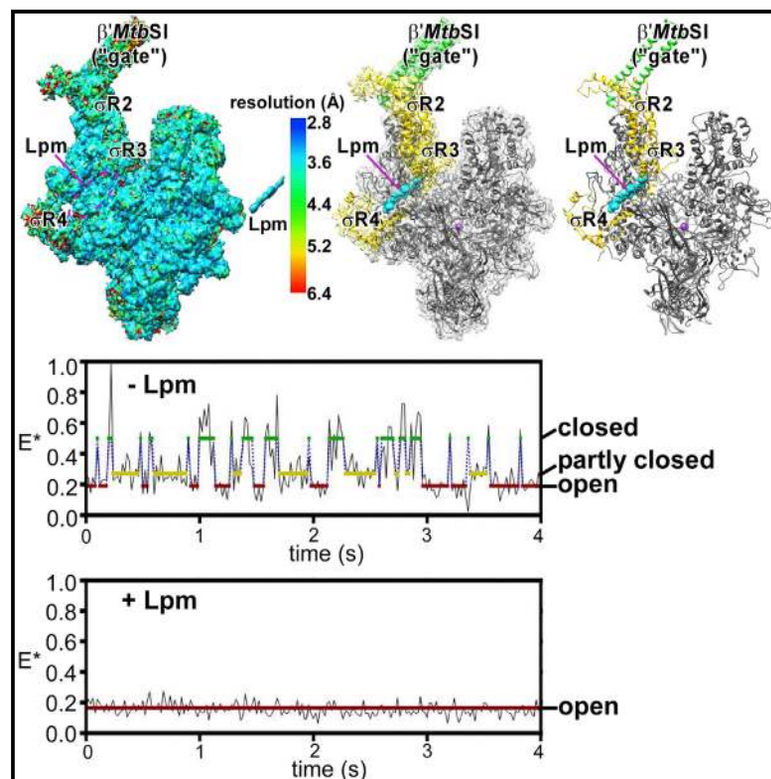


# Molecular Cell

## Structural Basis of Transcription Inhibition by Fidaxomicin (Lipiarmycin A3)

### Graphical Abstract



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### In Brief

Lin et al. report cryo-EM and single-molecule spectroscopic analyses of the antibacterial drug fidaxomicin bound to its molecular target, RNA polymerase. The results define the structure of the drug-target complex, show that fidaxomicin traps the RNA polymerase clamp in an open conformational state, and enable structure-based design of improved fidaxomicin analogs.

### Highlights

- Cryo-EM structure of drug-target complex
- Single-molecule spectroscopic analysis of effects of drug on target conformation
- Resistance determinant of drug
- Structure-activity relationships and structure-based design of drug analogs



# Structural Basis of Transcription Inhibition by Fidaxomicin (Lipiarmycin A3)

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## SUMMARY

Fidaxomicin is an antibacterial drug in clinical use for treatment of *Clostridium difficile* diarrhea. The active ingredient of fidaxomicin, lipiarmycin A3 (Lpm), functions by inhibiting bacterial RNA polymerase (RNAP). Here we report a cryo-EM structure of *Mycobacterium tuberculosis* RNAP holoenzyme in complex with Lpm at 3.5-Å resolution. The structure shows that Lpm binds at the base of the RNAP “clamp.” The structure exhibits an open conformation of the RNAP clamp, suggesting that Lpm traps an open-clamp state. Single-molecule fluorescence resonance energy transfer experiments confirm that Lpm traps an open-clamp state and define effects of Lpm on clamp dynamics. We suggest that Lpm inhibits transcription by trapping an open-clamp state, preventing simultaneous interaction with promoter –10 and –35 elements. The results account for the absence of cross-resistance between Lpm and other RNAP inhibitors, account for structure-activity relationships of Lpm derivatives, and enable structure-based design of improved Lpm derivatives.

## INTRODUCTION

Fidaxomicin is an antibacterial drug in clinical use for treatment of *Clostridium difficile* diarrhea (Lancaster and Matthews, 2012; Venugopal and Johnson, 2012). The active pharmaceutical ingredient of fidaxomicin, lipiarmycin A3 (Lpm; Kaufmann et al., 2015; Serra et al., 2017; Figure S1A), is a macrocyclic antibiotic with bactericidal activity against Gram-positive bacteria and

efflux-deficient strains of Gram-negative bacteria (Lancaster and Matthews, 2012; Venugopal and Johnson, 2012; Srivastava et al., 2011). Lpm functions by inhibiting bacterial RNA polymerase (RNAP; Talpaert et al., 1975; Sonenshein et al., 1977; Sonenshein and Alexander, 1979). Lpm exhibits no cross-resistance with the classic RNAP inhibitor rifampin (Rif; Sonenshein et al., 1977; Ebright, 2005) and inhibits transcription initiation at an earlier step than Rif (Sonenshein et al., 1977; Ebright, 2005; Tupin et al., 2010; Artsimovitch et al., 2012), suggesting that the binding site and mechanism of Lpm differ from those of Rif. Efforts spanning more than a decade to obtain structural information on RNAP-Lpm by use of X-ray crystallography have been unsuccessful.

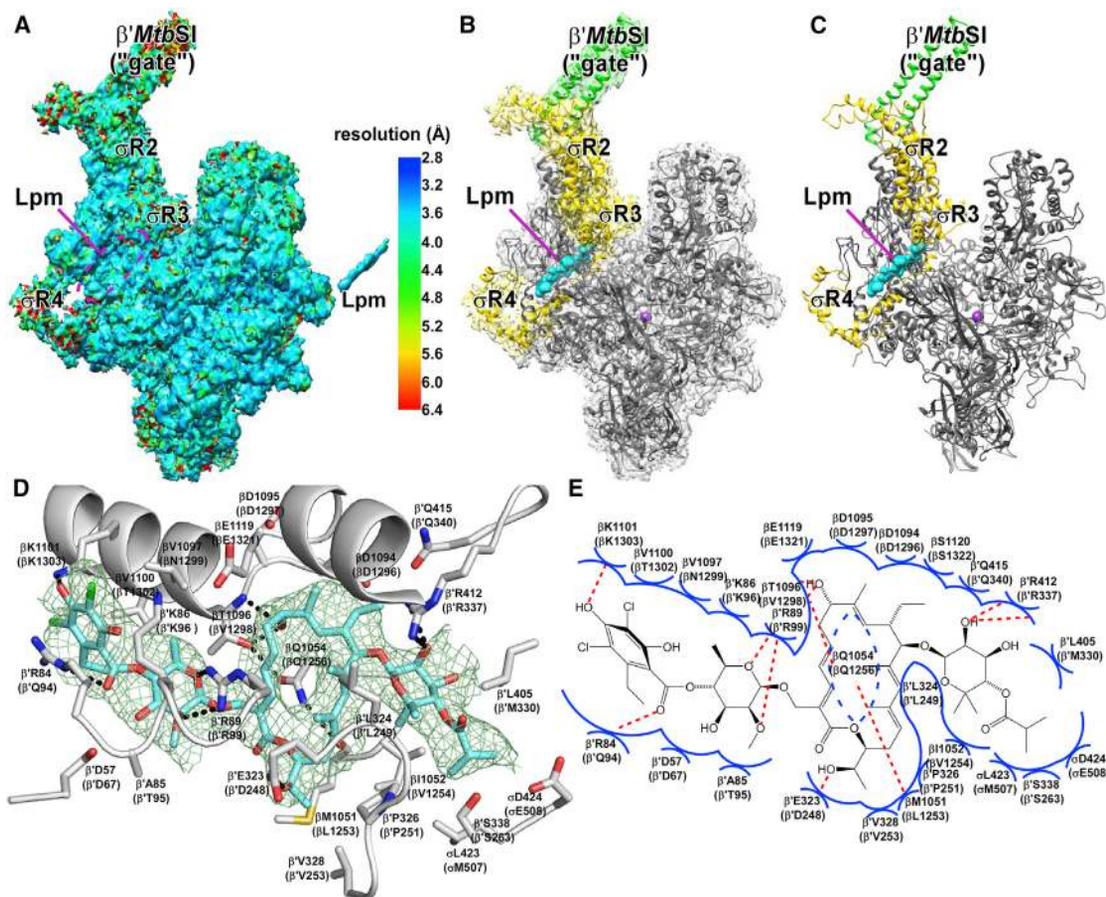
## RESULTS AND DISCUSSION

### Structure of *Mtb* RNAP-Lpm

Structural information on RNAP is available for three bacterial genera: the Gram-positive bacterial genus *Mycobacterium* (Hubin et al., 2017; Lin et al., 2017), the Gram-negative bacterial genus *Escherichia* (Murakami, 2013), and the *Thermus/Deinococcus*-clade bacterial genus *Thermus* (Zhang et al., 1999; Vassilyev et al., 2002). Lpm potently inhibits RNAP from genus *Mycobacterium* and genus *Escherichia* (concentrations resulting in 50% inhibition [ $IC_{50}$ ] = 0.3  $\mu$ M and 8  $\mu$ M, respectively; Talpaert et al., 1975; Kurabachew et al., 2008; Figure S1B) but only poorly inhibits RNAP from genus *Thermus* ( $IC_{50}$  = 200  $\mu$ M; Figure S1B). Accordingly, efforts in our laboratory to obtain structural and functional information on RNAP-Lpm focused on RNAP from genus *Mycobacterium* and genus *Escherichia*.

We first sought to obtain structural information on RNAP-Lpm by use of X-ray crystallography. However, we were unable to prepare crystals of RNAP-Lpm, despite numerous attempts at crystal soaking (assessing all reported crystal forms of RNAP from genus *Mycobacterium* and genus *Escherichia*) and





### Figure 1. Structure of *Mtb* RNAP-Lpm

(A) Density map for *Mtb* RNAP-Lpm (left) and Lpm (right), colored by local resolution.

(B) Density and atomic model for *Mtb* RNAP-Lpm. Gray, RNAP core other than  $\beta'$ MtbSI; green,  $\beta'$ MtbSI; yellow,  $\sigma$ ; cyan, Lpm; violet sphere, RNAP active-center  $Mg^{2+}$ .

(C) Atomic model for *Mtb* RNAP-Lpm. Colors are as in (B).

(D) RNAP-Lpm interactions (residues numbered as in *Mtb* RNAP and, in parentheses, as in *Escherichia coli* RNAP). Gray ribbons, RNAP backbone; gray sticks, RNAP carbon atoms; green mesh, density map for Lpm; cyan sticks, Lpm carbon atoms; red, blue, and green sticks, oxygen, nitrogen, and chlorine atoms, respectively; dashed lines, H-bonds.

(E) Summary of RNAP-Lpm interactions. Blue arcs, van der Waals interactions; red dashed lines, H-bonds.

See also Figures S1–S4.

co-crystallization (assessing crystallization conditions for all reported crystal forms of RNAP from genus *Mycobacterium* and genus *Escherichia*, and assessing more than  $10^4$  additional crystallization conditions by robotic screening).

We then sought to obtain structural information on RNAP-Lpm by use of cryo-electron microscopy (cryo-EM) with single-particle reconstruction (Merino and Raunser, 2017; Boland et al., 2017). Pilot experiments suggested that cryo-EM particle-orientation distributions using RNAP from genus *Mycobacterium* were more favorable for high-resolution structure determination than cryo-EM particle-orientation distributions using RNAP from genus *Escherichia*. Accordingly, our cryo-EM analysis focused on RNAP from genus *Mycobacterium*.

We determined a structure of *Mycobacterium tuberculosis* (*Mtb*) RNAP  $\sigma^A$  holoenzyme in complex with Lpm by use of cryo-EM with single-particle reconstruction (Merino and Raunser,

2017; Boland et al., 2017; Figures 1 and S1–S3; Table 1). The cryo-EM density map shows unambiguous density for RNAP, including the taxon-specific, *Mycobacterium*-specific sequence insertion ( $\beta'$ MtbSI or “gate”; Lin et al., 2017), for  $\sigma$  conserved regions 2, 3, and 4 ( $\sigma$ R2,  $\sigma$ R3, and  $\sigma$ R4, respectively), and for Lpm. The mean resolution of the structure is 3.5 Å (Figure S1F). Local resolution ranges from  $\sim$ 2.5–3.8 Å in central parts of the structure, including RNAP residues close to Lpm, to  $\sim$ 5–6.5 Å in peripheral parts of the structure, including  $\beta'$ MtbSI and  $\sigma$ R4 (Figures 1A and S1G and S2A). Local B-factors range from  $\sim$ 8–100 Å<sup>2</sup> in central regions to  $\sim$ 300–400 Å<sup>2</sup> in peripheral regions (Figure S2B). The map shows clear density for backbone and side-chain atoms of RNAP and individual functional groups of Lpm (Figures S2C and S2D).

