# Structural basis of transcription activation by Rob, a pleiotropic AraC/XyIS family regulator

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# ABSTRACT

Rob, which serves as a paradigm of the large AraC/XvIS family transcription activators, regulates diverse subsets of genes involved in multidrug resistance and stress response. However, the underlying mechanism of how it engages bacterial RNA polymerase and promoter DNA to finely respond to environmental stimuli is still elusive. Here, we present two cryo-EM structures of Rob-dependent transcription activation complex (Rob-TAC) comprising of Escherichia coli RNA polymerase (RNAP), Robregulated promoter and Rob in alternative conformations. The structures show that a single Rob engages RNAP by interacting with RNAP  $\alpha$ CTD and  $\sigma^{70}$ R4, revealing their generally important regulatory roles. Notably, by occluding  $\sigma^{70}$ R4 from binding to -35 element, Rob specifically binds to the conserved Rob binding box through its consensus HTH motifs, and retains DNA bending by aid of the accessory acidic loop. More strikingly, our ligand docking and biochemical analysis demonstrate that the large Rob C-terminal domain (Rob CTD) shares great structural similarity with the global Gyrl-like domains in effector binding and allosteric regulation, and coordinately promotes formation of competent Rob-TAC. Altogether, our structural and biochemical data highlight the detailed molecular mechanism of Robdependent transcription activation, and provide favorable evidences for understanding the physiological roles of the other AraC/XylS-family transcription factors.

# INTRODUCTION

In order to survive, bacteria have successfully evolved complex transcription initiation mechanisms to timely and finely evoke adaptive responses to a myriad of environmental threats, through transcribing key genes involved in stress responses by RNA polymerase (1-3). In Escherichia coli, RNA polymerase (RNAP) is assembled by a multisubunit RNA polymerase core  $(\alpha_2 \beta \beta' \omega)$  and a principal promoterspecific factor  $\sigma^{70}$  (alternative  $\sigma$  factors will be required under stress conditions) (4). At transcription initiation stage, in combination with RNA polymerase core,  $\sigma^{70}$  usually makes specific interactions with consensus -35 element (TTGACA) and -10 element (TATAAT) of canonical promoters by  $\sigma^{70}$  region 4 ( $\sigma^{70}$ R4) and  $\sigma^{70}$  region 2 ( $\sigma^{70}$ R2), respectively (5–7). Subsequently, the double strands of promoter DNA melt, allowing the RNAP-promoter closed complex (RPc) isomerizes into a stable RNAP-promoter

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open complex (RPo) which enables transcription initiation without activators (8–12). In contrast, most promoters of bacterial stress response genes do not simultaneously contain the two canonical consensus promoter elements. Thus, transcription activators are required to strengthen the interactions between RNAP and such non-canonical promoters. These interactions remodel promoter DNA, sustain optimal promoter recognition, and finally promote RPc to isomerize into an initiation-competent transcription activation complex (TAC), which is virtually an activator-dependent RPo (13–18).

Rob, originally designated by its binding ability to the right border of the *E. coli* replication origin *oriC* (19), is characterized as a representative member of the global AraC/XylS family transcription activators, which regulate expression of diverse subsets of genes involved in multidrug resistance, virulence, and stress response in many clinically important pathogens, such as Vibrio cholerae, Salmonella typhimurium, Yersinia pestis, Pseudomonas putida, Citrobacter rodentium, enteroaggregative E. coli and Shigella spp. (20–23). The common feature of AraC/XylS family transcription activators is the composition of two conserved helix-turn-helix (HTH) motifs that interact with the upstream recognition element A (designated as A site) and the downstream recognition element B (designated as B site) of the highly degenerate corresponding promoters, respectively (20,23). Up to date, a growing number of genetic and biochemical studies have elucidated that Rob is a pleiotropic transcription activator targeting widely distributed Rob regulons. Overexpression of Rob not only causes antibiotics resistance through down-regulating the influx activity, but also leads to adaptive tolerance to organic solvents, superoxide-generating agents and heavy metals, mainly by upregulating the efflux activity to afford increased cellular detoxification (2, 24-27).

Though the HTH motif containing Rob N-terminal domain (Rob NTD) shares high sequence and structural similarities with those of the other AraC/XylS family regulators, various studies have proved that Rob also has its own distinct functional and structural characteristics: (i) Unlike SoxS and MarA, which are regulated by certain inducers, Rob is constitutively expressed and maintained at a high level in cell, as can be regulated by itself and other paralogs (19,28). (ii) Rob has a large Cterminal domain (Rob CTD), which was identified as a potential effector binding domain (25,26). Rob CTD also plays physiological roles in the activation of Rob by an effector through a 'sequestration-dispersal' mechanism, and in retaining stabilization by protecting Rob from cellular protease degradation by Lon and ClpYQ (29). However, the underlying mechanism of how Rob orchestrates its CTD and NTD to synergistically respond to changing environmental stimuli is still poorly understood. (iii) Comparative analysis of the co-crystal structures of Rob-micF double-stranded DNA (PDB ID: 1D5Y) (30) with MarAmar double-stranded DNA (PDB ID: 1BL0) (31) shows that Rob displays two major differences in DNA binding from MarA. The second HTH (helix 6) of Rob only binds to the DNA backbone instead of binding to the adjacent DNA major groove as MarA does, and thus orientation of the upstream DNA is unbent while MarA bends DNA with an angle of  $35^{\circ}$ . Whether this conformation of RobmicF structure is a functional state is still under dispute. Although previous genetic epistasis experiments and mutational analysis have identified that some key residues of Rob were involved in DNA binding and transcription activation (29,32), the detailed mechanism mediating Robdependent transcription activation is still elusive, largely due to the absence of accurate structural information for Rob-dependent transcription activation complex. Furthermore, whether Rob interacts with the C-terminal domain of RNAP alpha subunit ( $\alpha$ CTD) and activates transcription of Rob regulons through proposed 'pre-recruitment' mechanism for the other AraC/XylS-family transcription factors remain to be elucidated.

In the present study, we determined two functional cryo-EM structures of Rob-dependent transcription activation complex (Rob-TAC<sup>I</sup> and Rob-TAC<sup>II</sup>). The structures trap two alternative conformations of Rob in complex with E. coli RNAP and micF promoter DNA, whose downstream gene is involved in cellular influx of antibiotics, detergents, and toxins. In Rob-TAC, a single Rob molecule engages RNAP by simultaneously interacting with RNAP aCTD and  $\sigma^{70}$ R4, revealing their versatility and important regulatory roles. Notably, by occluding  $\sigma^{70}$ R4 from binding to promoter -35 element, Rob specifically binds to the conserved Rob binding box through its consensus HTH motifs, and retains DNA bending by aid of the accessory acidic loop (residues 187-193) from Rob CTD. More strikingly, our ligand docking and biochemical analysis demonstrate that the large Rob CTD shares great structural similarity with the global Gyrl-like domains in effector binding and allosteric regulation, and it coordinately promotes formation of competent Rob-TAC along with the DNA-binding Rob NTD. Our results systematically define the key interactions that mediate Rob-dependent transcription activation, provide a new mechanistic framework for bacterial transcription regulation, and may support further exploration on the physiological roles of the other AraC/XylS family transcription factors.

