or Tyr18, are critical for *in vivo* function as a result of their essential role in binding *boxB* RNA, whereas other positions such as Thr5, Glu9 and Glu13 can be readily substituted *in vitro* and *in vivo*^{19,22}. Previous *in vitro* selection work from our laboratory supported the idea that any substitutions for Arg6 and Arg7 disrupted stringent *boxB* RNA binding²³.

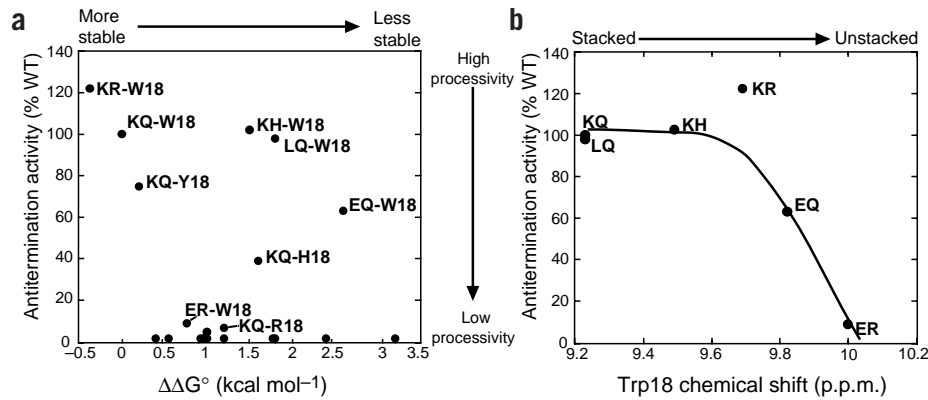


Figure 5 Energetic and structural correlation of antitermination activity. (a) No correlation is observed between the stability of the N-*boxB* complexes and their antitermination activity in solution. (b) Antitermination activity does correlate with Trp18-A7 stacking as detected by the chemical shift of Trp18 or the fluorescence of 2AP-7 (not shown).

activity in this assay, indicating that the signal does not result from *boxB*-independent antitermination by N (ref. 30).

Antitermination activity tested on agar plates expanded the lower dynamic range of the assay and revealed that a number of our variants have temperature-sensitive phenotypes (Fig. 4c). All variants that were positive at 37 °C also showed full activity at 23 °C. However, many variants that were inactive at 37 °C showed some level of antitermination activity at 23 °C. In particular, the fact that the E14R15 mutant has activity at lower temperature suggests that it binds *boxB* specifically *in vivo* and that its lack of functionality at higher temperature is not due to nonspecific RNA binding. Previous work has shown that some Nus proteins are dispensable for λ growth at lower temperatures (30 °C)³¹, consistent with our observations that many more of the complexes are functional at the permissive temperature.

Our data demonstrated that the structure of the N-*boxB* complex interface, rather than its stability, is the principal determinant of functional antitermination (Fig. 5). Our structural data suggested that the wild-type and mutant peptides have structural variation at the Trp18-A7 interface, while having essentially the same stem-helix interaction and overall loop folding (A9 extruded). The relative

stability of the complexes showed little correlation with antitermination activity (Fig. 5a). Six of the complexes we examined showed stability within 1 kcal/mol of that of the wild-type complex but <10% antitermination activity, whereas one (E14Q15-W18), which binds more than 2.5 kcal mol⁻¹ more weakly than the wild-type complex, had 63% activity *in vivo* (Fig. 5a). By contrast, complexes containing a stacked Trp18-A7 interface were fully active for antitermination (Fig. 5b). Complexes in which Trp18 is unstacked (E14R15, Fig. 5b) seemed to hinder or block processive antitermination. These data led us to conclude that there are different levels of control on the full dynamic range of transcription antitermination by N. The structure of the Trp18-A7 interface is critical in forming a fully functional antitermination switch, in addition to the effect of simple approximation of components²⁸.

Role of NusA

Our conclusion raised the questions of how the N-*boxB* conformation is sensed by the elongating polymerase, and how a single stacking interaction can dictate the processivity of a 400-kDa molecular machine. NusA is the most likely suspect as a conformational sensor.