The cryo-EM structure shows that the Lpm binding site is located at the base of the RNAP clamp (Landick, 2001; Cramer

**Table 1. Cryo-EM Data Collection and Data Analysis Statistics**

Structure	<i>Mtb</i> RNAP-Lpm	
EMDB and PDB Codes	EMDB: 4230	PDB: 6FBV
Data Collection		
Number of grids used	2	
Grid type	lacey carbon	
Microscope/detector	Titan Krios/ Gatan K2	
Voltage (kV)	300 kV	
Magnification	130,000×	
Recording mode	counting mode	
Dose rate (e-/s)	8	
Pixel size (Å/pix)	1.061	
Total dose (e-/Å <sup>2</sup> )	72.05	
Number of frames/movie	50	
Total exposure time (s)	10	
Defocus range (μm)	1.0–2.0	
Data processing	grid I	grid II
Number of micrographs	1,238	1,029
Number of particles picked	452,912	366,594
Number of particles after 2D/3D	126,977	97,412
Combined Dataset		
Particles used for final map calculation	68,895	
Map resolution (FSC 0.143; Å)	3.52	
Map resolution (ResMap median; Å)	3.5	
Structure/Map Fitting		
Resolution (Å)	3.5	
Map sharpening B factor (Å <sup>2</sup> )	–109	
Experimental map/model correlation	0.83	
Experimental map/2Fo-Fc correlation	0.98	
Average B factor (Å <sup>2</sup> )	95.3	
Total number of atoms	25,156	
Clash Score	13	
Ramachandran Plot		
Favored (%)	94.5	
Outliers (%)	0.03	
Root-mean-square deviations (RMSDs)		
Bond length (Å)	0.006	
Bond angle (°)	1.027	
Structure factors are from the cryo-EM map, and the 2Fo-Fc map is from the final model.		

et al., 2001; Chakraborty et al., 2012) and encompasses RNAP clamp  $\alpha$  helices  $\beta$ a16 $\alpha$ 1 and  $\beta$ '4 $\alpha$ 1 and RNAP switch-region (Cramer et al., 2001; Srivastava et al., 2011) switches SW2, SW3, and SW4 (Figures 1 and S3). Lpm binds to RNAP in a fully extended conformation similar to conformations in crystal struc-

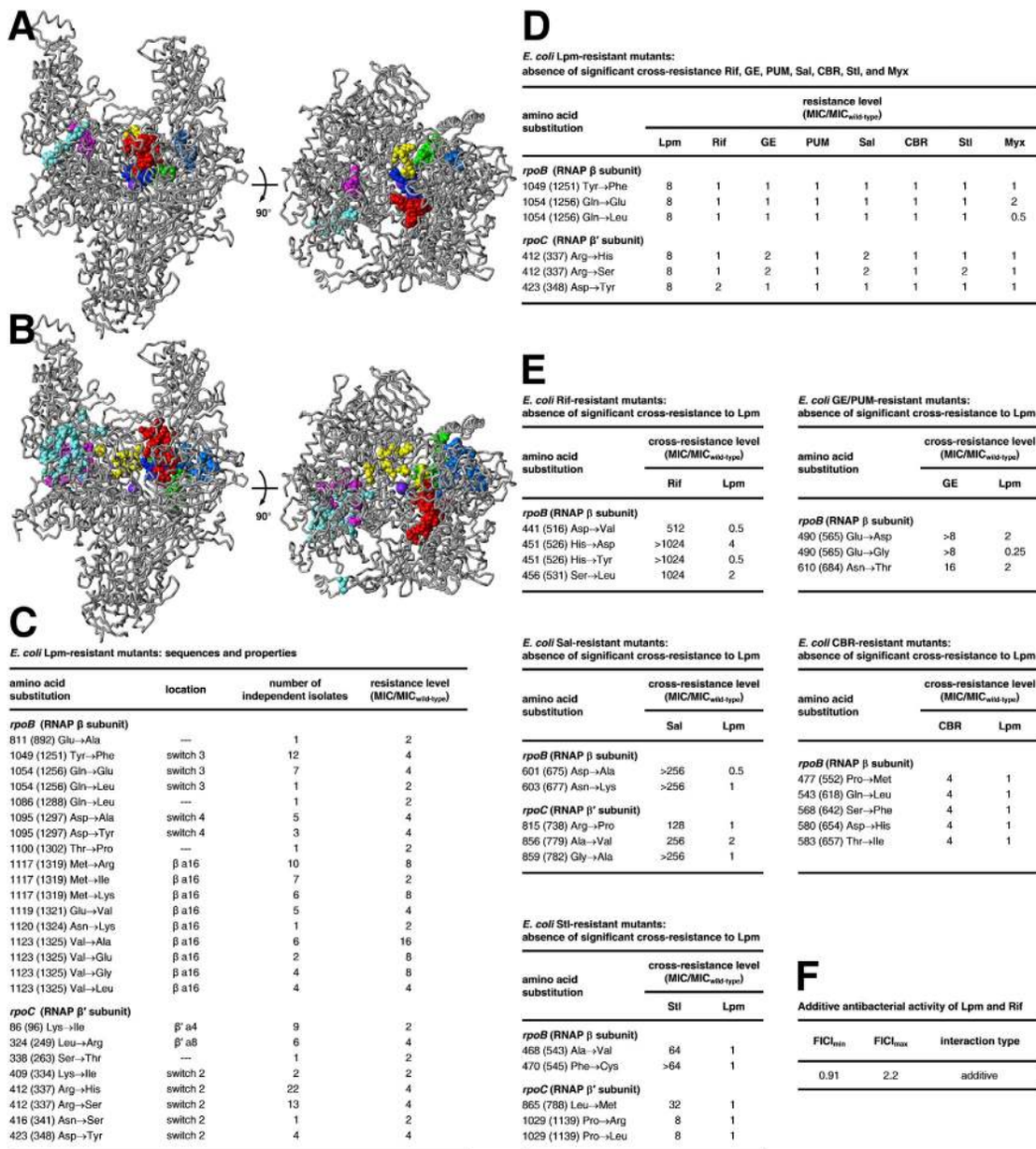
tures of Lpm alone (Figure S3A; Ihle et al., 2000; Serra et al., 2017). All five structural moieties of Lpm—*isobutyryl*,  $\beta$ -D-noviosyl, macrocycle,  $\beta$ -D-rhamnosyl, and homodichloro-orsellinyl—interact with RNAP (Figures 1D and 1E and S1A, S3B, and S3C). Lpm makes H-bonds with six RNAP residues:  $\beta$ T1096 (V1298),  $\beta$ K1101 (K1303),  $\beta$ 'R84 (Q94),  $\beta$ 'R89 (R99),  $\beta$ 'E323 (D248), and  $\beta$ 'R412 (R337) (residues numbered as in *Mtb* RNAP [UniProtKB: P9WGY9, UniProtKB: P9WGY7] and, in parentheses, as in *Escherichia coli* RNAP [UniProtKB: P0A8V2, UniProtKB: P0A8T7]). Alanine substitutions of five of these RNAP residues result in Lpm resistance, confirming that the interactions are functionally relevant (Figure S3D).

Sequence alignments show that residues of RNAP that contact Lpm are conserved in Gram-positive and Gram-negative bacterial RNAP, but are not conserved in human RNAP I, II, and III (Figures S4A and S4B), accounting for the ability of Lpm to inhibit both Gram-positive and Gram-negative bacterial RNAP but not human RNAP I, II, and III (Figure S4C).

The cryo-EM structure accounts for structure-activity relationships of Lpm analogs produced by metabolism (Babakhani et al., 2011), precursor feeding (Hochlowski et al., 1997), mutasynthesis (Xiao et al., 2011; Niu et al., 2011), and semi-synthesis (Figures 1D and 1E and S3B, S3C, and S5). The structure accounts for structure-activity relationships indicating functional importance of the Lpm *isobutyryl* moiety (“b1” versus Lpm),  $\beta$ -D-noviosyl moiety (“c1” versus “b2” and “a6”); macrocycle and C25 hydroxyl therein (“a5” versus Lpm),  $\beta$ -D-rhamnosyl moiety and C6' methyl therein (“e1” versus “d1” and “a7”); “a6” versus Lpm, and homodichloro-orsellinyl moiety and C8' methyl, C3' chlorine, and C5' chlorine therein (“d1” versus Lpm; “a2” to “a4” versus Lpm; “a8” versus “a6”). The observation that the Lpm C4' hydroxyl is exposed to solvent at the exterior of RNAP (Figures S3B and S3C) predicts that substituents, including large substituents, may be introduced at the C4' hydroxyl oxygen without eliminating the ability to inhibit RNAP, provided the H-bond-acceptor character of the oxygen is maintained. Validating this prediction, we have prepared a Lpm analog having a large substituent, benzyl, appended at the C4' hydroxyl oxygen by semi-synthesis from Lpm and have found the analog to retain high activity in inhibiting RNAP (Figure S5; analog “a1”); potencies within approximately an order of magnitude of that of Lpm for *Mtb* and *S. aureus* RNAP and equal to that of Lpm for *Escherichia coli* RNAP).

### Relationship between Binding Site and Resistance Determinant of Lpm and Binding Sites and Resistance Determinants of Other RNAP Inhibitors

The binding site on RNAP for Lpm (cyan in Figure 2A) does not overlap the binding sites of other RNAP inhibitors, including Rif and sorangicin (Sor) (Campbell et al., 2001, 2005), GE23077 (GE) and pseudoridimycin (PUM) (Zhang et al., 2014; Maffioli et al., 2017), CBR703 (CBR) and D-AAP-1 (AAP) (Lin et al., 2017; Feng et al., 2015; Bae et al., 2015b), salinamide (Sal) (Degen et al., 2014), streptolydigin (Stl) (Tuske et al., 2005; Termiakov et al., 2005), and myxopyronin (Myx), coralopyronin (Cor), ripostatin (Rip), and squaramide (SQ) (Mukhopadhyay et al., 2008; Belogurov et al., 2009; Molodtsov et al., 2015) (Figure 2A). The binding site for Lpm is close to, but does not overlap,



**Figure 2. Relationship between Binding Site and Resistance Determinant of Lpm and Binding Sites and Resistance Determinants of Other RNAP Inhibitors**

(A) Binding positions of Lpm (cyan; Figure 1), Rif and Sor (red; PDB: 1I6V, PDB: 1YNJ, PDB: 2A68, PDB: 2A69, PDB: 4KN4, PDB: 4KN7, PDB: 4OIR, and PDB: 5UHB), GE and PUM (dark blue; PDB: 4OIN, PDB: 4OIR, and PDB: 5X21), CBR and AAP (light blue; PDB: 4XSY, PDB: 4XSH, PDB: 4ZH2, PDB: 4ZH3, PDB: 4ZH4, PDB: 5UHE, and PDB: 5UHF), Sal (green; PDB: 4MEX), StI (yellow; PDB: 1ZYR), and Myx and SQ (magenta; PDB: 3DXJ, PDB: 3EQL, PDB: 4YFK, PDB: 4YFN, and PDB: 4YFX) mapped onto the structure of *Mtb* RNAP (gray; two orthogonal views; β'*Mtb*SI and σ omitted for clarity). Violet sphere, RNAP active-center Mg<sup>2+</sup>.

(B) Resistance determinants of Lpm (cyan; C), Rif and Sor (red; Campbell et al., 2001, 2005), GE and PUM (dark blue; Zhang et al., 2014; Maffioli et al., 2017), CBR703 and AAP (light blue; Lin et al., 2017; Feng et al., 2015; Bae et al., 2015b), Sal (green; Degen et al., 2014), StI (yellow; Tuske et al., 2005; Temiakov et al., 2005), and Myx, Cor, Rip, and SQ (magenta; Mukhopadhyay et al., 2008; Belogurov et al., 2009; Molodtsov et al., 2015) mapped onto the structure of *Mtb* RNAP.

(C) Sequences and properties of *E. coli* Lpm-resistant mutants (residues numbered as in *Mtb* RNAP and, in parentheses, as in *Escherichia coli* RNAP).

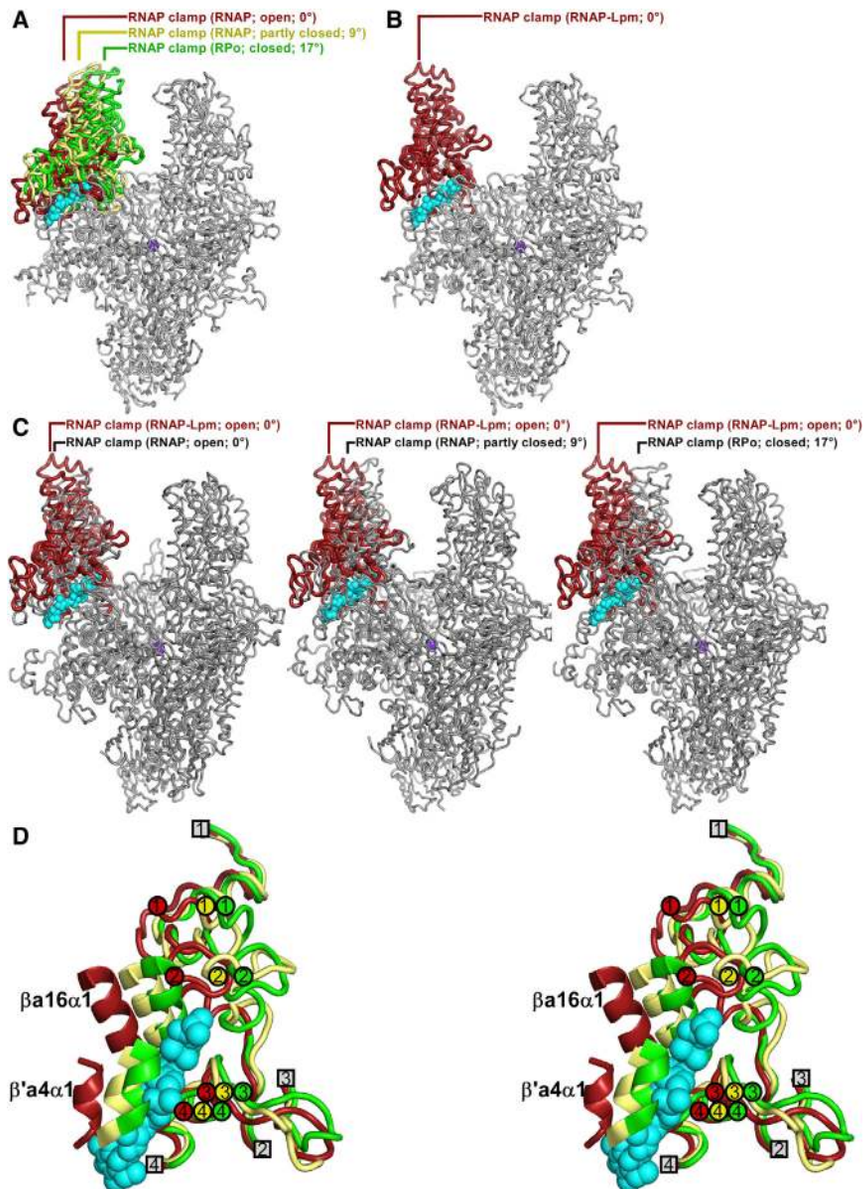
(D) Absence of significant cross-resistance of *E. coli* Lpm-resistant mutants to Rif, GE, PUM, Sal, CBR, StI, and Myx.