#### MATERIALS AND METHODS

#### **Plasmid construction**

Plasmid pET28a-rob encoding C-terminal His6 tagged E. coli Rob under the control of T7 promoter was synthesized by Sangon Biotech, Inc. Linear micF\_mango DNA fragment corresponding to -85 to +50 of E. coli micF promoter followed by Mango III coding sequence was generated by de novo PCR using primers of micF\_mango F1, R1 and R2 (Supplementary Table S1) (16,33-36), purified using the QIAquick PCR Purification Kit (Qiagen, Inc.), and stored at -80°C. Two strands of micF DNA fragment corresponding to -63 to +20 of micF promoter were synthesized and annealed in a mole ratio of 1:1. Plasmids carrying rob amino acid substitutions were constructed using sitedirected mutagenesis (QuikChange Site-Directed Mutagenesis Kit, Agilent, Inc.). Primers of Rob mutants used in this study are shown in Supplementary Table S1. Sequences of micF\_mango DNA and micF DNA are shown in Supplementary Figure S1.

# **Rob protein purification**

E. coli strain BL21(DE3) (Invitrogen, Inc.) was transformed with plasmid pET28a-rob or pET28a-rob derivatives. Single colonies of the resulting transformants were used to inoculate 11 LB broth containing 50 µg/ml Active Ingredient, and cultures were incubated at  $37^{\circ}$ C with shaking until OD<sub>600</sub> reached 0.6. Protein expression was induced by addition of IPTG to 0.5 mM, and cultures were incubated 14 h at 20°C. The cells were harvested by centrifugation (5422 g; 10 min at 4°C), resuspended in 20 ml buffer A (20 mM Tris-HCl, pH 8.0, 0.2 M NaCl, 5% glycerol), and lysed using a highpressure homogenizer NanoGenizer (Genizer LLC, Irvine, CA, USA). The lysate was centrifuged (13000 g; 40 min at 4°C), and the supernatant was loaded onto a 3 ml Ni-NTA agarose column (Oiagen, Inc.) equilibrated with buffer A. The column was subsequently washed with 30 ml buffer A containing 0.04 M imidazole and eluted with 15 ml buffer A containing 0.2 M imidazole. The eluate was loaded onto a HiLoad 16/600 Superdex 200 column (GE Healthcare, Inc.) equilibrated in buffer B (20 mM Tris-HCl, pH 8.0, 75 mM NaCl, 5 mM MgCl<sub>2</sub>), and the column was eluted with 120 ml of the same buffer. Fractions containing E. coli Rob were pooled and stored at -80°C. E. coli Rob deriva-tives were expressed and purified using the same procedure as the wild type protein. The yields were 4 mg/l, and the purities were > 95%.

## E. coli RNAP purification

E. coli RNAP was prepared from E. coli strain BL21(DE3) (Invitrogen, Inc.) transformed with plasmids of pGEMD (37) and pIA900 (38). Single colonies of the resulting transformants were used to inoculate 50 ml LB broth containing 100 µg/ml Active Ingredient, and cultures were incubated 16 h at 37°C with shaking. Aliquots (10 ml) were used to inoculate 1 L LB broth containing 100 µg/ml Active Ingredient, cultures were incubated at 37°C with shaking until  $OD_{600} = 0.6$ , cultures were induced by addition of IPTG to 1 mM, and cultures were incubated 15 h at 20°C. Then cells were harvested by centrifugation (5000 g; 15 min at 4°C), resuspended in 20 ml lysis buffer C (50 mM Tris-HCl, pH 7.9, 0.2 M NaCl, 2 mM EDTA, 5% glycerol and 5 mM DTT) and lysed using a high-pressure homogenizer NanoGenizer (Genizer LLC, Irvine, CA, USA). The lysate was cleared by centrifugation (13 000 g; 30 min at  $4^{\circ}$ C), and the supernatant was precipi-tated by 0.7% (g/ml) Polymin P (pH 7.9), incubated for 10 min at 4°C with stirring, followed by centrifugation (8000 g; 15 min at 4°C). The precipitate was washed with buffer D (20 mM Tris-HCl pH 7.9 and 5% glycerol) containing 0.5 M NaCl, followed by centrifugation (8000 g; 15 min at 4°C). Protein was extracted with buffer D containing 1 M NaCl, followed by centrifugation (8000 g; 20 min at 4°C). Then the extract is precipitated by addition of 30 g ammonium sul-fate, incubated for 30 min at 4°C with stirring, followed by centrifugation (13 000 g; 30 min at 4°C). Subsequently, the pellet was resuspended in buffer D containing 0.5 M NaCl, and loaded onto a 5 ml column of Ni-NTA agarose (Qiagen, Inc.) pre-equilibrated with the same buffer. The column was washed with 25 ml buffer D containing 20 mM imidazole and eluted with 25 ml buffer D containing 0.15 M imida-zole. The eluate was diluted in buffer E (20 mM Tris-HCl,

pH 7.9, 5% glycerol, 1 mM EDTA and 1 mM DTT) and loaded onto a Mono Q 10/100 GL column (GE Healthcare, Inc.) equilibrated in buffer E and eluted with a 160 ml linear gradient of 0.3-0.5 M NaCl in buffer E. Fractions containing *E. coli* RNAP were pooled and applied to a HiLoad 16/600 Superdex 200 column (GE Healthcare, Inc.) equilibrated in buffer B, and the column was eluted with 120 ml of the same buffer. Fractions containing *E. coli* RNAP were pooled and stored at -80°C. Yield was 2.5 mg/l, and purity was >95%.

# Assembly of E. coli Rob-TAC

DNA oligonucleotides *micF* scaffold (sequence is shown in Figure 1A) were synthesized (Sangon Biotech, Inc.) and dissolved in nuclease-free water to 1 mM. Template strand DNA and nontemplate strand DNA were annealed at 1:1 ratio in 10 mM Tris–HCl, pH 7.9, 0.2 M NaCl. Then, *E. coli* Rob-TAC was assembled by incubating *E. coli* RNAP, *micF* scaffold, and *E. coli* Rob in a molar ratio of 1: 1.1: 10 at 4°C overnight. Subsequently, the mixture was applied to a HiLoad 16/600 Superdex 200 pg column (GE Healthcare Life Sciences) equilibrated in buffer B. After identification by SDS-PAGE and electrophoretic mobility shift assay (EMSA), the fractions containing *E. coli* Rob-TAC were concentrated to 18.7 mg/ml using Amicon Ultra centrifugal filters (10 kDa MWCO, Merck Millipore, Inc.).

# **Cryo-EM grid preparation**

Immediately before freezing, 6 mM CHAPSO was added to the freshly purified *E. coli* Rob-TAC. C-flat grids (CF-1.2/1.3-4C; Protochips, Inc.) were glow-discharged for 60 s at 15 mA (PELCO/Easiglow apparatus) or with a cleaning time for 120 s (Gatan/SOLARUS 950 plasma cleaning system) prior to the application of 3  $\mu$ l of *E. coli* Rob-TAC complex, then plunge-frozen in liquid ethane using a Vitrobot (FEI, Inc.) with 95% chamber humidity at 10°C.