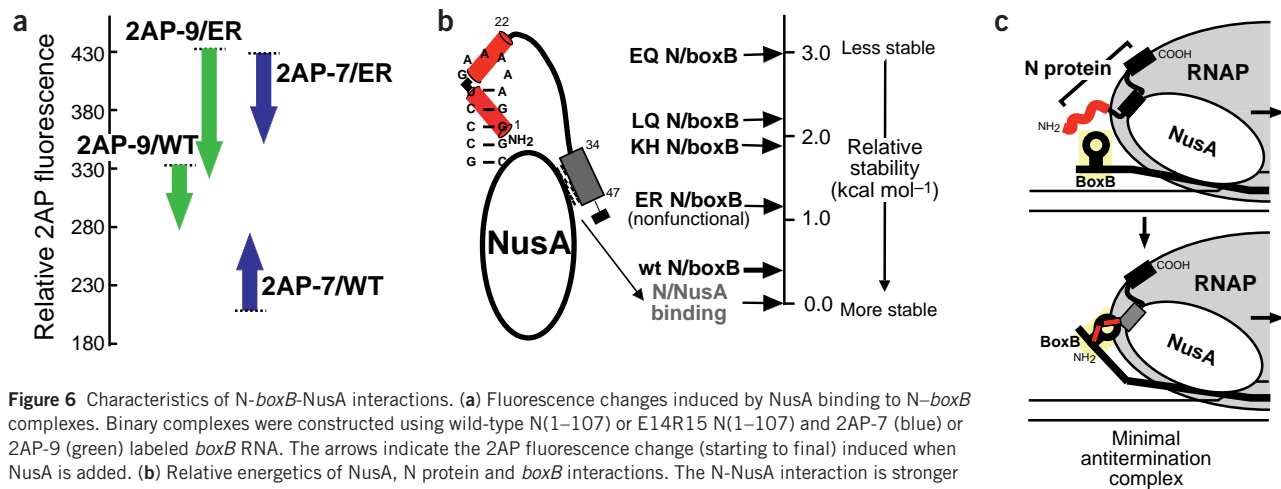


Figure 6 Characteristics of N-*boxB*-NusA interactions. (a) Fluorescence changes induced by NusA binding to N-*boxB* complexes. Binary complexes were constructed using wild-type N(1-107) or E14R15 N(1-107) and 2AP-7 (blue) or 2AP-9 (green) labeled *boxB* RNA. The arrows indicate the 2AP fluorescence change (starting to final) induced when NusA is added. (b) Relative energetics of NusA, N protein and *boxB* interactions. The N-NusA interaction is stronger than N binding to the *boxB* RNA. The wild-type (WT), K14H15, L14Q15 and E14Q15 complexes show >50% wild-type antitermination function *in vivo*, whereas ER shows <10% activity. (c) Model for formation of the minimal antitermination complex *in vivo*. The tight binding of N to NusA implies that N is prebound to polymerase during elongation and scans the message as it emerges from the transcription elongation complex. As the *boxB* hairpin emerges from the transcription bubble, the N-terminal domain of N engages the message to generate the minimal antitermination complex.

NusA has an important role in regulating the processivity of RNA polymerase^{32–35}. Previous work has indicated that NusA can discriminate between N–*boxB* complexes differing in a single nucleotide^{13,17} and can bind to N with high affinity at a site distinct from the RNA-binding domain¹². We therefore examined the binding and fluorescence effects of NusA on various 2AP-labeled N–*boxB* complexes (Fig. 6).

NusA showed no detectable interaction with complexes constructed from an N peptide (either wild-type or E14R15 N(1–22)) and a 2AP-labeled *boxB* (data not shown), indicating that the N–*boxB* interface alone may not be sufficient to recruit NusA binding with high affinity. NusA bound to complexes containing full-length N protein (wild-type or E14R15 N(1–107)) (Fig. 6a). In the wild-type N(1–107)–*boxB* complex, NusA quenched the fluorescence of the extruded base (2AP-9), but modestly enhanced the emission from 2AP at the crucial Trp18-A7 interface. In the E14R15 N(1–107)–*boxB* complex, NusA quenched both the extruded base (2AP-9) and the base located at the top of the loop (2AP-7). Thus, NusA seems to drive the binary N protein–*boxB* complexes toward similar, but distinct, final conformational states. Furthermore, NusA binds with similar, high-affinity K_d values to both wild-type and E14R15 complexes (14 ± 6 versus 9 ± 2 nM, respectively; see Supplementary Fig. 2 online), suggesting that NusA affinity is dictated by protein–protein contacts between N protein and NusA, rather than the N–*boxB* interface, in line with previous observations¹².

Notably, the K_d for NusA and N protein (70 nM) determined by van Gilst and von Hippel under more stringent temperature and salt conditions¹² actually indicates stronger binding than that of N protein to *boxB* RNA (K_d 127 nM under the same conditions; see Supplementary Fig. 3 online) (Fig. 6b). An energetic ranking of a number of functional and nonfunctional N–*boxB* interactions indicates that the N–NusA interaction is as much as 3 kcal mol^{–1} more favorable than the RNA–protein interactions needed to construct a functional antitermination switch *in vivo* (Fig. 6b).