(E) Absence of significant cross-resistance of *E. coli* Rif-, GE/PUM-, Sal-, CBR-, and StI-resistant mutants to Lpm.

(F) Additive antibacterial activity of Lpm and Rif.

(Resistance and cross-resistance levels for Lpm-, CBR-, and StI-resistant mutants are from strains having the mutant RNAP subunit gene on a plasmid and the corresponding wild-type RNAP subunit gene on the chromosome [merodiploid strains; see STAR Methods]. Resistance and cross-resistance levels for

(legend continued on next page)



**Figure 3. Effects of Lpm on RNAP Clamp Conformation: Cryo-EM Data**

(A) RNAP open (red), partly closed (yellow), and closed (green) clamp conformational states. *Mtb* RNAP-Lpm main mass (view orientation as in Figures 1 and 2) and, superimposed, RNAP clamps from crystal structures of *Tth* RNAP in different crystal lattices (*Tth* RNAP core enzyme with open clamp [red; structure determined in this work to enable comparison of open clamp, partly closed clamp, and closed clamp states of RNAP from a single bacterial species; Table 2], *Tth* RNAP holoenzyme with partly open clamp [yellow, PDB: 5TMC], and *Tth* RPO [green; PDB: 4G7H]). (B) RNAP open clamp state in *Mtb* RNAP-Lpm. Red, clamp in *Mtb* RNAP-Lpm. Other colors are as in (A).

(C) Comparisons of RNAP clamp conformation in *Mtb* RNAP-Lpm to RNAP clamp conformations in crystal structures of *Tth* RNAP core enzyme with open clamp (left; Table 2), *Tth* RNAP holoenzyme with partly closed clamp (center; PDB: 5TMC), and *Tth* RPO with closed clamp (right; PDB: 4G7H). Each image shows *Mtb* RNAP-Lpm (clamp in red; view orientation and other colors as in Figures 1 and 2) and, superimposed, the RNAP clamp of the comparator structure (gray).

(D) Stereodiagram of the Lpm binding site showing conformations of the RNAP structural elements SW1, SW2, SW3, SW4, βa16α1, and β'a4α1 in open-clamp (red; *Mtb* RNAP-Lpm), partly-closed-clamp (yellow; PDB: 5TMC), and closed-clamp (green; PDB: 4G7H) states. RNAP switches SW1, SW2, SW3, and SW4 are shown with ends that connect to the RNAP clamp as numbered circles and ends that connect to the RNAP main mass as numbered squares. Cyan, Lpm.

the binding site of Myx, Cor, Rip, and SQ, which bind to the RNAP switch region, contacting SW1 and the face of SW2 opposite the face contacted by Lpm. The binding site for Lpm is far from the binding sites of other RNAP inhibitors.

Amino acid substitutions conferring resistance to Lpm were identified by sequencing spontaneous *Bacillus subtilis* Lpm-resistant mutants (Sonenshein et al., 1977; Ebright, 2005); additional substitutions were identified by sequencing induced Lpm-resistant mutants of *E. coli* RNAP (Ebright, 2005); and, in this work, a full set of substitutions was identified by extending the latter method (Figures 2B and 2C). Lpm-resistant substitutions were

obtained at ten sites in the RNAP β subunit and seven sites in the RNAP β' subunit (Figures 2B and 2C). All sites of Lpm-resistant substitutions are located in the immediate vicinity of the Lpm binding site (Figure 2B), and all high-level (≥4-fold) Lpm-resistant substitutions involve RNAP residues that make direct contact with Lpm or that contact other RNAP residues that, in turn, make direct contact with Lpm (Figure S5A). The resistance determinant for Lpm does not overlap the resistance determinants of other RNAP inhibitors (Figure 2B). Consistent with the absence of overlap of binding sites and resistance determinants, mutants resistant to Lpm are not cross-resistant to other RNAP inhibitors (Figures 2D and S6B), and, reciprocally, mutants resistant to other RNAP inhibitors are not cross-resistant to Lpm (Figures 2E and S6C). Also consistent with the absence of overlap of binding sites and resistance determinants, Lpm and Rif exhibit additive antibacterial activity when co-administered (Figure 2F).

Lpm-, CBR-, and StI-resistant mutants are from strains having the mutant RNAP subunit gene on the chromosome and no corresponding wild-type RNAP subunit gene [non-merodiploid strains; see STAR Methods]. Resistance and cross-resistance levels are expected to be lower for merodiploid strains than for non-merodiploid strains.)

See also Figure S5.

**Table 2. X-Ray Crystallography Data Collection and Data Analysis Statistics**

Structure	<i>Tth</i> RNAP core enzyme
PDB code	PDB: 6ASG
Data Collection	
Data collection source	APS-19ID
Space group	R3
Cell Dimensions	
a, b, c (Å)	280.7, 280.7, 184.9
$\alpha$ , $\beta$ , $\gamma$ (°)	90.0, 90.0, 120.0
Resolution	50.0–3.8 (4.0–3.8)
Number of Unique Reflections	61,395
R <sub>merge</sub>	0.245 (0.704)
R <sub>meas</sub>	0.275 (0.830)
R <sub>pim</sub>	0.124 (0.430)
Highest resolution shell CC <sub>1/2</sub>	0.418
I/ $\sigma$ (I)	5.0 (1.2)
Completeness (%)	99.2 (92.3)
Redundancy	4.8 (3.2)
Refinement	
Resolution (Å)	42.5–3.8
Number of unique reflections	50,511
Number of test reflections	2,611
ssR <sub>work</sub> /R <sub>free</sub>	0.22/0.27 (0.27/0.32)
Number of Atoms	
Protein	23,551
Ligand/ion	3
RMSDs	
Bond length (Å)	0.003
Bond angle (°)	0.658
MolProbity Statistics	
Clash score	15.2
Rotamer outliers (%)	1.0
C $\beta$ outliers (%)	0
Ramachandran Plot	
Favored (%)	94.1
Outliers (%)	0.4
Data for the highest-resolution shells are shown in parentheses.	

### Effects of Lpm on RNAP Clamp Conformation: Cryo-EM Data

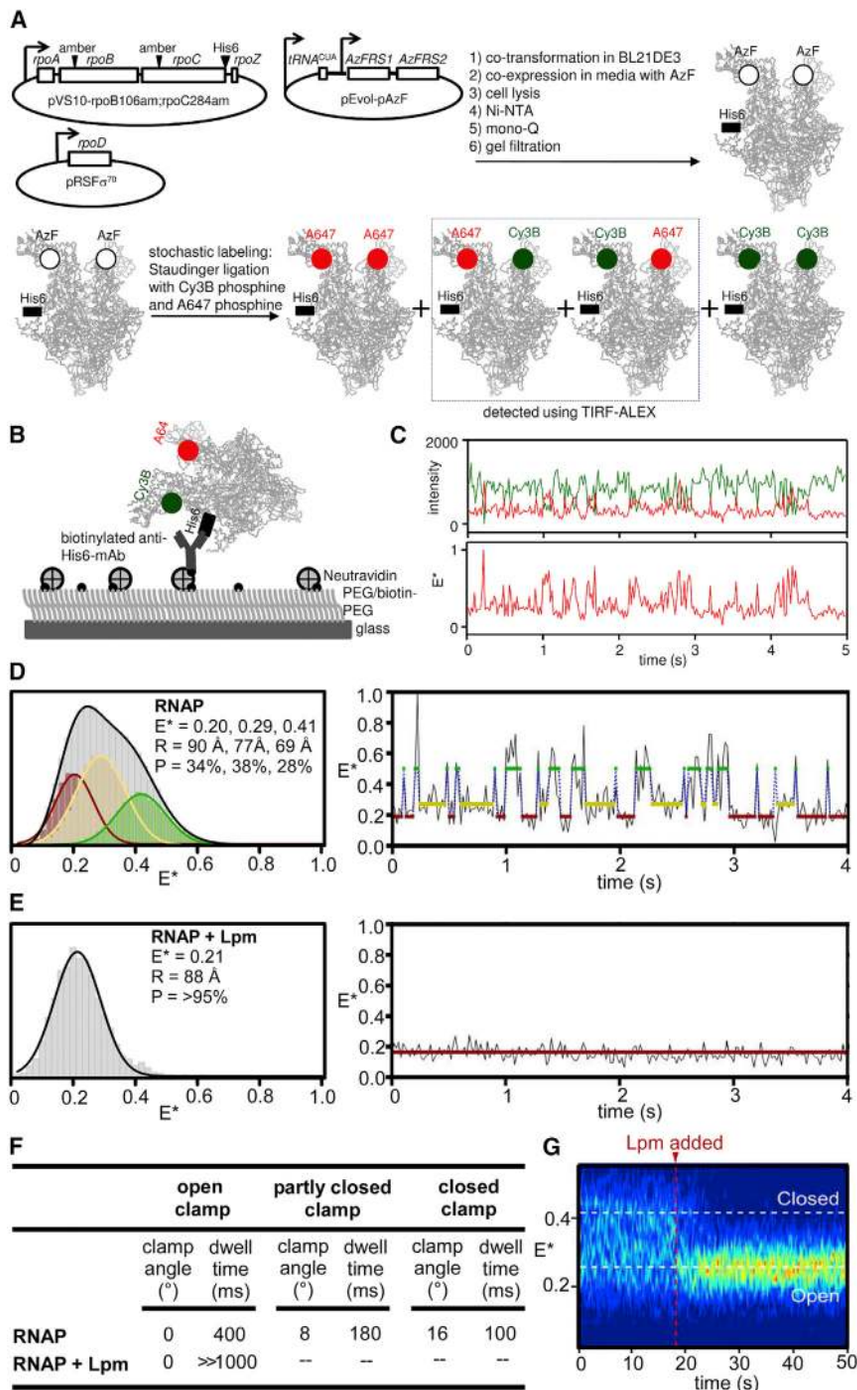
The RNAP clamp adopts different conformations—ranging from open, to partly closed, to closed—in different crystal structures (Landick, 2001; Cramer et al., 2001; Chakraborty et al., 2012; Figure 3A). Differences in clamp conformation arise from rigid-body rotations about a hinge formed by the RNAP switch region (Landick, 2001; Cramer et al., 2001; Chakraborty et al., 2012). Importantly, differences in clamp conformation are observed even in crystal structures of RNAP from a single bacterial species, indicating that differences in clamp conformations are state differences and not species differences (Figure 3A; Table 2).

In the cryo-EM structure of *Mtb* RNAP-Lpm, the clamp adopts an open conformational state, superimposable on the open state in a crystal structure of *Thermus thermophilus* (*Tth*) RNAP core enzyme (Table 2), but differing by 9° from the partly closed state in a crystal structure of *Tth* RNAP holoenzyme (PDB: 5TMC), and differing by 17° from the closed state in crystal structures of catalytically competent RNAP-promoter transcription initiation complexes (*Tth* RPo and *Mtb* RPo; PDB: 4G7H and PDB: 5UH5, respectively) (Figures 3B and 3C). The interactions Lpm makes with  $\beta$ a16 $\alpha$ 1,  $\beta'$ a4 $\alpha$ 1, SW2, SW3, and SW4 require an open-clamp conformation (Figure 3D). Each of these regions is positioned to accommodate Lpm in the open state, but not in partly closed and closed states. In particular,  $\beta$ a16 $\alpha$ 1 and  $\beta'$ a4 $\alpha$ 1 would severely clash with Lpm in partly closed and closed states (Figure 3D). We infer that Lpm, through its interactions with its binding site, induces or stabilizes an open-clamp conformation.

### Effects of Lpm on RNAP Clamp Conformation: Single-Molecule FRET Data

RNAP clamp conformation in solution can be monitored by single-molecule fluorescence resonance energy transfer (FRET) experiments assessing distances between fluorescent probes incorporated specifically at the tips of the two walls of the RNAP active-center cleft (Chakraborty et al., 2012; Figures 4A–4C). Single-molecule FRET experiments analyzing the RNAP holoenzyme in the absence of Lpm show a broad multimodal distribution of FRET efficiencies, indicative of open, partly closed, and closed states (Figure 4D, left, gray curve). In contrast, analogous experiments in the presence of Lpm show a narrow unimodal peak with a FRET efficiency, indicative of the open state (Figure 4E, left). We conclude that Lpm traps the open-clamp state in solution. Single-molecule time traces in the absence of Lpm show static and dynamic molecules; the dynamic molecules (34%, n = 207) show transitions between three distinct FRET efficiency levels—corresponding to open-clamp, partly-closed-clamp, and closed-clamp states, with dwell times on the 100- to 400-ms timescale—indicating dynamic transitions between interconverting clamp states (Figures 4D, left, colored curves, 4D, right, and 4F). In contrast, analogous single-molecule time traces in the presence of Lpm show only the FRET efficiency level corresponding to the open-clamp state, indicating stable trapping of the open-clamp state (Figures 4E, right, and 4F). Single-molecule time traces before and after addition of Lpm show a rapid (< 2 s) transition from the multimodal, dynamic clamp profile of RNAP in the absence of Lpm to the unimodal, non-dynamic clamp profile of RNAP in the presence of Lpm, demonstrating rapid trapping of the open-clamp state (Figure 4G). We conclude, based both on the cryo-EM structure and FRET, that Lpm traps the open-clamp state.