# Cryo-EM data acquisition and processing

The grids with E. coli Rob-TAC were imaged using a 300 kV Titan Krios (FEI, Inc.). Images were recorded with Serial EM(39) in counting mode with a physical pixel size of 1.100 Å or 1.307 Å and a defocus range of 1.4–2.2  $\mu$ m. Data were collected with a dose of 10 e/pixel/s. Images were recorded with a 10 s exposure and 0.25 s subframes. Subframes were aligned and summed using MotionCor2 (40). The contrast transfer function was estimated for each summed image using CTFFIND4 (41). From the summed images,  $\sim 1000$ particles were manually picked and subjected to 2D classification in RELION (42). 2D averages of the best classes were used as templates for auto-picking in RELION. Autopicked particles were manually inspected, then subjected to 2D classification in RELION. Poorly populated classes were removed, resulting in a dataset of 791 759 particles from 4,372 movies for *E. coli* Rob-TAC<sup>I</sup> or a dataset of 417 732 particles from 2099 movies for E. coli Rob-TAC<sup>II</sup>. These particles were 3D-classified in RELION using a map of E. coli RPo (PDB ID: 6CA0) (10) low-pass filtered to 40 Å resolution as a reference. For *E. coli* Rob-TAC<sup>I</sup>, 3D classification resulted in 4 classes. Particles in Classes 2 and 3

were combined and further performed 3D classification focused in the region of Rob. After focused 3D classification, particles in Classes 2, 3, 4 were combined and processed by CTF refinement, Bayesian polishing and 3D auto-refined, and the best-resolved 386,188 particles was post-processed in RELION. The Gold-standard Fourier-shell-correlation analysis indicated a mean map resolution of 4.06 Å of *E. coli* Rob-TAC<sup>1</sup> (Supplementary Figure S3 and S5). Likewise, For *E. coli* Rob-TAC<sup>II</sup>, 3D classification resulted in 3 classes. Particles in Class 3 were processed by CTF refinement, Bayesian polishing, 3D auto-refinement, the bestresolved 335,021 particles were post-processed in RELION. And the Gold-standard Fourier-shell-correlation analysis indicated a mean map resolution of 4.57 Å of *E. coli* Rob-TAC<sup>II</sup> (Supplementary Figure S4 and S6).

#### Cryo-EM model building and refinement

The model of *E. coli* RNAP RPo (PDB ID: 6CA0) (10), the crystal structure of Rob in complex with *E. coli micF* DNA (PDB ID: 1D5Y) (30), and the ternary structure of *E. coli* MarA, DNA and RNAP  $\alpha$ CTD (PDB ID: 1XS9) (43) were manually fitted into the cryo-EM density maps in Coot (44), followed by adjustment of main- and sidechain conformations in Coot and real-space refinement using Phenix (45). The coordinates were real-space refined with secondary structure restraints in Phenix.

#### In vitro transcription assay

In vitro Mango-based transcription assays were carried out by incubating E. coli RNAP, micF-mango scaffold, with or without E. coli Rob or its derivatives. Transcription assay was performed in a 96-well microplate format. Reaction mixtures (80 µl) contained: 0 or 4 µM Rob or Rob derivatives, 0.1 µM E. coli RNAP, 50 nM micF-mango scaffold, 1 µM TO1-Biotin, 0.1 mM NTP mix (ATP, UTP, GTP and CTP), with or without ligand (100  $\mu$ M chenodeoxycholic acid or 5 mM 2, 2'-bipyridine), 40 mM Tris-HCl, pH 8.0, 50 mM NaCl, 10 mM MgCl<sub>2</sub>, 5% glycerol. First, E. coli RNAP, Rob and micF-mango scaffold were incubated for 15 min at 37°C, then mixtures were supplemented with 0.030 mg/ml Active Ingredient and incubated for 2 min at 22°C, then NTP mix and TO1-biotin were added into the mixture and incubated for 10 min at 37°C. Finally, fluorescence emission intensities were measured using a multimode plate reader (EnVision, PerkinElmer Inc.; excitation wavelength = 510nm; emission wavelength = 535 nm). Relative transcription activities of Rob derivatives were calculated using:

$$A = (I - I_0) / (I_{WT} - I_0)$$
(1)

where  $I_{WT}$  and I are the fluorescence intensities in the presence of Rob and Rob derivatives;  $I_0$  is the fluorescence intensity in the absence of Rob.

#### Electrophoretic mobility shift assay

To further clarify whether Rob derivatives affect the formation of Rob-TAC or not, we also carried out electrophoretic mobility shift assays in reaction mixtures (20  $\mu$ l) containing:1  $\mu$ M Rob or Rob derivatives, 50 nM *E. coli* RNAP, 15 nM micF DNA in EMSA buffer (40 mM Tris-HCl, pH 8.0, 100 mM NaCl, 10 mM MgCl<sub>2</sub>, 5% glycerol). E. coli RNAP was firstly incubated with micF DNA for 10 min at 37°C, then incubated with Rob or Rob derivatives for 20 min at 37°C. Reaction mixtures were supplemented with 0.050 mg/ml Active Ingredient and incubated for 2 min at 22°C. The reaction mixtures were finally applied to 5% polyacrylamide slab gels (29:1 acrylamide/bisacrylamide), electrophoresed in 90 mM Tris-borate, pH 8.0, and 0.2 mM EDTA, and stained with 4S Red Plus Nucleic Acid Stain (Sangon Biotech, Inc.) according to the procedure of the ana-lyzed by ImageJ manufacturer, and (https:// imagej.nih.gov/ij/).

Electrophoretic mobility shift assays (EMSA) of *E. coli* Rob and RNAP were performed in reaction mixtures (20  $\mu$ l) containing: 1  $\mu$ M Rob or Rob derivatives, 2  $\mu$ M *E. coli* RNAP, 15 nM *micF* DNA in EMSA buffer. Rob was incubated with *E. coli* RNAP for 10 min at 25°C and then incubated with (or without) DNA scaffold for 15 min at 25°C. Reaction mixtures were supplemented with 0.050 mg/ml Active Ingredient and incubated 2 min at 22°C. The reaction mixtures were applied to 5% polyacrylamide slab gels (29:1 acrylamide/bisacrylamide), electrophoresed in 90 mM Tris-borate, pH 8.0, and 0.2 mM EDTA, and stained with 4S Red Plus Nucleic Acid Stain (Sangon Biotech, Inc.) or coomassie brilliant blue staining solution (Sangon Biotech, Inc.) according to the procedure of the manufacturer.

# Molecular Docking analysis and the burying surface area measurement

All molecular docking studies were performed using Autodock4.2 package (46). Briefly, Rob from the cryo-EM structure of Rob-TAC<sup>II</sup> was docked with its potential ligands (chenodeoxycholic acid, 2,2'-bipyridine, sodium decanoate and 4,4'-bipyridine). The molecule was added with non-polar hydrogens and assigned partial atomic charges using AutoDockTools (ADT) (46). The coordinates of potential ligands in Rob structure were generated based on the coordinates of rhodamine 6G (R6G) from the crystal structure of the Gyrl-like family protein SAV2435 (PDB ID: 5KAW) (53) in combination with CORINA Classic online service. A grid box with  $40 \times 40 \times 40$  grid points and 0.2 grid spacing centered roughly at the potential ligand binding position was used as the searching space. 100 runs of Larmarckian Genetic Algorithm were performed to search the protein-ligand interactions. The results were clustered and ranked. Result analyses and figure rendering were per-formed using PyMOL.

To measure the potential burying surface areas in the Rob-TAC<sup>I</sup> or Rob-TAC<sup>II</sup>, we used the online web server (http://sts.bioe.uic.edu/castp) and followed the descriptions as illustrated before (54).

#### Data analysis

Data for *in vitro* transcription assays or EMSA are means of three technical replicates. Error bars represent mean  $\pm$ SEM of n = 3 experiments. Asterisk (\*\*\*) or (\*\*) indicates highly significant (*P* value < 0.001) or significant (*P* value < 0.01) difference from the wild-type Rob analyzed by oneway ANOVA with Tukey's multiple comparison test, respectively.