DISCUSSION

A novel model for antitermination

In vivo, the N–*boxB* RNA complex is believed to serve the bipartite function of tethering NusA to the *nut* site RNA and sequestering N to interact directly with RNA polymerase^{36,37}. The relative energetics of N–NusA and N–*boxB* interactions lead us to propose a new model for assembly of the minimal antitermination complex (Fig. 6c). The very high affinity between N and NusA implies that N can be bound to the elongating polymerase, via interactions with NusA, where NusA associates with elongating polymerase with similar affinity (30–100 nM)^{32,33}. This complex can be assembled stably before the exit of the *nut* site from polymerase. This view differs from the model generally described for antitermination, whereby N is recruited to the transcription elongation complex (TEC) via interactions with *boxB* (Fig. 1b)^{13,36,38,39}. Previous structural modeling of TECs supports the idea that polymerase constantly scans the nascent transcript for termination signals via the flap-tip helix and the action of bound NusA⁴⁰. Our work implies that N protein modifies NusA, altering the function of RNA polymerase through NusA and RNAP interactions when it properly engages an appropriate RNA structure. This view of N action is consistent with experiments showing that N supports efficient transcription antitermination even in the absence of *boxB*^{16,35} and with modeling efforts focused on understanding transcription termination⁴⁰. We conclude that conformational sensing of the Trp18-A7 interface occurs in the context of a TEC, containing N, NusA and RNA polymerase itself, after N has engaged the nascent *boxB* hairpin. Essentially, the unfolded N acts as a template onto which the compo-

nents of the antitermination complex assemble. This action of N is apparently dependent on the exact conformation of the interface of Trp18-A7, where the presence or absence of the stacking directs NusA and polymerase into either different orientations and spatial arrangements or different physical contacts. Supporting this view is previous work indicating that the quaternary NusA–N–*nut* polymerase complex can be formed and is stable with respect to competition from exogenous components³⁵. Although these components represent a minimal antitermination complex, assembly of the complete complex (containing N, NusA, NusB, NusE, NusG, *boxA*, *boxB* and RNAP) is essential for processive antitermination under stringent conditions *in vivo*¹⁴.

Implications

Our results obtained under stringent conditions suggest that proper assembly of the RNA–protein interface is indispensable for full antitermination activity *in vivo*. One consequence of our observations is the conclusion that genetic screens using antitermination to isolate RNA-binding peptides²⁸ will likely be influenced by constraints of functional antitermination.

The N–*boxB* interface provides a clear example of adaptive recognition in which the sequence context dictates the structural meaning of interfacial residues. *In vivo*, the various conformations we observed had profound effects on polymerase function and demonstrated that approximation of components is not sufficient to generate a functional transcriptional switch. Rather, the transcription elongation complex has the means to sense the conformation of *cis*-acting regulatory elements, much like a stockbroker reading a ticker tape, profoundly altering its processivity accordingly. Similarly, it seems likely that a more complicated eukaryotic elongation complex could distinguish between various *cis*-acting regulatory signals such as those provided by HIV Tat^{41,42} and Nrd1p^{43–45}. In addition to revealing the structural origin of this control process, our work indicates that ligands targeting elements of elongation complexes⁴⁶ offer potential tools to better understand and control the mechanisms of gene expression in both prokaryotic and eukaryotic systems.

METHODS

***In vitro* peptide selection.** Generation and purification of mRNA–peptide fusions and selection cycles were done as previously described²³. Approximately 5 pmol of purified ³⁵S-labeled fusions were incubated with 200 pmol of biotinylated *boxB* RNA (5′-GGCCUGAAAAGGGCCAAA-biotin-3′) in the selection step in the presence of tRNA competitor.

Synthesis of RNAs and peptides. The 2-aminopurine (2AP)-labeled *boxB* RNAs, 5′-GGCCUGAAAAGGGCC-3′ (with underlined nucleotides individually labeled 2AP-7, 2AP-8, 2AP-9), were generated by solid-phase synthesis using standard phosphoramidite chemistry, and then deprotected and purified by 20% urea-PAGE. RNA without the 2AP label for NMR experiments (5′-GCCUGAAAAGGGCC-3′) was synthesized using T7 RNA polymerase and a synthetic DNA template⁴⁷. Peptides MDAQTRERRAEXXAQXKAAN(gy) (where X represents variable residues and gy represents extra residues for quantification purposes for peptides lacking Trp18) were made by automated Fmoc synthesis (using an ABI 432), deprotected and purified by reversed-phase HPLC.