Strikingly, the effect of Lpm on RNAP clamp conformation is *opposite* the effects of Myx, Cor, and Rip on RNAP clamp conformation. Lpm, through interactions with RNAP switch-region SW2, SW3, and SW4, traps an open-clamp state (Chakraborty et al., 2012); in contrast, Myx, Cor, and Rip, through interactions with SW1 and the opposite face of SW2, trap a closed-clamp state (Figures 3 and 4). The different effects of these inhibitors on RNAP clamp conformation presumably underlie their different effects on isomerization of RPo to RPe



**Figure 4. Effects of Lpm on RNAP Clamp Conformation: Single-Molecule FRET Data**

(A) Use of unnatural amino-acid mutagenesis, Staudinger ligation with Cy3B-phosphine and Alexa 647-phosphine, and total-internal-reflection fluorescence microscopy with alternating-laser excitation microscopy (TIRF-ALEX) to obtain FRET data for single molecules of RNAP having fluorescent probes at the tips of the walls of the RNAP active-center cleft (STAR Methods). Green, fluorescent donor probe Cy3B; red, fluorescent acceptor probe Alexa 647; black square, hexahistidine tag.

(B) Surface immobilization of fluorescently probe-labeled RNAP for TIRF-ALEX.

(C) Time trace of donor emission intensity (green) and acceptor emission intensity (red) (top) and corresponding time trace of donor-acceptor FRET efficiency (bottom).

(D) Single-molecule FRET data for RNAP holoenzyme in the absence of Lpm. Left: histogram. Gray, all states; colors, hidden Markov model (HMM)-assigned open, partly closed, and closed states (red, yellow, and green, respectively). Right: time trace with HMM-assigned open, partly closed, and closed states (red, yellow, and green, respectively).

(E) Single-molecule FRET data for the RNAP holoenzyme in the presence of Lpm (histogram and time trace as in D).

(F) Summary of RNAP clamp angles and dwell times for open, partly closed, and closed clamp states in the absence and presence of Lpm.

(G) Time trace for RNAP holoenzyme before and after addition of Lpm.

RNA hairpins, trigger RNAP clamp opening by binding to the site.

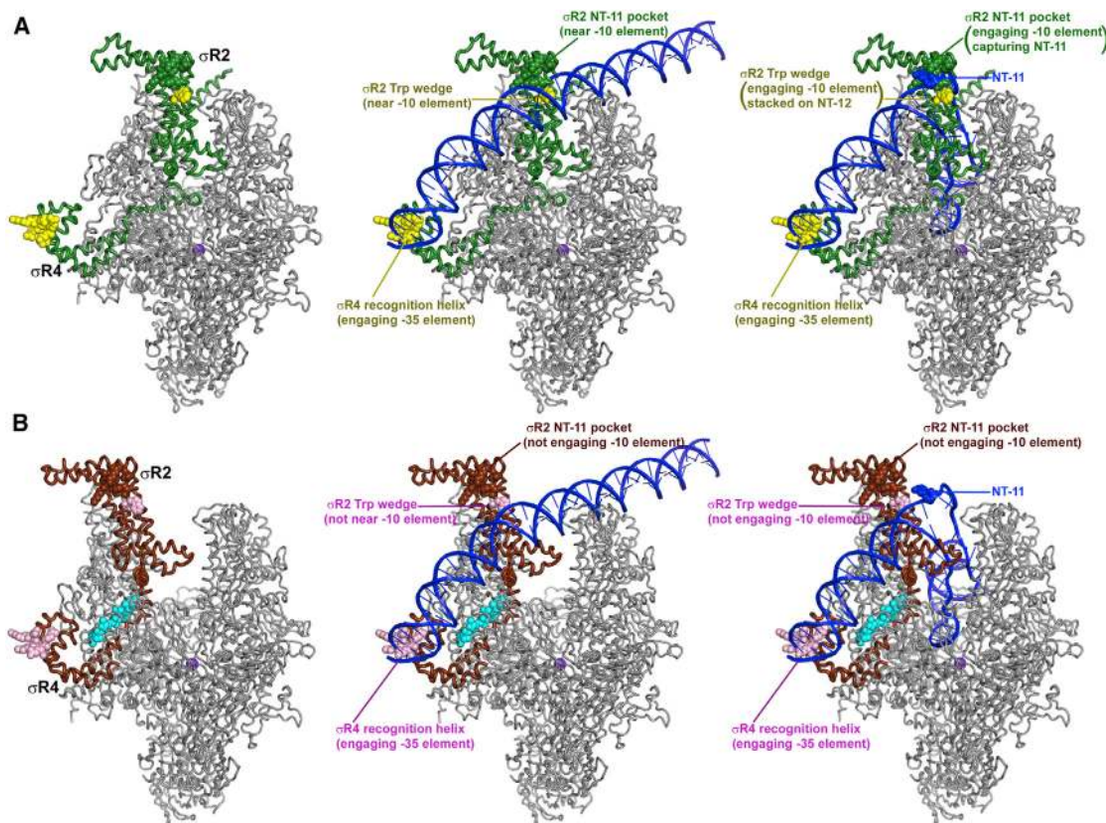
### Mechanism of Transcription Inhibition by Lpm

The effects of Lpm on RNAP clamp conformation suggest a model for the mechanism by which Lpm inhibits isomerization of R<sub>Pc</sub> to R<sub>Po</sub> (Sonenshein and Alexander, 1979; Ebright, 2005; Tupin et al., 2010; Artsimovitch et al., 2012) in transcription initiation (Figure 5). Because  $\sigma$ R2 (the  $\sigma$  module that recognizes promoter  $-10$  elements; Ruff et al., 2015; Zhang et al., 2012; Zuo and Steitz, 2015; Bae et al., 2015a; Feng et al., 2016) interacts with the RNAP clamp and  $\sigma$ R4 (the  $\sigma$  module that recognizes promoter  $-35$  elements; Ruff et al., 2015; Zuo and Steitz, 2015; Bae et al., 2015a; Feng et al., 2016) interacts with the main mass of RNAP, differences in RNAP clamp conformation necessarily result in differences in the spatial relationship of  $\sigma$ R2 and  $\sigma$ R4 (Figure 5, left). We propose that, in the closed-clamp state,  $\sigma$ R2 and  $\sigma$ R4 can engage simultaneously with promoter  $-10$  and  $-35$  elements during formation of R<sub>Po</sub> (Figure 5A, center and right), but, in the open-clamp state,  $\sigma$ R2 and  $\sigma$ R4

(inhibition of an early step in isomerization by Lpm; inhibition of a later step in isomerization by Myx, Cor, and Rip; Tupin et al., 2010; Artsimovitch et al., 2012).

The binding site on RNAP for Lpm overlaps the proposed binding site on RNAP for pause-inducing and termination-inducing RNA hairpins (Hein et al., 2014). It is attractive to speculate that the Lpm binding site is an “allosteric trigger” for RNAP clamp opening and that ligands of this site, including not only Lpm but also pause-inducing and termination-inducing

interacts with the RNAP clamp and  $\sigma$ R4 (the  $\sigma$  module that recognizes promoter  $-35$  elements; Ruff et al., 2015; Zuo and Steitz, 2015; Bae et al., 2015a; Feng et al., 2016) interacts with the main mass of RNAP, differences in RNAP clamp conformation necessarily result in differences in the spatial relationship of  $\sigma$ R2 and  $\sigma$ R4 (Figure 5, left). We propose that, in the closed-clamp state,  $\sigma$ R2 and  $\sigma$ R4 can engage simultaneously with promoter  $-10$  and  $-35$  elements during formation of R<sub>Po</sub> (Figure 5A, center and right), but, in the open-clamp state,  $\sigma$ R2 and  $\sigma$ R4



### Figure 5. Mechanism of Transcription Inhibition by Lpm

(A) RNAP holoenzyme (left), RPc (center; DNA as in Ruff et al., 2015), and RPo (right; DNA as in Zuo and Steitz, 2015; Bae et al., 2015a; Feng et al., 2016) for RNAP in the absence of Lpm (closed clamp).  $\sigma$ R2 and  $\sigma$ R4 simultaneously engage promoter -10 and -35 elements in RPc and RPo. Atomic coordinates are from PDB: 5UH5. Green ribbons,  $\sigma$ ; green surface,  $\sigma$ R2 NT-11 pocket; yellow surfaces,  $\sigma$ R2 Trp wedge and  $\sigma$ R4 recognition helix; blue, DNA. Other colors are as in Figures 1 and 2. (B) As (A), but for RNAP in the presence of Lpm (open clamp).  $\sigma$ R2 and  $\sigma$ R4 cannot simultaneously engage promoter -10 and -35 elements in RPc and RPo. Brown ribbons,  $\sigma$ ; brown surface,  $\sigma$ R2 NT-11 pocket; pink surface,  $\sigma$ R2 Trp wedge and  $\sigma$ R4 recognition helix; cyan, Lpm. Other colors are as in (A).

cannot engage simultaneously with promoter -10 and -35 elements (Figure 5B, center and right). In particular, we propose that the  $\sigma$ R2 “Trp wedge,” which intercalates into DNA and stacks on the non-template-strand nucleotide at promoter position -12, nucleating promoter unwinding in formation of RPo (Bae et al., 2015a; Feng et al., 2016), can engage DNA only in the closed-clamp state, and that the  $\sigma$ R2 “NT-11 pocket,” which captures an unstacked non-template-strand nucleotide at promoter position -11, stabilizing promoter unwinding upon formation of RPo (Zhang et al., 2012; Zuo and Steitz, 2015; Bae et al., 2015a; Feng et al., 2016), can engage DNA only in the closed-clamp state (Figure 5, center and right). We suggest that the inability of the open-clamp state to engage simultaneously with -10 and -35 elements occurs during late steps in the formation of RPo, when interactions of DNA downstream of the -10 element with the RNAP active-center cleft constrain the -10 element to be near the RNAP active-center cleft, in a position unreachable by  $\sigma$ R2 in the open-clamp state (Figure 5, right), and possibly also occurs during earlier steps in formation of RPo, if, as proposed by Vassylyev et al. (2002) and NandyMazumdar et al. (2016), interactions of DNA downstream of the -10 element with the opposite wall of the RNAP active-center cleft constrain the

-10 element to be near the RNAP active-center cleft, in a position unreachable by  $\sigma$ R2 in the open-clamp state (Figure 5, center). The proposal that Lpm inhibits transcription initiation by trapping an open-clamp state, thereby preventing simultaneous engagement of -10 and -35 elements, is consistent with recently proposed models for transcription initiation (Feklistov et al., 2017).

### Prospect

The methods and results of this report provide a strategy to obtain structural information for any RNAP inhibitor, irrespective of species selectivity and effects on RNAP conformation. The use of cryo-EM to determine the structure of RNAP-Lpm overcame two obstacles that had precluded structure determination by X-ray crystallography: (1) the species selectivity of Lpm (which only poorly inhibits *Thermus* RNAP; Figure S1B) and (2) trapping by Lpm of an RNAP open-clamp conformation (which is incompatible with known crystal lattices of non-*Thermus* bacterial RNAP).

The results of this report provide fundamental information regarding the binding site and mechanism of Lpm and provide insights for antibacterial drug development involving Lpm. The primary limitation of Lpm as an antibacterial drug is that Lpm

has no systemic bioavailability and, thus, is ineffective against systemic infections such as tuberculosis and *S. aureus* blood and lung infections (Srivastava et al., 2011; Lancaster and Matthews, 2012; Venugopal and Johnson, 2012). The chemical strategy and synthetic procedure demonstrated here for appending substituents at the Lpm C4' hydroxyl (Figure S5; "a1") provides a means to overcome this limitation. It should be possible to append substituents improving physical and pharmacokinetic properties, nanoparticle drug-delivery modules, or even other antibacterial-agent pharmacophores, at the Lpm C4' hydroxyl while retaining RNAP-inhibitory activity.

## STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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## SUPPLEMENTAL INFORMATION

Supplemental Information includes six figures and can be found with this article online at <https://doi.org/10.1016/j.molcel.2018.02.026>.

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## AUTHOR CONTRIBUTIONS

W.L. and A.M. prepared RNAP derivatives. W.L., K.D., Y.L., R.Y., Z.Z., E.T.E., and D.T. performed structure determination. D. Degen, R.Y.E., E.S., M.G., A.S., S.M., and Y.J. assessed RNAP-inhibitory activities, antibacterial activities, resistance, and cross-resistance. A.M., D. Duchi, D.W., and A.N.K. assessed FRET. Y.W.E., S.D., H.Z., and C.Z. prepared RNAP inhibitors. R.H.E. designed the study, analyzed data, and wrote the paper.