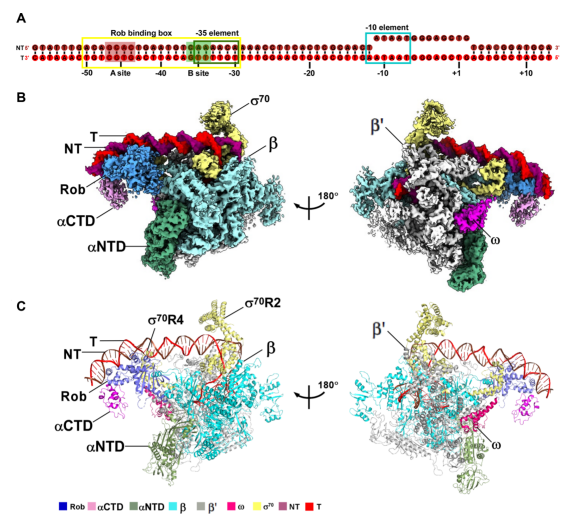


Figure 1. The overall structure of *E. coli* Rob-TAC. (A) DNA scaffold used in structure determination of *E. coli* Rob-TAC<sup>II</sup>. (B, C) Two views of the cryo-EM density map (B) and structure model (C) of *E. coli* Rob-TAC<sup>II</sup>. The EM density maps and cartoon representations of Rob-TAC<sup>II</sup> are colored as indicated in the color key. NT, non-template-strand promoter DNA; T, template-strand promoter DNA.

## RESULTS

#### **Overall structure of Rob-TAC**

In order to obtain the structure of Rob-TAC, we used a nucleic-acid scaffold corresponding to positions -57 to +13 of micF promoter (positions numbered relative to the transcription start site; Figure 1A). MicF encodes an antisense RNA involved in multidrug resistance by repressing expression of the major membrane porin, OmpF (47,48). The *micF* DNA scaffold is composed of a 20 bp Rob binding box robbox (including recognition element A site with sequence of GCAC and recognition element B site with sequence of CAAA which overlaps with the -35 element), and a 13 bp non-complementary transcription bubble (-11 to +2) with a consensus -10 element, which was positioned at the catalytic center of RNAP for stabilizing core RNAP and allowing RNA synthesis (Figure 1A) (12,16,18,49). Meanwhile, our in vitro Mango-based transcription assay showed that, in the presence of Rob, E. coli RNAP efficiently activated transcription of micF-mango promoter, reflecting that these purified proteins are biologically relevant (Supplementary Figure S2A and B). By combing *E. coli* RNAP, Rob and *micF* DNA scaffold, we successfully assembled Rob-TAC complex in a stoichiometric manner as evidenced from SDS-PAGE and native PAGE analyses (Supplementary Figure S2A, C and D).

The frozen Rob-TAC complex was subsequently analyzed on a Titan Krios cryo-EM, and the data was sequentially processed, modelled and refined. Fortunately, we finally trapped two structures of Rob-TAC, namely Rob-TAC<sup>I</sup> and Rob-TAC<sup>II</sup>, at nominal resolutions of 4.06 and 4.57 Å, respectively (Figure 1B and C; Supplementary Figures S3–S7; Table 1). Local resolution calculation exhibits two central core RNAP at 3.5–4.5 Å and peripheral RNAP αCTD and Rob at 5.5–7.0 Å (Supplementary Figure S5C and 6C). There is only one Rob molecule present in each complex, simultaneously engaging RNAP  $\alpha$ CTD, RNAP  $\sigma^{70}$  R4 and *robbox* of promoter DNA, as is different from the reported dimeric transcription activators (13, 14). Though the two structures superimpose onto each other well at the densities of E. coli RNAP core enzyme, promoter DNA, and Rob NTD, the obvious extra density of

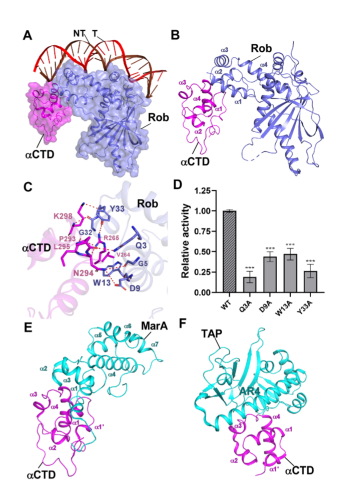
	Rob-TAC <sup>II</sup>	Rob-TAC <sup>I</sup>
Data collection and processing		
Microscope	Titan Krios	Titan Krios
Voltage $(kV)$	300	300
Detector	K2 summit	K3 summit
Electron exposure $(e/Å^2)$	50	50
Defocus range (µm)	1.4-2.2	1.4-2.2
Data collection mode	Count	Super resolution
Physical pixel size (Å/pixel)	1.307	1.100
Symmetry imposed	C1	C1
Initial particle images	478 388	962 734
Final particle images	335 021	386 188
Map resolution (Å) <sup>a</sup>	4.57	4.06
Refinement		
Map sharpening <i>B</i> -factor (Å)	-89	-101
Root-mean-square deviation		
Bond length (Å)	0.006	0.007
Bond angle ( <sup>0</sup> )	0.904	0.993
MolProbity statistics		
Clashscore	12.00	10.00
Rotamer outliers (%)	1.10	1.00
Cβ outliers (%)	0.0	0.0
Ramachandran plot		
Favored (%)	91.84	91.38
Outliers (%)	0.20	0.20

**Table 1.** Single particle cryo-EM data collection, processing, and modelbuilding for *E. coli* Rob-TAC<sup>II</sup>, Rob-TAC<sup>I</sup>

<sup>a</sup>Gold-standard FSC 0.143 cutoff criteria.

Rob CTD in Rob-TAC<sup>I</sup> exhibits significant differences from that of Rob-TAC<sup>II</sup>, with a rotation of  $\sim 35^{\circ}$  (Supplementary Figures S7D). In Rob-TAC<sup>I</sup>, Rob CTD retains in close proximity to RNAP aCTD and helix 1 from Rob NTD, possibly plays a potential role in RNAP recruitment and facilitates DNA binding (Supplementary Figures S7B and C). While in Rob-TAC<sup>II</sup>, Rob CTD moves away from RNAP  $\alpha$ CTD and helix 1 of Rob NTD, mainly interacts with the C-terminal portion of Rob NTD and the long linker connecting Rob NTD and Rob CTD, rendering the acidic loop of Rob CTD close to the downstream promoter (Figure 1A and B). Since the density and occupancy of Rob CTD in Rob-TAC<sup>I</sup> seem to be significantly weaker and lower than that in Rob-TAC<sup>II</sup>, we infer that the non-stable Rob-TAC<sup>I</sup> is most probably a pre-state of the stable Rob-TAC<sup>II</sup>. Therefore, we choose Rob-TAC<sup>II</sup> to be described in the following section.