Binding studies. K_d values for N-peptide binding to *boxB* RNA were determined by monitoring the fluorescence change of RNAs labeled with 2AP at the seventh, eighth or ninth position upon addition of concentrated peptide and were fitted as previously described²³. NusA binding was performed by titrating NusA solution into preformed N–*boxB* complex. Free energies were calculated on the basis of complex stability at 20 °C in standard binding buffer (50 mM potassium acetate, 20 mM Tris, pH 7.5). Individual K_d values were reproducible with an error of ± 0.2 kcal mol^{–1}.

NMR spectroscopy. Spectra were collected on a Varian Inova 600 at 25 °C in a buffer of 50 mM NaCl, 10 mM sodium phosphate, 0.5 mM EDTA, pH 6. Complex formation was monitored by inspecting imino protons of the RNA. The concentrations of the complexes were 150–300 μM.

Circular dichroism. CD spectra were collected on an Aviv 62 DS CD spectrofluorimeter in 10 mM potassium phosphate buffer at 20 °C. Spectra for the peptide portion in the complex were calculated by subtracting spectra for free RNA and excess peptide from that of the complex. Helical content was calculated as described previously⁴⁶.

Antitermination assay. Strains expressing wild-type and mutant N protein were constructed using the two plasmid reporter system previously described²². All sequences were verified by sequencing. N-mutant strains were plated on tryptone agar supplemented with 0.05 mM IPTG, 0.08 mg ml⁻¹ X-gal and the appropriate antibiotics. Plates were incubated at either 23 °C (permissive temperature) or 37 °C (nonpermissive temperature) and scored for blue color. Temperature sensitivity was defined here as the loss of relative β-galactosidase units in comparing the blue color of colonies grown at permissive and nonpermissive temperatures. All assays were performed in triplicate. Intensities of blue color are shown for sample N-mutant strains K14Q15E18 (-), K14Q15A18 (+), Nun-N fusion (+++) and K14Q15W18 (++++).

Protein expression and purification. Wild-type and E14R15 N protein were expressed from plasmid pET-N1 in BL21 (DE3) gold cells as described¹², except that an SP-cation exchange column (Amersham) was used for the final purification step and the protein was dialyzed into fluorescence binding buffer (50 mM KCl, 20 mM Tris-HCl, pH 7.5, 1 mM β-mercaptoethanol). The NusA gene was amplified from plasmid pMS7, subcloned into vector pET19b (Novagen) and verified by sequencing. BL21 (DE3) cells harboring the resulting plasmid were induced at an OD₆₀₀ of 0.4 with 1 mM IPTG, grown for 3 h, harvested by centrifugation and resuspended 20 ml buffer A (50 mM Tris-HCl pH 8.0, 100 mM NaCl, 1 mM β-mercaptoethanol, 1 mM imidazole) per liter of culture. The cells were lysed by three passes through a French press and clarified by two centrifugations at 18,000g for 30 min at 4 °C. The supernatant was loaded on a Superflow Ni-NTA column (Qiagen), washed with 5 column volumes of buffer A and then with 5 column volumes of buffer A containing 500 mM NaCl, and eluted with 400 mM imidazole in buffer A. The eluted protein was dialyzed against fluorescence binding buffer.

N protein expression level. Expression of wild-type N protein, E14R15 and N(23–107) variants was accomplished as above except that the cells were induced with 0.5 mM IPTG at an OD₆₀₀ of 0.3 and grown for 3 h, and 0.1 OD units of culture was concentrated and resuspended in 100 μl of 6× SDS load buffer (0.28 M Tris-Cl/SDS, pH 6.8, 30% (v/v) glycerol, 1% (w/v) SDS, 0.5 M DTT). Five microliters of this solution was separated by 15% SDS-PAGE and then transferred to nitrocellulose (Bio-Rad) in transfer buffer (10 mM NaHCO₃ –3 mM Na₂CO₃, pH 9.9, 20% (v/v) methanol, 0.05% (w/v) SDS) at 4 °C. The nitrocellulose was blocked at room temperature in wash buffer (10 mM Tris-HCl, pH 7.5; 100 mM NaCl; 1 mM EDTA, pH 8.0; 0.01% (v/v) NP-40) containing 5% (w/v) nonfat dry milk and 50 μg per milliliter of yeast tRNA, and then 100 nM biotinylated *boxB* RNA (5'-GGCCUGAAAAGGCCAAA-biotin-3') was added. The nitrocellulose was washed three times with wash buffer, probed with 0.5 μg ml⁻¹ streptavidin-horseradish peroxidase (Amersham) in wash buffer containing 0.5% (w/v) BSA, washed four times and visualized using Supersignal West Pico substrate (Pierce).

Note: Supplementary information is available on the Nature Structural Biology website.

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COMPETING INTERESTS STATEMENT

The authors declare that they have no competing financial interests.

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