## DECLARATION OF INTERESTS

R.H.E. has patent filings on the Lpm target (US8206898 and US8697354). The other authors declare no competing interests.

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## STAR★METHODS

## KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Penta-His Biotin Conjugate	QIAGEN	Cat# 34440
Bacterial and Virus Strains		
<i>Actinoplanes deccanensis</i> ATCC 21983	Parenti et al., 1975	N/A
<i>Actinomadura</i> sp. DSMZ 13491	Ciciliato et al., 2004	N/A
<i>Dactylosporangium aurantiacum hamdenensis</i> NRRL 18085	ARS Culture Collection	NRRL 18085
<i>Escherichia coli</i> BL21(DE3)	EMD Millipore	Cat# C600003
<i>E. scherichia coli</i> D21f2tolC ( <i>rfa</i> tolC)	Fralick and Burns-Keliher, 1994	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -D516V	Zhang et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -H526D	Zhang et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -H526Y	Zhang et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -S531L	Zhang et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -E565D	Maffioli et al., 2017	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -E565G	Maffioli et al., 2017	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -D675A	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -N677K	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -N684T	Zhang et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -Y1251F	this paper	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -Q1256E	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoB</i> -Q1256L	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -R337H	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -R337S	this paper	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -D348Y	this paper	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -R738P	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -A779V	Degen et al., 2014	N/A
<i>Escherichia coli</i> D21f2tolC <i>rpoC</i> -G782A	Degen et al., 2014	N/A
<i>Escherichia coli</i> XE54	Tang et al., 1994	N/A
<i>Escherichia coli</i> XL1-Blue	Agilent / Stratagene	Cat# 200249
<i>Streptomyces</i> sp. ID38640	Maffioli et al., 2017	N/A
<i>Tth</i> HB-8	DSMZ	DSM 579
Chemicals, Peptides, and Recombinant Proteins		
lipiarmycin A3 (Lpm)	gift from A.L. Sonenschein	N/A
lipiarmycin A3 (Lpm)	this paper	N/A
lipiarmycin A3 (Lpm)	BioAustralis	Cat# BIA-F1356
lipiarmycin A3 (Lpm)	Biorbyt	Cat# orb146075
4"-O-benzyl-Lpm (Lpm analog "a1")	this paper	N/A
5"-deschloro-Lpm (Lpm analog "a2")	this paper	N/A
3,"5"-dideschloro-Lpm (50-1d; Lpm analog "a3")	this paper	N/A
5"-bromo-3,"5"-dideschloro Lpm (Lpm analog "a4")	this paper	N/A
69-1d (Lpm analog "a5")	Xiao et al., 2011 and Niu et al., 2011	N/A
70-9d (Lpm analog "a6")	Xiao et al., 2011 and Niu et al., 2011	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
70-10d (Lpm analog “a7”)	<a href="#">Xiao et al., 2011</a> and <a href="#">Niu et al., 2011</a>	N/A
70-11d (Lpm analog “a8”)	<a href="#">Xiao et al., 2011</a> and <a href="#">Niu et al., 2011</a>	N/A
OP-1118 (Lpm analog “b1”)	Toronto Research Chemical	N/A
53-1d (Lpm analog “b2”)	<a href="#">Xiao et al., 2011</a> and <a href="#">Niu et al., 2011</a>	N/A
57-4d (Lpm analog “c1”)	<a href="#">Xiao et al., 2011</a> and <a href="#">Niu et al., 2011</a>	N/A
desaryl-Lpm (Lpm analog “d1”)	this paper	N/A
54-3d (Lpm analog “e1”)	<a href="#">Xiao et al., 2011</a> and <a href="#">Niu et al., 2011</a>	N/A
rifampin (Rif)	Sigma-Aldrich	Cat# R3501
GE23077 (GE)	<a href="#">Ciciliato et al., 2004</a>	N/A
pseudouridimycin (PUM)	<a href="#">Maffioli et al., 2017</a>	N/A
salinamide A (Sal)	gift from W. Fenical	N/A
CBR703 (CBR)	Maybridge	Cat# RF05117SC
streptolydigin (Stl)	gift from E. Steinbrecher	N/A
myxopyronin B (Myx)	<a href="#">Ebright and Ebright, 2012</a>	N/A
corallopyronin A (Cor)	gift from H. Irshick and R. Jansen	N/A
ripostatin A (Rip)	gift from H. Irshick and R. Jansen	N/A
<i>Escherichia coli</i> RNA polymerase (RNAP) $\sigma^{70}$ holoenzyme	this paper	N/A
Alexa647, Cy3B-labeled <i>E. coli</i> RNA polymerase (RNAP) $\sigma^{70}$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> RNA polymerase (RNAP) $\sigma^A$ holoenzyme	<a href="#">Lin et al., 2017</a>	N/A
<i>Mycobacterium tuberculosis</i> [Ala1101] $\beta$ -RNA polymerase (RNAP) $\sigma^A$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala84] $\beta'$ -RNA polymerase (RNAP) $\sigma^A$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala89] $\beta'$ -RNA polymerase (RNAP) $\sigma^A$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala323] $\beta'$ -RNA polymerase (RNAP) $\sigma^A$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala412] $\beta'$ -RNA polymerase (RNAP) $\sigma^A$ holoenzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> RNA polymerase (RNAP) core enzyme	<a href="#">Lin et al., 2017</a>	N/A
<i>Mycobacterium tuberculosis</i> RNA polymerase (RNAP) core enzyme	<a href="#">Lin et al., 2017</a>	N/A
<i>Mycobacterium tuberculosis</i> [Ala1101] $\beta$ -RNA polymerase (RNAP) core enzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala84] $\beta'$ -RNA polymerase (RNAP) core enzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala89] $\beta'$ -RNA polymerase (RNAP) core enzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> [Ala323] $\beta'$ -RNA polymerase (RNAP) core enzyme	this paper	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
<i>Mycobacterium tuberculosis</i> [Ala412]β'-RNA polymerase (RNAP) core enzyme	this paper	N/A
<i>Mycobacterium tuberculosis</i> σ <sup>A</sup>	<a href="#">Lin et al., 2017</a>	N/A
<i>Staphylococcus aureus</i> RNA polymerase (RNAP) σ <sup>A</sup> holoenzyme	<a href="#">Maffioli et al., 2017</a>	N/A
<i>Staphylococcus aureus</i> RNA polymerase (RNAP) core enzyme	<a href="#">Maffioli et al., 2017</a>	N/A
<i>Staphylococcus aureus</i> σ <sup>A</sup>	<a href="#">Maffioli et al., 2017</a>	N/A
<i>Tth</i> RNA polymerase (RNAP) σ <sup>A</sup> holoenzyme	<a href="#">Zhang et al., 2012</a>	N/A
<i>Tth</i> RNA polymerase (RNAP) core enzyme	<a href="#">Zhang et al., 2012</a>	N/A
<i>Tth</i> σ <sup>A</sup>	<a href="#">Zhang et al., 2012</a>	N/A
HeLaScribe Nuclear Extract	Promega	Cat# E309A
γ-[2'-(2-benzothiazoyl)-60-hydroxybenzothiazole]-ATP	Jena Biosciences	Cat# NU-1700
fermentation growth media containing limiting chloride	<a href="#">Hochlowski et al., 1997</a>	N/A
H-top-agar	<a href="#">Sambrook and Russell, 2001</a>	N/A
LB broth	<a href="#">Sambrook and Russell, 2001</a>	N/A
LB agar	<a href="#">Sambrook and Russell, 2001</a>	N/A
ampicillin	Sigma-Aldrich	Cat# A0166
chloramphenicol	Sigma-Aldrich	Cat# C0378
kanamycin	Sigma-Aldrich	Cat# K4000
L-arabinose	Sigma-Aldrich	Cat# A3256
isopropyl-β-D-thiogalactoside (IPTG)	Gold Biotechnology	Cat# I2481C50
polyethyleneimine (Polymin P)	Sigma-Aldrich	Cat# P3143
4-azido-L-phenylalanine	Chem-Impex International	Cat# 06162
Alexa647-pentanoyl-ethylenediaminyl-phosphine	<a href="#">Chakraborty et al., 2010, 2015</a>	N/A
Cy3B-carboyl-ethylenediaminyl-phosphine	<a href="#">Chakraborty et al., 2010, 2015</a>	N/A
benzyl bromide	Sigma-Aldrich	Cat# B17905
n-octyl-β-D-glucopyranoside	Biosynth	Cat# O-2000
poly(acrylic acid sodium salt) 5,100 Da	Hampton Research	Cat# HR2-773
Vectabond aminosilane reagent	Vector Labs	Cat# SP-1800
methoxy-PEG succinimidyl valerate (mPEG-SVA)	Laysan Bio	Cat# MPEG-SVA-5000
biotinyl-PEG succinimidyl valerate (Biotin-PEG-SVA)	Laysan Bio	Cat# Biotin-PEG-SVA-5000
NeutrAvidin	ThermoFisher	Cat# 31000
Trolox	Sigma-Aldrich	Cat# 238813
glucose oxidase	Sigma-Aldrich	Cat# 345386
catalase	Sigma-Aldrich	Cat# C30
<b>Critical Commercial Assays</b>		
QuantiChromT Chloride Assay Kit	BioAssay Systems	Cat# DICL-250
HeLaScribe Nuclear Extract <i>in vitro</i> Transcription System	Promega	Cat# E3110
QuikChange site-directed mutagenesis kit	Agilent	Cat# 200519
QuikChange site-directed mutagenesis kit	Agilent	Cat# 200523
QuikChange site-directed mutagenesis kit	Agilent	Cat# 210518
Wizard Genomic DNA Purification Kit	Promega	Cat# A1125
Gel/PCR DNA Fragments Extraction Kit	IBI Scientific	Cat# IB47030
Wizard Classic 1 crystal screening solution kit	Emerald / Rigaku	Cat# 1009530
Wizard Classic 2 crystal screening solution kit	Emerald / Rigaku	Cat# 1009531
Wizard Classic 3 crystal screening solution kit	Emerald / Rigaku	Cat# 1009533
Wizard Classic 4 crystal screening solution kit	Emerald / Rigaku	Cat# 1009534