Comparative structural analysis shows that E. coli RNAP containing  $\sigma^{70}$  in Rob-TAC structures maintains in a similar conformation to that of E. coli RNAP-promoter open complex (PDB ID: 6CA0) (10) with root mean square deviation (RMSD) of 1.410 Å (3144 Cas aligned) and 1.492 Å (3237 C $\alpha$ s aligned) in Rob-TAC<sup>I</sup> and Rob-TAC<sup>II</sup>, respectively. By aid of  $\beta'$  coiled coil,  $\sigma^{70}$ R2 contacts the consensus -10 element. RNAP accommodates the transcription bubble and downstream dsDNA in the same way as those in RPo (10) (Supplementary Figures S8 and S9). One distinguishing feature of Rob-TAC<sup>II</sup> is that Rob occludes  $\sigma^{70}R4$ from binding to the -35 element and specifically binds robbox. Furthermore, the co-crystal structure of Rob complexed with *micF* double stranded DNA could be well fitted into the upstream promoter density map, except for the obvious DNA bending towards Rob CTD (Supplementary Figure S7E), especially the acidic loop of Rob. Such DNA



**Figure 2.** The interactions between Rob and RNAP  $\alpha$ CTD. (A) Relative locations of Rob, *E. coli* RNAP  $\alpha$ CTD, and the upstream double-stranded DNA. (B) Rob interacts with *E. coli* RNAP  $\alpha$ CTD. (C) Detailed interactions between *E. coli* RNAP  $\alpha$ CTD and Rob. Salt-bridges are shown as red dashed lines. (D) Substitutions of Rob residues involved in interactions with *E. coli* RNAP  $\alpha$ CTD decreased *in vitro* transcription activity. Data for *in vitro* transcription assays are means of three technical replicates. Error bars represent  $\pm$  SEM of n = 3 experiments. Asterisk (\*\*\*) indicates highly significant (*P* value < 0.001) difference from the wild-type Rob analyzed by one-way ANOVA with Tukey's multiple comparison test, respectively. (E) Interface between *E. coli* MarA and RNAP  $\alpha$ CTD (PDB ID: 1XS9). (F) Interface between *T. thermophilus* RNAP  $\alpha$ CTD (PDB ID: 512D).

conformational change shares some similarities with that of MarA-*mar* binary complex (PDB ID: 1BL0) (31) (Figure 1B and C; Supplementary Figure S7B and C).

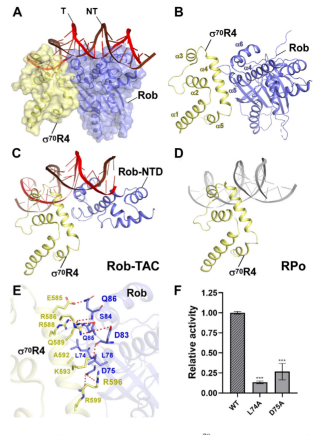
# RNAP $\alpha$ CTD is required for Rob-dependent transcription activation of class II promoters

Distinct from previous *in vitro* transcription assay that inferred RNAP  $\alpha$ CTD was not required for transcription activation of class II promoters (50), in Rob-TAC<sup>II</sup>, there exists abundant interactions between Rob NTD and RNAP  $\alpha$ CTD, burying a surface area of 149.39 Å<sup>2</sup> (Figure 2A-B). Evidently, residues Q3, G5, D9 and W13 from helix  $\alpha$ 1 of Rob form hydrogen bonds with residues P293 and N294 in the N-terminal loop of helix  $\alpha$ 1, and helix  $\alpha$ 4 of RNAP  $\alpha$ CTD, respectively. Residues G5 and G32 of Rob form van der Waals contacts with residues V264 and L295 of RNAP  $\alpha$ CTD (Figure 2C). Moreover, residue Y33 in the loop connecting helix  $\alpha 2$  and  $\alpha 3$  of Rob generates hydrogen bonds with residues from the DNA binding '265 determinant' (R265 and K298) of RNAP  $\alpha$ CTD (Figure 2C), indicative of DNA binding deficiency of RNAP  $\alpha$ CTD targeting at promoter UP elements (AT-rich DNA regions locate approximately from -40 to -60 of a promoter) (51). Consistent with this, mutation of the key residues of Rob-RNAP  $\alpha$ CTD interaction led to obvious decrease in both transcription activation activities by Mango-based transcription assay and Rob-TAC formation activities evidenced by EMSA (Figure 2D, Supplementary Figure S10A and B), suggesting the necessity of these residues to Rob-dependent transcription activation.

In comparison, structural analysis shows that interactions between Rob and RNAP aCTD in Rob-TAC<sup>II</sup> are similar to those between MarA and RNAP aCTD (MarA-RNAP  $\alpha$ CTD, PDB ID: 1XS9) (43) (with DNA omitted for clarity, Figure 2E) as well as between the class II transcription activator protein TAP (T. thermophilus TTHB099) and RNAP αCTD from TAP-dependent transcription activation complex (TAP-TAC, PDB ID: 5I2D, Figure 2F) (13), with buried surface areas being 123.81  $\text{\AA}^2$  and 182.54  $\text{\AA}^2$ , respectively. In Rob-TAC<sup>II</sup> and MarA-RNAP aCTD complexes, each activator uniquely contacts the DNA binding '265 determinant' of RNAP  $\alpha$ CTD by the homologous helices and loops. Likewise, in TAP-TAC, activating region 4 (AR4) of one TAP monomer interacts with a cluster of residues in the N-terminus of helix  $\alpha 4$ , the loop connecting helix  $\alpha 3$  and  $\alpha 4$ , and '265 determinant' of RNAP  $\alpha CTD$ (Figure 2F). These implies the important regulatory role of the '265 determinant' of RNAP αCTD.

# Rob occludes $\sigma^{70}$ R4 from binding to promoter -35 element and synergistically redirects RNAP to Rob regulons

Despite the critical interactions between  $\sigma^{70}R2$  and promoter consensus -10 element, interactions between  $\sigma^{70}R4$ and promoter consensus -35 element have also been extensively proved to be essential for RPo formation and transcription activation (3,5,10,12). While in case of class II promoters that contain activator binding box overlapping with -35 element to some extent,  $\sigma^{70}$ R4 remodeling is required to maintain successful transcription initiation. As expected, in Rob-TAC<sup>II</sup>, Rob forms several types of interactions with the long helix  $\alpha 5$  of  $\sigma^{70}$ R4, with a buried surface area of 40.46 Å<sup>2</sup> (Figure 3A–C). Thus, helix  $\alpha$ 5 and the adjacent loop of Rob in Rob-TAC<sup>II</sup> occlude  $\sigma^{70}$ R4 from binding to -35 element, rendering it makes weaker interactions with DNA (Figure 3C and D). Residues S84, Q85, Q86 in the loop connecting helix  $\alpha 5$  and  $\alpha 6$  of Rob form three hydrogen bonds with residues Q589, R588 and E585 in  $\sigma^{70}$  R4, respectively. Residues D75 and D83 of Rob simultaneously make ionic bonds with residues R596, R599, K593 and R586 from  $\sigma^{70}$ R4. Furthermore, residues of L74 and L78 from Rob form van der Waals contacts with the hydrophobic residue A592 from  $\sigma^{70}$ R4. In accordance with the above interactions (Figure 3E), our transcription and EMSA analyses of mutants also showed that the key residues are functionally relevant (Figure 3F; Supplementary Figure S10B). This observation is consistent with the previous genetic and bio-



**Figure 3.** The interactions between Rob and  $\sigma^{70}$ R4. (A) Relative locations of Rob,  $\sigma^{70}$ R4, and upstream double-stranded DNA in *E. coli* Rob-TAC. (**B**) Rob interacts with the  $\sigma^{70}$ R4.  $\sigma^{70}$ R4 and Rob are represented as yellow or blue cartoon, respectively. (**C**) Relative locations of Rob-NTD, *E. coli* RNAP  $\sigma^{70}$ R4 and the upstream double-stranded DNA. (**D**) Relative locations of *E. coli* RNAP  $\sigma^{70}$ R4 and the upstream typical –35 element DNA (PDB ID: 6CA0). (**E**) Detailed interactions between  $\sigma^{70}$ R4 and Rob. Hydrogen bonds are shown as red dashed lines. (**F**) Substitutions of the residues involved in Rob- $\sigma^{70}$  interactions reduced *in vitro* transcription activity. Data for *in vitro* transcription assays are means of three technical replicates. Error bars represent mean  $\pm$  SEM of n = 3 experiments. Asterisk (\*\*\*) indicates highly significant (*P* value < 0.001) difference from the wild-type Rob analyzed by one-way ANOVA with Tukey's multiple comparison test, respectively.

chemical experiments which suggested that Rob possibly occludes  $\sigma^{70}$ R4 from binding to promoter -35-hexamer and synergistically redirect RNAP to specific *robbox*-containing promoters (32).