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Wizard Cryo 1 crystal screening solution kit	Emerald / Rigaku	Cat# 1009536
Wizard Cryo 2 crystal screening solution kit	Emerald / Rigaku	Cat# 1009537
Crystal Screen Cryo crystal screening solution kit	Hampton Research	Cat# HR2-122
Crystal Screen 2 Cryo crystal screening solution kit	Hampton Research	Cat# HR2-121
Crystal Screen crystal screening solution kit	Hampton Research	Cat# HR2-110
Crystal Screen 2 crystal screening solution kit	Hampton Research	Cat# HR2-112
Detergent Screen HT crystal screening solution kit	Hampton Research	Cat# HR2-406
Additive Screen HT crystal screening solution kit	Hampton Research	Cat# HR2-138
Index crystal screening solution kit	Hampton Research	Cat# HR2-144
Natrix crystal screening solution kit	Hampton Research	Cat# HR2-116
Natrix 2 crystal screening solution kit	Hampton Research	Cat# HR2-117
PEG/Ion crystal screening solution kit	Hampton Research	Cat# HR2-126
PEG/Ion 2 crystal screening solution kit	Hampton Research	Cat# HR2-098
PEGRx 1 crystal screening solution kit	Hampton Research	Cat# HR2-082
PEGRx 2 crystal screening solution kit	Hampton Research	Cat# HR2-084
SaltRx 1 crystal screening solution kit	Hampton Research	Cat# HR2-107
SaltRx 2 crystal screening solution kit	Hampton Research	Cat# HR2-109
NeXtal Protein Complex Suite crystal screening solution kit	QIAGEN	Cat# 130715
NeXtal Nucleix Suite crystal screening solution kit	QIAGEN	Cat# 130719
<b>Deposited Data</b>		
<i>Mycobacterium tuberculosis</i> RNA polymerase $\sigma^A$ holoenzyme in complex with lipiarmycin A3 (Lpm)	this paper	EMDB: 4230 PDB: 6FBV
<i>Tth</i> RNA polymerase core enzyme	this paper	PDB: 6ASG
<b>Recombinant DNA</b>		
pACYC- <i>rpoA</i>	<a href="#">Banerjee et al., 2014</a>	N/A
pACYC- <i>rpoA</i> -sigA	<a href="#">Banerjee et al., 2014</a>	N/A
pACYCDuet-Sau-H10-A	<a href="#">Maffioli et al. (2017)</a>	N/A
pCDF- <i>rpoZ</i>	<a href="#">Banerjee et al., 2014</a>	N/A
pCDFDuet-Sau-Z	<a href="#">Maffioli et al. (2017)</a>	N/A
pCOLA-Duet-1	Novagen / EMD Millipore	Cat# 71406
pCOLADuet-Mtb- <i>rpoB-rpoC</i>	this paper	N/A
pCOLADuet-Mtb- <i>rpoB</i> -K1101A- <i>rpoC</i>	this paper	N/A
pCOLADuet-Mtb- <i>rpoB-rpoC</i> -R84A	this paper	N/A
pCOLADuet-Mtb- <i>rpoB-rpoC</i> -R89A	this paper	N/A
pCOLADuet-Mtb- <i>rpoB-rpoC</i> -E323A	this paper	N/A
pCOLADuet-Mtb- <i>rpoB-rpoC</i> -R412A	this paper	N/A
pCOLADuet- <i>rpoB-rpoC</i>	<a href="#">Banerjee et al., 2014</a>	N/A
pCOLADuet-Sau-BC	<a href="#">Maffioli et al., 2017</a>	N/A
pET21a-Sau-H6-sigA	<a href="#">Maffioli et al., 2017</a>	N/A
pET28a-Tt- $\sigma^A$	<a href="#">Zhang et al., 2012</a>	N/A
pET30a-Mtb- $\sigma^A$	<a href="#">Jacques et al., 2006</a>	pSR01
pETDuet- <i>rpoB-rpoC</i>	<a href="#">Banerjee et al., 2014</a>	N/A
pEVOL-pAzF	<a href="#">Chin et al., 2002</a>	N/A
pJF09	<a href="#">Jacques et al., 2006</a>	N/A
pJF10	<a href="#">Jacques et al., 2006</a>	N/A
pREII-NH $\alpha$	<a href="#">Niu et al., 1996</a>	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
pRL663	<a href="#">Wang et al., 1995</a>	N/A
pRL663-K96I	this paper	N/A
pRL663-L249R	this paper	N/A
pRL663-S263T	this paper	N/A
pRL663-K334I	this paper	N/A
pRL663-R337H	this paper	N/A
pRL663-R337S	this paper	N/A
pRL663-N341S	this paper	N/A
pRL663-K345N	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-K345Q	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-K345R	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-K345T	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-D348Y	this paper	N/A
pRL663-L788M	<a href="#">Tuske et al., 2005</a>	N/A
pRL663-P1139R	<a href="#">Tuske et al., 2005</a>	N/A
pRL663-P1139L	<a href="#">Tuske et al., 2005</a>	N/A
pRL663-G1346D	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-V1351F	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-I1352N	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-I1352S	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL663-G1354C	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706	<a href="#">Severinov et al., 1997</a>	N/A
pRL706-A543V	<a href="#">Tuske et al., 2005</a>	N/A
pRL706-F545C	<a href="#">Tuske et al., 2005</a>	N/A
pRL706-P552M	<a href="#">Feng et al., 2015</a>	N/A
pRL706-Q618L	<a href="#">Feng et al., 2015</a>	N/A
pRL706-S642F	<a href="#">Feng et al., 2015</a>	N/A
pRL706-D654H	<a href="#">Feng et al., 2015</a>	N/A
pRL706-T657I	<a href="#">Feng et al., 2015</a>	N/A
pRL706-E892A	this paper	N/A
pRL706-M1232I	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-Y1251F	this paper	N/A
pRL706-T1255I	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-Q1256E	this paper	N/A
pRL706-Q1256L	this paper	N/A
pRL706-V1275F	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-V1275M	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-L1278V	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-E1279G	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-E1279K	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-A1283V	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-Y1285D	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-Q1288L	this paper	N/A
pRL706-L1291F	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-D1297A	this paper	N/A
pRL706-D1297Y	this paper	N/A
pRL706-V1298L	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-T1302P	this paper	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
pRL706-M1315L	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-P1317L	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-M1319I	this paper	N/A
pRL706-M1319K	this paper	N/A
pRL706-M1319R	this paper	N/A
pRL706-P1320A	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-E1321V	this paper	N/A
pRL706-S1322P	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-S1322T	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-S1322V	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-S1322Y	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRL706-N1324K	this paper	N/A
pRL706-V1325A	this paper	N/A
pRL706-V1325E	this paper	N/A
pRL706-V1325G	this paper	N/A
pRL706-V1325L	this paper	N/A
pRL706-L1326W	<a href="#">Mukhopadhyay et al., 2008</a>	N/A
pRSFduet-sigma	<a href="#">Hudson et al., 2009</a>	N/A
pVS10	<a href="#">Belogurov et al., 2007</a>	N/A
pVS10-rpoB106am;rpoC284am	this paper	N/A
<b>Software and Algorithms</b>		
Coot	<a href="#">Emsley et al., 2010</a>	<a href="https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/">https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/</a>
ebFRET	<a href="#">van de Meent et al., 2014</a>	<a href="http://ebfret.github.io/">http://ebfret.github.io/</a>
FSC	<a href="#">Henderson et al., 2012</a>	<a href="http://www.imagescience.de/fsc.html">http://www.imagescience.de/fsc.html</a>
Gctf	<a href="#">Zhang, 2016</a>	<a href="https://www.mrc-lmb.cam.ac.uk/kzhang/Gctf/">https://www.mrc-lmb.cam.ac.uk/kzhang/Gctf/</a>
HKL2000	<a href="#">Otwinowski and Minor, 1997</a>	<a href="http://www.hkl-xray.com/">http://www.hkl-xray.com/</a>
Leginon	<a href="#">Suloway et al., 2005</a>	<a href="http://www.legion.org">http://www.legion.org</a>
MotionCor2	<a href="#">Zheng et al., 2017</a>	<a href="http://msg.ucsf.edu/em/software/motioncor2.html">http://msg.ucsf.edu/em/software/motioncor2.html</a>
Molrep	<a href="#">Vagin and Teplyakov, 1997</a>	<a href="http://www.ccp4.ac.uk/html/molrep.html">http://www.ccp4.ac.uk/html/molrep.html</a>
MATLAB	Mathworks	<a href="https://www.mathworks.com/products/matlab.html">https://www.mathworks.com/products/matlab.html</a>
Origin	OriginLab	<a href="https://www.originlab.com/index.aspx?go=PRODUCTS/Origin">https://www.originlab.com/index.aspx?go=PRODUCTS/Origin</a>
Phenix	<a href="#">Adams et al., 2010</a>	<a href="https://www.phenix-online.org/">https://www.phenix-online.org/</a>
Relion v2.0.5	<a href="#">Fernandez-Leiro and Scheres, 2017</a>	<a href="https://www2.mrc-lmb.cam.ac.uk/relion/index.php?title=Main_Page">https://www2.mrc-lmb.cam.ac.uk/relion/index.php?title=Main_Page</a>
ResMap	<a href="#">Kucukelbir et al., 2014</a>	<a href="http://resmap.sourceforge.net/">http://resmap.sourceforge.net/</a>
Scipion v1.1	<a href="#">de la Rosa-Trevin et al., 2016</a>	<a href="https://github.com/I2PC/scipion/wiki">https://github.com/I2PC/scipion/wiki</a>
SGE Spiral Gradient Endpoint v1.3	Spiral Biotech	N/A
SigmaPlot	Systat Software	<a href="https://systatsoftware.com/products/sigmaplot/">https://systatsoftware.com/products/sigmaplot/</a>
TwoTone-ALEX	<a href="#">Holden et al., 2010</a>	<a href="https://groups.physics.ox.ac.uk/genemachines/group/Main.Software.html">https://groups.physics.ox.ac.uk/genemachines/group/Main.Software.html</a>
<b>Other</b>		
Titan Krios 300 keV cryo-electron microscope	FEI / ThermoFisher	N/A
K2 Summit direct electron detection camera	Gatan	N/A
GIF Quantum imaging filter	Gatan	N/A

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
PELCO easiGlow glow discharge cleaning system	Ted Pella	N/A
EM GP grid plunger	Leica	N/A
Ubuntu 16.04 Linux GPU workstation	Titan Computers	N/A
GeForce GTX 1080 Ti graphics card	Nvidia	N/A
Gryphon liquid handling system	Art Robbins Instruments	N/A
Total-internal-reflection fluorescence (TIRF) microscope	<a href="#">Holden et al., 2010</a> ; <a href="#">Duchi et al., 2016, 2018</a>	N/A
Samba 532 nm green laser	Cobolt	N/A
CUBE 635-30E red laser	Coherent	N/A
100x oil immersion objective (numerical aperture 1.4)	Olympus	N/A
1205C acousto-optical modulator	Isomet	N/A
Dichroic mirror (545 nm/650 nm)	Semrock	N/A
Dichroic mirror (630 nm DRLP)	Omega	N/A
545 nm LP emission filter	Chroma	Cat# HQ545LP
633/25 nm notch filter	Semrock	Cat#NF03-633E-25
iXon 897 EMCCD camera	Andor Technology	N/A
MS-2000 motorized x/y-scanning stage	ASI	N/A
Hitachi 2000 series HPLC system with L2455 detector	Hitachi	N/A
Hitachi 7000 series HPLC system with L7450 detector	Hitachi	N/A
ÄKTApurifier 10 FPLC system	GE Healthcare	Cat# 28406264
Luna C18 HPLC column (5 $\mu$ m; 100 Å; 250 $\times$ 4.6 mm)	Phenomenex	Cat# 00G-4041-E0
Supelco Discovery BIO Wide Pore C18 semi-prep HPLC column (10 $\mu$ m; 300 Å; 250 $\times$ 10 mm)	Sigma-Aldrich	Cat# 567209-U
HiLoad 16/60 Superdex 200 column	GE Healthcare	Cat# 17-1069-01
HiPrep Heparin FF 16/10 column	GE Healthcare	Cat# 28-9365-49
Mono Q 10/100 GL column	GE Healthcare	Cat# 17-5167-01
Mono S HR 10/10 column	GE Healthcare	Cat# 17-0557-01
SP Sepharose FF column	GE Healthcare	Cat# 17-0729-01
Silica gel, pore size 60 Å, 200-425 mesh particle size	Sigma-Aldrich	Cat# 236772
Ni-NTA agarose	QIAGEN	Cat# 30230
Amicon Ultra-15 30K centrifugal filters	EMD Millipore	Cat# UFC903024
EmuSiFlex-C5 cell disruptor	Avestin	N/A
Lambda 25 UV/Vis spectrophotometer	PerkinElmer	N/A
GENios Pro microplate reader	Tecan	N/A
GeneAmp 9700 PCR thermal cycler	Applied Biosystems	N/A
Autoplate 4000 spiral plater	Spiral Biotech	N/A
SGE transparent plastic template	Spiral Biotech	N/A
Lacey carbon film grids	Electron Microscopy Sciences	Cat#LC300-CU-100
Borosilicate glass coverslips #1.5	Menzel / ThermoFisher	Cat# 15654786
CultureWell 6 mm silicone gasket	Grace Bio-Labs	Cat# 103280
Whatman filter paper, Grade 1	Sigma-Aldrich	Cat# WHA1001150
Whatman filter paper, Grade 541	Sigma-Aldrich	Cat# WHA1541070

**CONTACT FOR REAGENT AND RESOURCE SHARING**

Requests for further information or reagents should be directed to Richard H. Ebright ([ebright@waksman.rutgers.edu](mailto:ebright@waksman.rutgers.edu)).

## METHOD DETAILS

### Lpm

Lpm was the kind gift of A.L. Sonenschein (Tufts University, Medford MA), was prepared by fermentation of *Actinoplanes deccanensis* ATCC 21983 (Parenti et al., 1975), as in Coronelli et al. (1975), or was purchased from BioAustralis or Biorbyt.

### Lpm analogs: previously described Lpm analogs

Lpm analog “a4” (3’-bromo-3,’5’-dideschloro-Lpm; compound 3 of Hochlowski et al., 1997) was prepared by fermentation of *Dactylosporangium aurantiacum hamdenensis* NRRL 18085 (Theriat et al., 1987; ARS Patent Culture Collection, Peoria IL) with precursor feeding, as in Hochlowski et al. (1997). Lpm analogs “a5” (69-1d), “a6” (70-9d), “a7” (70-10d), “a8” (70-11d), “b2” (53-11d), “c1” (54-4d), and “e1” (54-3d) were prepared by fermentation of mutants of *Dactylosporangium aurantiacum hamdenensis* NRRL 18085, as in Xiao et al. (2011) and Niu et al. (2011). Lpm analog “b1” (OP-1118; Babakhani et al., 2011) was purchased from Toronto Research Chemicals.

### Lpm analogs: Lpm analogs “a2” (5’-deschloro-Lpm) and “d1” (desaryl-Lpm)

Lpm analogs “a2” (5’-deschloro-Lpm) and “d1” (desaryl-Lpm) were prepared by fermentation of *Dactylosporangium aurantiacum hamdenensis* NRRL 18085 (Theriat et al., 1987; ARS Patent Culture Collection, Peoria IL) in growth media containing limiting chloride [growth media with KCl replaced by KBr; residual chloride concentration = 40  $\mu$ M, quantified using QuantiChromT Chloride Assay Kit (BioAssays Systems)]. First-stage (0.2 L) and second-stage (10 L) cultures were prepared as in Hochlowski et al. (1997), and second-stage culture broths were harvested as in Hochlowski et al. (1997). Culture broths were adjusted to pH 7 by addition of 1 mL 12 N NaOH, supplemented with 5 L acetone, shaken 1 h at 22°C, and extracted with 3x4 L ethyl acetate. The extracts were evaporated, and the resulting material (20 g) was re-dissolved in 300 mL ethyl acetate, filtered through Whatman filter paper (Grade 1; Sigma-Aldrich), and evaporated. The resulting material (14 g) was partitioned in 900 mL 1:1:1 (v/v/v) chloroform-methanol-water, and the lower phase was evaporated. The resulting material (11 g) was chromatographed on silica gel (Sigma-Aldrich; 3 cm x 30 cm; stepwise elution with 2%–50% methanol in chloroform, followed by elution with 100% methanol). UV-absorbant (254 nm) fractions were collected, evaporated, and assayed for antibacterial activity by spotting on H-top-agar/LB-agar (Sambrook and Russell, 2001) plates seeded with  $10^9$  cfu *E. coli* D21f2toIC (Fralick and Burns-Keliiher, 1994), incubating 16 h at 37, and assessing growth inhibition. Active fractions eluting at ~5% to ~10% methanol were pooled, further purified by re-chromatography on silica gel (procedures essentially as above), further purified by reversed-phase HPLC [Hitachi 7000 with L7450 detector; Supelco Discovery BIO Wide Pore C18 semi-prep column; 1:1 (v/v) acetonitrile-water isocratic elution; flow rate = 2 mL/min], and evaporated, yielding 80 mg Lpm analog “a3” [MALDI-MS m/z: calculated 1011.41 (M + Na<sup>+</sup>); found 1011.07] and 8 mg Lpm analog “a2” [MALDI-MS m/z: calculated 1045.59 (M + Na<sup>+</sup>); found 1044.60, 1046.70]. Material eluting at 100% methanol was further purified by reversed-phase HPLC (procedures essentially as above), and evaporated, yielding 0.2 mg Lpm analog “d1” [MALDI-MS m/z: calculated, 846.98 (M + Na<sup>+</sup>); found, 847.90].