#### Rob specifically interacts with two DNA major grooves of *rob*box and the acidic loop facilitates such interaction

As a representative example of the AraC/XylS family transcription activators, Rob harbors two highly conserved DNA-binding HTH motifs (designated as HTH<sup>A</sup> and HTH<sup>B</sup>, respectively) in Rob NTD (Figure 4A). Unlike MarA, which can insert its two conserved HTH motifs into the corresponding DNA major grooves of *mar* promoter and bends DNA by 35°, the co-crystal structure of Rob– *micF* showed that Rob could only insert helix  $\alpha$ 3 of HTH<sup>A</sup> into the A site of *robbox*, with helix  $\alpha$ 6 of HTH<sup>B</sup> contacting the phosphate backbone of B site, and thus rendering pro-

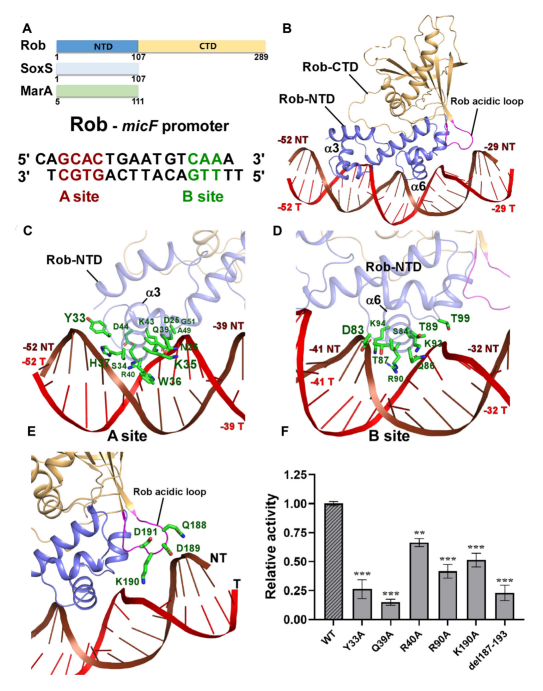


Figure 4. The interactions between Rob and the *micF* promoter DNA. (A) Domain architecture of Rob, SoxS and MarA (top panel); the sequences of the *micF* promoter for Rob protein with the A- and B-box sequences highlighted in red and green, respectively (bottom panel). (B) Rob in complex with the *micF* promoter DNA. Rob-NTD and Rob-CTD are represented as blue or orange cartoon, respectively. (C, D) Detailed interactions between Rob and the *micF* promoter DNA. Residues from Rob involved in interacting with A- or B-site sequences of *micF* promoter DNA are shown in green sticks. (E) Zoom-in view of the interactions between the acid loop and the *micF* promoter DNA. The acidic loop (residues 187-193) connecting strands  $\beta$ 3 and  $\beta$ 4 in the extra C-terminal domain of the Rob protein is highlighted in magenta. Residues from the acidic loop that contact DNA are shown in green sticks. (F) Substitutions of residues involved in Rob-DNA interactions suppressed *in vitro* transcription activity. Data for *in vitro* transcription assays are means of three technical replicates. Error bars represent  $\pm$  SEM of n = 3 experiments. Asterisk (\*\*\*) or (\*\*) indicates highly significant (*P* value < 0.001) or significant (*P* value < 0.01) difference from the wild-type Rob analyzed by one-way ANOVA with Tukey's multiple comparison test, respectively.

moter DNA unbent (Supplementary Figures S11A) (30). However, some other *in vivo* and *in vitro* assays on Rob held different opinions (32, 52) and it needs to be further clarified. Surprisingly, both helix  $\alpha 3$  of HTH<sup>A</sup> and helix  $\alpha 6$  of HTH<sup>B</sup> in Rob-TAC<sup>II</sup> insert into the DNA major grooves of robbox and make the DNA bent with an orientation similar to that of MarA (Figure 4A–D; Supplementary Figures S11B), revealing a general promoter remodeling mode for the AraC/XylS family transcription activators. By using helix  $\alpha 3$  of HTH<sup>A</sup> and helix  $\alpha 6$  of HTH<sup>B</sup>, Rob makes abundant interactions with the corresponding DNA major grooves of robbox (Figure 4C and D; Supplementary Figures S11C). As to the A site of robbox, residues W36 and S34 of Rob form hydrogen bonds with -45A and -47G of the non-template strand DNA, respectively. Residues Y33, H37 of Rob make contacts with the DNA phosphate backbone of the non-template strand DNA. Both of residues Q39 and R40 form two hydrogen bonds with the template strand DNA, indicating their critical roles in stabilizing DNA. Besides, several residues (D25, N26, K35, K43, D44, A49, G51) from helix  $\alpha$ 3 and the adjacent loops interacts with the DNA phosphate backbone of the template strand DNA. In respect to the B site of *robbox*, residues Q86 and R90 of Rob insert into the major groove and specifically recognize bases of -34T, -35T from the template strand DNA, and -37T, -38G from the non-template strand DNA (Figure 4D). In addition, residues of R55, D83, S84, T87, K94 make extensive contacts with the non-template strand DNA, while residues of Q86, K93, T89, T99 contact template strand DNA. Another striking feature of Rob-TAC<sup>II</sup> is that the acidic loop of Rob (spanning from S187 to E193) moves in close proximity to the bent micF promoter DNA. Structural analysis shows that residue K190 makes obvious ionic bonds with the DNA phosphate backbone of -31G and -32T from the template strand DNA (Figure 4E). In good agreement with these observations, substitutions of the above key residues involved in Rob-DNA interactions (Y33A, Q39A and R90A) and acidic loop (K190A, del187-193) confer significant defects in transcription activation activities verified by our Mango-based transcription assay (Figure 4F) and Rob-TAC formation activities (Supplementary Figure S10A and B), suggesting the importance of their functions. This is in accordance with the recently reported molecular dynamics simulation experiment which proposed that the acidic loop of Rob might facilitate interconversion between the distinct DNA binding modes observed in MarA and Rob (52). These evidences reveal the necessity and accessory role of the acidic loop for Robdependent transcription activation.

#### Rob CTD activates transcription through allosteric coordination with Rob NTD

Rob CTD is the most obvious feature that distinguish itself from the other subfamily AraC/XylS factors, and has been evidenced to be crucial for stability and activation activity of Rob (29). *In vivo* analysis inferred that Rob CTD possibly stabilizes Rob by protecting its vulnerable DNA-binding NTD from proteolytic degradation by ClpYQ and Lon proteases (32). Consistent with this, Rob CTD in our Rob-TAC structures forms 'ridge-like' projection sitting atop the N- terminal surface of Rob NTD in Rob-TAC<sup>I</sup> (Supplementary Figure S7B and C) or embracing the C-terminal surface of Rob NTD in Rob-TAC<sup>II</sup> (Figure 1B and C). This equilibrium enables Rob NTD well protected by Rob CTD and promoter DNA.

In E. coli, Rob is postulated as a global regulator of multidrug resistance. This is mainly attributed to the distinct Rob CTD which forms a global GyrI-like domain capable of binding small molecules (25,26,53). However, the detailed molecular mechanism is poorly determined. To gain insight into the ligand binding properties and the evoked conformation changes of Rob, we compared Rob from Rob-TAC<sup>II</sup> with other GyrI-like domains in complex with ligands from other transcription factors involved in bacterial stress response (18,53,55). Intriguingly, comparative structural analysis shows that Rob CTD superimposes well with the evolutionarily conserved GyrI-like domain family regulators, such as Staphylococcus aureus SAV2435 (PDB ID: 5KAW) with an RMSD of 2.21 Å (133 Cα aligned) (Figure 5A) (53). These GyrI-like domains are mainly composed of antiparallel β-strands, connecting loops and long characteristic helices, and have been proved to be adaptive for broad selectivity of ligand binding and biological signaling. These ligand binding pockets exhibit similar stereochemical properties and some connecting loops were also verified to play regulatory roles in the allosteric regulation process. Consistent with the above points, our molecular docking analysis of Rob with rhodamine 6G (R6G) and other four potential ligands (chenodeoxycholic acid, 4,4'-bipyridine, sodium decanoate and 2,2'-bipyridine) exhibits well fitted ligand binding pockets (Figure 5A-C; Supplementary Figure S12), indicating similar allosteric regulation mode. In accordance with the well-defined nature of ligand binding pockets, all the four potential ligands are mainly surrounded by cluster of aromatic and heterocyclic amino acids (Figure 5C and D; Supplementary Figure S12B and D), and mutation of the involved conserved residues largely compromises Rob-dependent transcription activity and Rob-TAC formation, especially Rob W164A and E262A (Figure 5E; Supplementary Figure S12E and F; Supplementary Figure S10E). This implies their biological significance in maintaining stereochemical conformation of Rob CTD and activating Rob-dependent transcription activity. As allosteric regulation model has been proposed for GyrI-like domain containing factors upon effector binding, it is therefore tempting to speculate that Rob may allosterically activate transcription through synergistical coordination with its effector sensor CTD and effector responder NTD.

# DISCUSSION

Bacteria has established diverse transcription regulation strategies to respond to a variety of environmental stresses and transcription activation is a well-known strategy. Though both *in vivo* and *in vitro* experiments have elucidated that the AraC/XylS family transcription activators play fundamental roles in multidrug resistance, metabolism of carbohydrate, heavy metals sensing, and virulence (20,22,23), long-standing questions regarding to their molecular mechanism still remain. Based on our structural and biochemical analysis of Rob-TAC, several key fea-

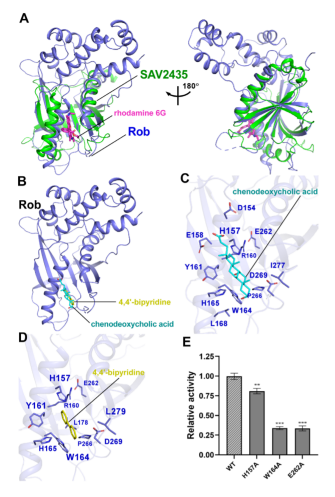


Figure 5. Molecular docking analysis of Rob with potential ligands. (A) Structural superimposition of the Gyrl-like family protein SAV2435 (PDB ID: 5KAW) onto the Rob from Rob-TAC. Rob and the SAV2435 are shown as blue and green cartoon. The ligand rhodamine 6G(R6G) is shown in magenta stick. (B) Rob docked with chenodeoxycholic acid and 4,4'-bipyridine. (C) Predicted binding pockets for chenodeoxycholic acid on Rob protein. Residues potentially interacted with chenodeoxycholic acid are shown in blue sticks. (D) Predicted binding pockets for 4.4'-bipyridine on Rob protein. Residues potentially interacted with 4.4'bipyridine are shown in blue sticks. (E) Substitutions of conserved residues involved in Rob-ligand interactions suppressed in vitro transcription activity in the presence of 100 µM chenodeoxycholic acid. Data for in vitro transcription assays are means of three technical replicates. Error bars represent  $\pm$  SEM of n = 3 experiments. Asterisk (\*\*\*) indicates highly significant (P value < 0.001) difference from the wild-type Rob analyzed by one-way ANOVA with Tukey's multiple comparison test, respectively.

tures of Rob as a member of AraC/XylS subfamily transcription activators can be delineated: (i) Rob acts as a monomer in Rob-TAC. This is different from the reported class II transcription activators, which usually function in the dimer form (Figure 6A) (13,15,18,56). (ii) Rob remodels RNAP by specific Rob-RNAP  $\alpha$ CTD interactions and Rob- $\sigma^{70}$ R4 interactions. These interactions synergistically redirect RNAP to target at specific Rob regulons. Similar to SoxS, MarA and TAP, Rob interacts with the DNA binding '265 determinant' of RNAP  $\alpha$ CTD and UP elements, and stabilizes the transcription activation complex. This is also in agreement with the mutational analysis that different determinants of RNAP aCTD are required for CRP-dependent transcription activation (13,43,56,57). The Rob- $\sigma^{70}$ R4 interactions occlude  $\sigma^{70}$ R4 from binding to canonical promoter elements, make it acting as a co-sigma factor. This also resembles the efficient  $\sigma$  appropriation mechanism mediated by MotA and AsiA (49). Sequence alignment of Rob and its homologs (MarA, SoxS, TetD and RamA) shows that the key residues (D9, W13, Y33 and D75) of Rob involved in the above interfaces are highly conserved (Supplementary Figure S13), implying similar significant proteinprotein interactions for the AraC/XylS-family transcription factors to remodel RNAP. (iii) Rob remodels promoter DNA not only by its two conserved HTH motifs, but also by aid of the acidic loop, which possibly facilitates DNA bending and promotes formation and stability of Rob-TAC. (iv) The global Gyrl-like domain containing Rob CTD serves as an environmental stimuli sensor, which structurally stabilizes and activates transcription through allosteric coordination with the stress responder Rob NTD.