### Lpm analogs: Lpm analog “a1” (4’-O-benzyl-Lpm)

Lpm analog “a1” (4’-O-benzyl-Lpm) was prepared from Lpm by semi-synthesis as follows: To Lpm (Biorbyt: 3 mg; 2.8  $\mu$ mol) in 50  $\mu$ L anhydrous dimethylformamide (Sigma-Aldrich), was added anhydrous potassium carbonate (Sigma-Aldrich: 3 mg, 22  $\mu$ mol), and the reaction mixture was stirred 30 min at 50°C and then allowed to cool to room temperature. An aliquot (10  $\mu$ L) of 0.28 M benzyl bromide in anhydrous dimethylformamide [2.8  $\mu$ mol; prepared by dissolving 3.4  $\mu$ L benzyl bromide (Sigma-Aldrich) in 50  $\mu$ L anhydrous dimethylformamide immediately before use] was added, and the reaction mixture was stirred 30 min at 50°C. The reaction mixture was evaporated to remove dimethylformamide and then re-suspended in 600  $\mu$ L 100 mM monobasic sodium phosphate (ThermoFisher). Precipitated solids were collected by centrifugation, rinsed with water, re-dissolved in methanol, and purified by reversed-phase HPLC (Hitachi 2000 with L2455 detector; Phenomenex Luna C18, 100 Å, 25 mm x 4.6 mm column; phase A = water; phase B = acetonitrile; gradient = 40% B at 0 min, 100% B at 20 min; flow rate = 1 mL/min). Lpm analog “a1” eluted at 20 minutes. Yield: 0.35 mg, 10%. MALDI-MS m/z: calculated, 1171.44 (M + Na<sup>+</sup>); found 1169.64, 1171.64.

### Other RNAP inhibitors

Cor A and Rip A were the kind gift of H. Irschik and R. Jansen (Helmholtz Institut, Braunschweig, Germany), StI was the kind gift of E. Steinbrecher (Upjohn-Pharmacia, Kalamazoo, MI), and Sal A was the kind gift of W. Fenical (The Scripps Research Institute, La Jolla, CA). Myx B was prepared by synthesis, as in Ebricht and Ebricht (2012); GE was prepared by fermentation of *Actinomadura* sp. DSMZ 13491, as in Ciciliato et al. (2004); and PUM was prepared by fermentation of *Streptomyces* sp. ID38640, as in Maffioli et al. (2017). Rif was purchased from Sigma-Aldrich, and CBR was purchased from Maybridge.

### *M. tuberculosis* RNAP $\sigma^A$ holoenzyme, RNAP core enzyme, and $\sigma^A$

*Mtb* RNAP  $\sigma^A$  holoenzyme was prepared by co-expression of genes for *Mtb* RNAP  $\beta'$  subunit, RNAP  $\beta$  subunit, N-terminally decahistidine-tagged RNAP  $\alpha$  subunit, RNAP  $\omega$  subunit, and N-terminally hexahistidine-tagged  $\sigma^A$  in *E. coli*, followed by cell lysis,

polyethylenimine precipitation, ammonium sulfate precipitation, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and anion-exchange chromatography on Mono Q (GE Healthcare), as in [Lin et al. \(2017\)](#).

For experiments in [Figures S1B, S3D, and S5](#), *Mtb* RNAP core enzyme was prepared by co-expression of genes for *Mtb* RNAP  $\beta'$  subunit, RNAP  $\beta$  subunit, N-terminally decahistidine-tagged RNAP  $\alpha$  subunit, and RNAP  $\omega$  subunit in *E. coli* [plasmids pCOLA-rpoB-rpoC, pACYC-rpoA, and pCDF-rpoZ ([Banerjee et al., 2014](#)); strain BL21(DE3) (EMD Millipore)], followed by cell lysis, polyethylenimine precipitation, ammonium sulfate precipitation, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and anion-exchange chromatography on Mono Q (GE Healthcare), as in [Lin et al. \(2017\)](#), but using plasmid pCOLADuet-Mtb-rpoB-rpoC [prepared by replacing the NcoI-BamHI segment of plasmid pCOLA-Duet (EMD Millipore) by the NcoI-BamHI segment of a DNA fragment carrying CCATGGTG followed by *Mtb* *rpoB* codons 3-1174 followed by TAAGGATCC, prepared by PCR using plasmid pJF09 ([Jacques et al., 2006](#); gift of S. Rodrigue, Université de Sherbrooke, Canada), and then replacing the NdeI-MfeI segment of the resulting plasmid by the NdeI-MfeI segment of a DNA fragment carrying CATATG followed by *Mtb* *rpoC* codons 2-1316 followed by TAGCAATTG, prepared by PCR using plasmid pJF10 ([Jacques et al., 2006](#); gift of S. Rodrigue, Université de Sherbrooke, Canada)], or derivatives thereof constructed using site-directed mutagenesis (QuikChange Site-Directed Mutagenesis Kit; Agilent), in place of plasmid pCOLADuet-rpoB-rpoC.

For experiments in [Figures S1B, S3D, and S5](#), *Mtb*  $\sigma^A$  was prepared by expression of a gene for N-terminally hexahistidine-tagged *Mtb*  $\sigma^A$  in *E. coli*, followed by cell lysis, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and anion-exchange chromatography on Mono Q (GE Healthcare), as in [Lin et al. \(2017\)](#).

### ***E. coli* RNAP $\sigma^{70}$ holoenzyme**

For experiments in [Figures S1B and S5](#), hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme was prepared from *E. coli* strain XE54 ([Tang et al., 1994](#)) transformed with plasmid pREII-NH $\alpha$  ([Tang et al., 1994](#)), using culture and procedures, cell lysis, polyethylenimine precipitation, ammonium sulfate precipitation, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and anion-exchange chromatography on Mono Q (GE Healthcare), as in [Degen et al. \(2014\)](#).

For experiments in [Figures 4C–4F](#), fluorescent-probe-labeled, hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme was prepared using unnatural-amino-acid mutagenesis ([Chin et al., 2002](#)) of co-expressed genes encoding RNAP  $\beta'$ ,  $\beta$ ,  $\alpha$ , and  $\omega$  subunits and  $\sigma^{70}$ , yielding an RNAP  $\sigma^{70}$  holoenzyme derivative containing 4-azido-L-phenylalanine (AzF) at position 284 of  $\beta'$  and position 106 of  $\beta$ , followed by azide-specific Staudinger ligation ([Saxon and Bertozzi, 2000](#)) to incorporate the fluorescent probes Cy3B and Alexa647 at position 284 of  $\beta'$  and position 106 of  $\beta$ , as follows ([Figure 4A](#)). Single colonies of *E. coli* strain BL21(DE3) (EMD Millipore) co-transformed with plasmid pVS10-rpoB106am;rpoC284am [constructed by use of site-directed mutagenesis (QuikChange Site-Directed Mutagenesis Kit; Agilent) to replace *rpoB* codon 106 and *rpoC* codon 284 by amber codons in plasmid pVS10 ([Belogurov et al., 2007](#))], plasmid pRSFduet-sigma ([Hudson et al., 2009](#)), and plasmid pEVOL-pAzF ([Chin et al., 2002](#)) were used to inoculate 20 mL LB broth ([Sambrook and Russell, 2001](#)) containing 100  $\mu$ g/mL ampicillin, 50  $\mu$ g/mL kanamycin, and 35  $\mu$ g/mL chloramphenicol, and cultures were incubated 16 h at 37°C with shaking. Culture aliquots (2x10 mL) were used to inoculate LB broth (2x1 L; [Sambrook and Russell, 2001](#)) containing 2 mM AzF (Chem-Impex International), 100  $\mu$ g/mL ampicillin, 50  $\mu$ g/mL kanamycin, and 35  $\mu$ g/mL chloramphenicol, cultures were incubated with shaking until OD<sub>600</sub> = 0.6; L-arabinose was added to 0.2% and IPTG was added to 1 mM, and cultures were further incubated 16 h at 16°C with shaking. Cells were harvested by centrifugation (4,000 x g; 20 min at 4°C), re-suspended in 20 mL buffer A (10 mM Tris-HCl, pH 7.9, 200 mM NaCl, and 5% glycerol), and lysed using an EmulsiFlex-C5 cell disrupter (Avestin). The lysate was cleared by centrifugation (20,000 x g; 30 min at 4°C), precipitated with polyethylenimine (Sigma-Aldrich) as in [Niu et al., 1996](#), and precipitated with ammonium sulfate as in [Niu et al., 1996](#). The sample was dissolved in 30 mL buffer A and loaded onto a 5 mL column of Ni-NTA-agarose (QIAGEN) pre-equilibrated in buffer A, and the column was washed with 50 mL buffer A containing 10 mM imidazole and eluted with 25 mL buffer A containing 200 mM imidazole. The sample was further purified by anion-exchange chromatography on Mono Q (GE Healthcare; 160 mL linear gradient of 300-500 mM NaCl in 10 mM Tris-HCl, pH 7.9, 0.1 mM EDTA, and 5% glycerol; flow rate = 2 mL/min). Fractions containing AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme were pooled, concentrated to ~1 mg/mL using 30 kDa MWCO Amicon Ultra-15 centrifugal ultrafilters (EMD Millipore), and stored in aliquots at -80°C. A reaction mixture containing 10  $\mu$ M AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme, 100  $\mu$ M Alexa647-pentanoyl-ethylenediaminyl-phosphine (prepared as in [Chakraborty et al., 2010, 2015](#)), and 100  $\mu$ M Cy3B-carboxyl-ethylenediaminyl-phosphine (prepared as in [Chakraborty et al., 2010, 2015](#)) in 1 mL buffer B (50 mM Tris-HCl, pH 7.9, 100 mM KCl, 5% glycerol, and 2% dimethylformamide) was incubated 1 h at 15°C, incubated 16 h on ice, and subjected to 5 cycles of buffer exchange (dilution with 5 mL buffer B, followed by concentration to 0.5 mL) using 30 kDa MWCO Amicon Ultra-15 centrifugal ultrafilters (EMD Millipore). The sample was further purified by gel-filtration chromatography on HiLoad 16/60 Superdex 200 prep grade (GE Healthcare) pre-equilibrated in buffer C (20 mM Tris-HCl, pH 8.0, 100 mM NaCl, 5 mM MgCl<sub>2</sub>, 1 mM  $\beta$ -mercaptoethanol, and 5% glycerol) and eluted in buffer C. Fractions containing fluorescent-probe-labeled hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme were pooled, concentrated to 1 mg/mL in buffer C using 30 kDa MWCO Amicon Ultra-15 centrifugal ultrafilters (EMD Millipore) and stored in aliquots at -80°C.

For experiments in [Figure 4G](#), fluorescent-probe-labeled, hexahistidine-tagged, FLAG-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme was prepared using unnatural-amino-acid mutagenesis ([Chin et al., 2002](#)) of genes encoding RNAP  $\beta'$  and  $\beta$  subunits to incorporate azidophenylalanine at position 284 of  $\beta'$  and position 106 of  $\beta$ , azide-specific Staudinger ligation ([Saxon and Bertozzi, 2000](#)) to incorporate Cy3B at

position 284 of the resulting  $\beta'$  derivative and Alexa647 at position 106 of the resulting  $\beta$  derivative, and *in vitro* reconstitution of RNAP (Tang et al., 1995) from the resulting  $\beta'$  and  $\beta$  derivatives, RNAP  $\alpha$  and  $\omega$  subunits, and  $\sigma^{70}$ , as in Chakraborty et al. (2010, 2012, 2015).

Efficiencies of incorporation of fluorescent probes were determined from UV/Vis-absorbance measurements and were calculated as:

$$\text{concentration of product} = [A_{280} - \epsilon_{\text{Cy3B},280} (A_{\text{Cy3B},559} / \epsilon_{\text{Cy3B},559}) - \epsilon_{\text{Alexa},280} (A_{\text{Alexa},652} / \epsilon_{\text{Alexa},647,652})] / \epsilon_{\text{P},280}$$

$$\text{Cy3B labelling efficiency} = 100\% [(A_{\text{Cy3B},559} / \epsilon_{\text{Cy3B},559}) / (\text{concentration of product})]$$

$$\text{Alexa647 labelling efficiency} = 100\% [(A_{\text{Alexa},652} / \epsilon_{\text{Alexa},652}) / (\text{concentration of product})]$$

where  $A_{280}$  is the measured absorbance at 280 nm,  $A_{\text{Cy3B},559}$  is the measured absorbance at the long-wavelength absorbance maximum of Cy3B (559 nm),  $A_{\text{Alexa},652}$  is the measured absorbance at the long-wavelength absorbance maximum of Alexa647 (652 nm),  $\epsilon_{\text{P},280}$  is the molar extinction coefficient of RNAP  $\sigma^{70}$  holoenzyme at 280 nm ( $240,000 \text{ M}^{-1} \text{ cm}^{-1}$ ),  $\epsilon_{\text{Cy3B},280}$  is the molar extinction coefficient of Cy3B at 280 nm ( $7,350 \text{ M}^{-1} \text{ cm}^{-1}$ ),  $\epsilon_{\text{Alexa},280}$  is the molar extinction coefficient of Alexa647 at 280 nm ( $10,400 \text{ M}^{-1} \text{ cm}^{-1}$ ), and  $\epsilon_{\text{Cy3B},559}$  is the extinction coefficient of Cy3B at its long-wavelength absorbance maximum ( $130,000 \text{ M}^{-1} \text{ cm}^{-1}$ ), and  $\epsilon_{\text{Alexa},652}$  is the extinction coefficient of Alexa647 at its long-wavelength absorbance maximum ( $245,000 \text{ M}^{-1} \text{ cm}^{-1}$ ). Labeling efficiencies were  $\sim 90\%$  for Cy3B and  $\sim 70\%$  for Alexa647.