It is worthwhile to mention several observations during our experiments. (i) Rob protein we purified exhibits obvious transcription activity; (ii) Rob CTD deletion mutant (residues 1-133) was mostly expressed as inclusion bodies and reduced greatly in solubility; (iii) The two alternative conformations of Rob-TAC we obtained show active 'dispersal' state, which may include an effector molecule during expression and purification. In agreement with this, the effectors play an insignificant role in enhancing transcription activities and Rob-TAC formation (Supplementary Figure S12E, S10A, B and E), and mutation of the potential ligand binding residues of Rob CTD apparently cause defects on transcription activities (Figure 5E; Supplementary Figure S12F). These data demonstrate that Rob CTD is essential for the stability and activity of Rob, and ligand is required for Rob-dependent transcription activity through binding to Rob CTD. Consistent with previous in vitro results, this also provides favorable evidence for the 'sequestration-dispersal' mechanism of Rob in vivo, which proposes Rob CTD as a novel off-on switch for the regulation of Rob's activation activity upon ligand binding (25,26,29,47). Thus, it is probably that in the 'sequestration' state, the residues involved in Rob-RNAP aCTD and Rob- $\sigma^{70}$ R4 interfaces might be enclosed by the large Rob CTD (and/or a cofactor) and retained in an inactive manner; while ligand binding is able to change the 'sequestration' conformation of Rob into a 'dispersal' one, and the above enclosed residues of Rob NTD will be exposed to engage RNAP and finally initiate transcription. (iv) To identify the two alternative conformations of Rob-TAC, further singlemolecule FRET assay was performed on the scaffold with a Cy5 fluorophore labelled on template strand of micFDNA2 and a Cy3B fluorophore on the 281-Cystine of Rob mutant (Supplementary Methods). The results showed only one FRET peak in each assay, and the FRET values were comparable and consistent with a distance of 5.5 nm from Rob-TAC<sup>II</sup> rather than 7.7 nm from Rob-TAC<sup>I</sup> (Supplementary Figure S14). Considering on this, we propose that the conformation of Rob-TACII is physiologically relevant and may have dominant significance for Rob-dependent transcription activation.

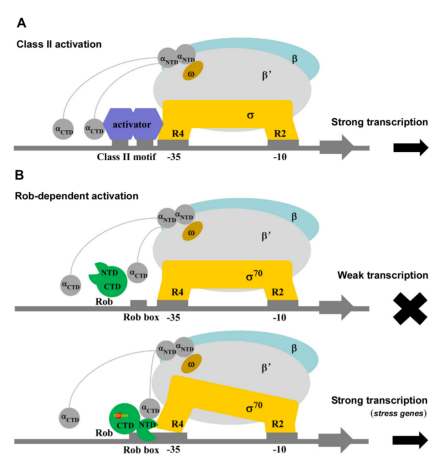


Figure 6. Proposed models for class-II-activator-dependent and Rob-dependent transcription activation. (A) Proposed working model for class-II-activator-dependent transcription activation. (B) Proposed working model for Rob-dependent transcription activation. Rob shows weak transcription activation activity in the absence of Rob-RNAP interactions, and activates the transcription of most genes with the promoter containing Rob binding box under stress.

In summary, a model for Rob-dependent transcription regulation can be proposed (Figure 6B). When environmental stimuli are absent, Rob CTD keeps in inactive sequestration state, and thus transcription cannot be initiated. Upon ligand binding, Rob CTD undergoes conformation change, turns Rob into de-sequestration active state, and relocates itself in cells. If Rob encounters a robbox-containing promoter, Rob CTD along with its acidic loop plays critical roles in the coordination with Rob NTD, promote two HTH motifs to insert into the corresponding DNA maior grooves, render DNA bent to form Rob-TAC<sup>II</sup> and finally initiate transcription efficiently. Consistent with the fact that Rob interacts with both RNAP  $\alpha$ CTD and  $\sigma^{70}$ R4 as revealed in the structure of Rob-TAC, Rob was found to form larger complex with RNAP in the absence of promoter DNA (Supplementary Figure S10C), suggestive of a 'pre-recruitment' mechanism, implying Rob possibly also activates transcription through a 'pre-recruitment' mechanism like SoxS (58). Besides, Rob also binds DNA in the absence of RNAP (Supplementary Figure S10D), which enables the DNA-binding Rob to recruit RNAP. Therefore, it is most probably that Rob-dependent transcription activation is accomplished through both a 'pre-recruitment' mechanism and a general 'recruitment' mechanism, and leading to enhanced formation of Rob-TAC (Supplementary Figure S10) (13,58). However, whether Rob activates transcription at the promoter clearance stage yet needs to be further clarified.

Comparative structural analysis shows that Rob CTD endows the evolutionarily conserved GyrI-like domain as the other GyrI-like domain containing transcription regulators (53). These GyrI-like domains possess similar residue composition, stereochemical properties, and potential allosteric regulation mechanism, implying their significant physiological roles. This resembles the recently reported *Bacil*lus subtilis BmrR (PDB: 7CKQ) (18) and E. coli EcmrR (PDB: 6XL5) involved in multidrug-sensing (55). However, there exists some big differences among them. Unlike the monomeric Rob, both BmrR and EcmrR function as dimers, sit on the distal face of promoter DNA, show no or few interactions with RNAP, and activate transcription mainly through promoter-distortion mechanism. This reveals that it is the two small conserved HTH motifs and the accessory acidic loop that accomplishes similar promoter remodeling functions to the above-mentioned dimers, which may reflect evolutionary significance to the AraC/XylS-family transcription factors.

By performing Pfam search using Rob sequence as a query, 2650 homologous protein sequences from various bacteria show the same architecture of 'HTH\_18 plus GyrI-

like' as Rob and are classified as transcription regulators, including AfrR and Caf1-R, revealing the global distribution and diverse functions of these 'Rob-like regulators'. Further sequence analysis shows that critical residues from HTH motifs of Rob NTD and aromatic residues in Rob CTD are highly conserved in these Rob-like regulators, which is in good agreement with the key residues defined in our Rob-TAC, implying their general regulatory roles during the physiological adaptation processes. However, new molecular transcription activation mechanism about them is yet to be explored experimentally. Meanwhile, it also unveils the potential role of Rob in providing a novel target for the antibacterial drug discovery.

During the review process of this manuscript, Hao et al. published the structure of RamA-class II activator complex. which also visualized a homologous activator interacts with RNAP  $\alpha$ CTD as well as  $\sigma^{70}$ R4 (59), exhibiting similar interfaces as those in TAP-TAC and our Rob-TAČII. Since CAP shares great similarity with TAP, though RNAP aCTD was invisible in CAP-TAC<sup>II</sup> due to its high flexibility (18), it might probably engage RNAP in a similar dynamic manner. These data collectively reveal a general regulation mode for class II activator-dependent transcription activation. Considering that RNAP  $\alpha$ CTD is conserved among all types of organisms, it could be generally applicable to other transcription factors in other organisms. However, not all class II activators remodel  $\sigma^{70}$  R4 of RNAP by occluding the interactions between  $\sigma^{70}$ R4 and -35 consensus element (such as TAP), this regulation mode of Rob might be an accessory way to efficiently remodel RNAP. As to class I promoters containing an activator binding site upstream of the -35 element, the corresponding activator is too far in distance to contact  $\sigma^{70}$ R4, and it may variably activate transcription by combination with the highly distinctive determinants of RNAP  $\alpha$ CTD (one or two),  $\sigma^{70}$ R4 and promoter DNA.

# DATA AVAILABILITY

Accession number for cryo-EM density map: EMD-32166 for of Rob-TAC<sup>I</sup> and EMD-32165 for Rob-TAC<sup>II</sup> (Electron Microscopy Data Bank). Accession number for atomic coordinates: 7VWZ for of Rob-TAC<sup>I</sup> and 7VWY for Rob-TAC<sup>II</sup> (Protein Data Bank).

# SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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Author contributions: J. S., F.L.W., F.F.L., L.W., Y.X., A.J.W., Y.L.J., S.J., F.G., Z.Z.F., J.C.L., Y.Z. performed the experiments. A.J.W., J.S., S.J. performed cryo-EM sample preparations and data collections. Y.F., W.L. performed cryo-EM structure determination. J.S., F.L.W. performed biochemical experiments. J.S., W.L. designed the study, J.S., W.L., S.W., Z.S. and Y. F. analyzed data and wrote the paper.

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