Specificities of incorporation of fluorescent probes were determined from the observed labeling efficiencies of (i) the labeling reaction with the AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme and (ii) a control labeling reaction with non-AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme, and were calculated as:

$$\text{labelling specificity} = 100\% [1 - ((\text{labelling efficiency with P}) / (\text{labelling efficiency with AzF} - \text{P}))]$$

where AzF-P is AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme, and P is non-AzF-derivatized hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme. Labeling specificities were  $> 90\%$ .

### ***S. aureus* RNAP $\sigma^A$ holoenzyme, RNAP core enzyme, and $\sigma^A$**

*S. aureus* RNAP core enzyme was prepared by co-expression of genes for *S. aureus* RNAP  $\beta'$  subunit, RNAP  $\beta$  subunit, N-terminally decahistidine-tagged RNAP  $\alpha$  subunit, and RNAP  $\omega$  subunit in *E. coli*, followed by cell lysis, polyethylenimine precipitation, ammonium sulfate precipitation, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and cation-exchange chromatography on HiPrep Heparin (GE Healthcare), as in Maffioli et al., 2017.

*S. aureus*  $\sigma^A$  was prepared by expression of a gene for N-terminally hexahistidine-tagged *S. aureus*  $\sigma^A$  in *E. coli*, followed by cell lysis, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN), and gel-filtration chromatography on Superdex 200 (GE Healthcare), as in Maffioli et al. (2017).

*S. aureus* RNAP  $\sigma^A$  holoenzyme was prepared by combination of *S. aureus* RNAP core enzyme and *S. aureus*  $\sigma^A$ , as in Maffioli et al. (2017).

### ***Tth* RNAP $\sigma^A$ holoenzyme, RNAP core enzyme, and $\sigma^A$**

*Tth* RNAP core enzyme was prepared from *Tth* strain H8 (DSM 579; DSMZ), using cell lysis, polyethylenimine precipitation, ammonium sulfate precipitation, cation-exchange chromatography on SP Sepharose FF (GE Healthcare), anion-exchange chromatography on Mono Q 10/100 GL (GE Healthcare), and cation-exchange chromatography on Mono S HR (GE Healthcare), as in Zhang et al. (2012) and Maffioli et al. (2017).

*Tth*  $\sigma^A$  was prepared by co-expression of a gene for N-terminally hexahistidine-tagged *Tth*  $\sigma^A$  in *E. coli*, followed by cell lysis, immobilized-metal-ion affinity chromatography on Ni-NTA agarose (QIAGEN) and anion-exchange chromatography on Mono Q (GE Healthcare), as in Zhang et al. (2012).

*Tth* RNAP  $\sigma^A$  holoenzyme was prepared by combining *Tth* RNAP core enzyme and *Tth*  $\sigma^A$ , followed by size-exclusion chromatography on Superdex 200 (GE Healthcare), as in Zhang et al. (2014).

### **RNAP-inhibitory activities**

Fluorescence-detected RNAP-inhibition assays with the profluorescent substrate  $\gamma$ -[2'-(2-benzothiazoyl)-60-hydroxybenzothiazole]-ATP (BBT-ATP; Jena Bioscience) were performed as in Feng et al. (2015), using 75 nM RNAP holoenzyme [*Mtb* RNAP  $\sigma^A$  holoenzyme (prepared by pre-incubating 75 nM *Mtb* RNAP core enzyme or core enzyme derivative and 300 nM *Mtb*  $\sigma^A$  in transcription buffer 10 min at  $0^\circ\text{C}$ ), *S. aureus* RNAP  $\sigma^A$  holoenzyme, *E. coli* RNAP  $\sigma^{70}$  holoenzyme, or *Tth* RNAP  $\sigma^A$  holoenzyme] and 20 nM DNA fragment containing bacteriophage T5 N25 promoter (prepared as in Zhang et al., 2014). The transcription buffer for *E. coli* RNAP  $\sigma^{70}$  holoenzyme was 50 mM Tris-HCl (pH 8.0), 100 mM KCl, 10 mM  $\text{MgCl}_2$ , 1 mM DTT, 10 mg/mL bovine serum albumin, and 5.5%

glycerol. The transcription buffer for *Mtb* RNAP  $\sigma^A$  holoenzyme, *S. aureus* RNAP  $\sigma^A$  holoenzyme, and *Tth* RNAP  $\sigma^A$  holoenzyme was 40 mM Tris-HCl (pH 8.0), 75 mM NaCl, 5 mM MgCl<sub>2</sub>, 2.5 mM DTT, 5% DMSO, and 12.5% glycerol.

Radical RNAP-inhibition assays with HeLa nuclear extract (human RNAP I/II/III) were performed as in [Degen et al. \(2014\)](#). Half-maximal inhibitory concentrations (IC<sub>50</sub>s) were calculated by non-linear regression in SigmaPlot (Systat Software).

### Lpm-resistant mutants

Lpm-resistant mutants were isolated using procedures analogous to those used for isolation of Myx-resistant mutants in [Mukhopadhyay et al. \(2008\)](#). Mutagenesis reactions were performed using the QuikChange Site-Directed Mutagenesis Kit (Agilent) with *E. coli* *rpoC* plasmid pRL663 ([Wang et al., 1995](#)) and oligodeoxyribonucleotide forward and reverse primers corresponding to *rpoC* codons 1-20, 67-68, 77-81, 93-100, 245-256, 259-265, 325-355, 378-382, 393-403, 425-433, 466-481, 1319-1327, and 1347-1360, or with *E. coli* *rpoB* plasmid pRL706 ([Severinov et al., 1997](#)) and oligodeoxyribonucleotide forward and reverse primers corresponding to *rpoB* codons 854-857, 890-899, 914-922, and 1246-1342 (primers at 75 nM; all other components at concentrations as specified by the manufacturer). Mutagenized plasmid DNA was introduced by transformation into *E. coli* XL1-Blue (Agilent). Transformants (~10<sup>4</sup> cells) were applied to LB-agar ([Sambrook and Russell, 2001](#)) plates containing 200 μg/mL ampicillin, plates were incubated 16 h at 37°C, and plasmid DNA was prepared from the pooled resulting colonies. The resulting pooled mutagenized plasmid DNA was introduced by transformation into uptake-proficient, efflux-deficient *E. coli* strain D21f2toIC ([Fralick and Burns-Keliher, 1994](#)). Transformants (~10<sup>3</sup> cells) were applied to LB-agar plates containing 5 μg/mL Lpm (the minimal concentration that prevents colony formation by wild-type transformants), 200 μg/mL ampicillin, and 1 mM IPTG, and plates were incubated 24-48 h at 37°C. Lpm-resistant mutants were identified by the ability to form colonies on this medium, were confirmed by re-streaking on the same medium, and were demonstrated to contain plasmid-linked Lpm-resistant mutations by preparing plasmid DNA, transforming *E. coli* D21f2toIC with plasmid DNA, and plating transformants on the same medium. Nucleotide sequences of *rpoB* and *rpoC* were determined by Sanger sequencing (eight primers per gene).

### Resistance and cross-resistance levels

Experiments in [Figures 2C](#) and [S6B](#) and [S6C](#) assessing resistance and cross-resistance levels of Lpm-resistant *E. coli* D21f2toIC pRL706 and *E. coli* D21f2toIC pRL663 derivatives (preceding section) and Myx/Cor/Rip-resistant *E. coli* D21f2toIC pRL706 and *E. coli* D21f2toIC pRL663 derivatives ([Mukhopadhyay et al., 2008](#)) were performed using spiral gradient endpoint assays ([Wallace and Corkill, 1989](#); [Paton et al., 1990](#); [Schalkowsky, 1994](#)) on 150 mm x 4 mm exponential-gradient plates containing LB agar ([Sambrook and Russell, 2001](#)), 0.05-50 μg/mL test compound (Lpm, Rif, Myx, Cor, or Rip), 200 mg/mL ampicillin, and 1 mM IPTG. Test compounds were applied to plates using an Autoplate 4000 spiral plater (Spiral Biotech). Single colonies of transformants of *E. coli* D21f2toIC ([Fralick and Burns-Keliher, 1994](#)) were inoculated into 5 mL LB broth ([Sambrook and Russell, 2001](#)) containing 200 μg/mL ampicillin and incubated at 37°C with shaking until OD<sub>600</sub> = 0.4-0.6, IPTG was added to 1 mM, and cultures were incubated 1 h at 37°C with shaking. Diluted aliquots (~1x10<sup>9</sup> cfu/mL for [Figure 2C](#) and ~1x10<sup>8</sup> cfu/mL for [Figure S6B](#)) were swabbed radially onto plates, and plates were incubated 16 h at 37°C. For each culture, the streak length was measured using a clear plastic template (Spiral Biotech), the test-compound concentration at the streak endpoint was calculated using the program SGE: Spiral Gradient Endpoint (v1.3; Spiral Biotech), and the minimum inhibitory concentration (MIC) was defined as the test-compound concentration at the streak endpoint.

Experiments in [Figure 2D](#) assessing resistance and cross-resistance levels of chromosomal Lpm-resistant mutants [mutations transferred from pRL706 or pRL663 derivatives of preceding section to chromosome of *E. coli* D21f2toIC ([Fralick and Burns-Keliher, 1994](#)) by λ-Red-mediated recombinering (procedures essentially as in [Datsenko and Wanner, 2000](#), but using transformation rather than electroporation)] were performed using broth microdilution assays ([Clinical and Laboratory Standards Institute, 2009](#)). Single colonies were inoculated into 5 mL LB broth ([Sambrook and Russell, 2001](#)) and incubated at 37°C with shaking until OD<sub>600</sub> = 0.4-0.8. Diluted aliquots (~5x10<sup>4</sup> cells) in 97 μL LB broth were dispensed into wells of a 96-well plate, and were supplemented with 3 μL methanol or 3 μL of a 2-fold dilution series of Lpm (MIC<sub>wild-type</sub> = 1.56 μg/mL), Rif (MIC<sub>wild-type</sub> = 0.20 μg/mL), CBR (MIC<sub>wild-type</sub> = 6.25 μg/mL), Sal (MIC<sub>wild-type</sub> = 0.049 μg/mL), Stl (MIC<sub>wild-type</sub> = 3.13 μg/mL), or Myx (MIC<sub>wild-type</sub> = 0.20 μg/mL), in methanol (final concentrations = 0 and 0.006-50 μg/mL), or diluted aliquots (~1x10<sup>5</sup> cells) in 50 μL LB broth were supplemented with 50 μL LB broth or 50 μL of a 2-fold dilution series of GE (MIC<sub>wild-type</sub> = 500 μg/mL) or PUM (MIC<sub>wild-type</sub> = 400 μg/mL) in LB broth (final concentrations = 0 and 25-2000 μg/mL). Plates were incubated 16 h at 37°C. The MIC was defined as the lowest tested concentration that inhibited bacterial growth by ≥ 90%.

Experiments in [Figure 2E](#) assessing Lpm-cross-resistance levels of Rif-, GE/PUM-, and Sal-resistant mutants were performed as described for experiments in [Figure 2D](#), but analyzing a panel of *E. coli* D21f2toIC derivatives ([Degen et al., 2014](#); [Zhang et al., 2014](#); [Maffioli et al., 2017](#)) comprising 4 chromosomal Rif-resistant mutants, 3 chromosomal GE/PUM-resistant mutants, and 5 chromosomal Sal-resistant mutants.

Experiments in [Figure 2E](#) assessing Lpm-cross-resistance levels of CBR- and Stl-resistant mutants were performed essentially as described for experiments in [Figure 2D](#), but analyzing a panel of *E. coli* D21f2toIC pRL706 and *E. coli* D21f2toIC pRL663 derivatives ([Tuske et al., 2005](#); [Feng et al., 2015](#)) comprising 5 CBR-resistant mutants and 5 Stl-resistant mutants. Single colonies were inoculated into 5 mL LB broth ([Sambrook and Russell, 2001](#)) containing 200 μg/mL ampicillin and incubated at 37°C with shaking until OD<sub>600</sub> = 0.4-0.6, IPTG was added to 1 mM, and cultures were incubated 1 h at 37°C with shaking. Diluted aliquots (~5x10<sup>4</sup> cells)

in 97  $\mu$ L LB broth containing 200  $\mu$ g/mL ampicillin and 1 mM IPTG were dispensed into wells of a 96-well plate, and were supplemented with 3  $\mu$ L methanol or 3  $\mu$ L of a 2-fold dilution series of CBR or StI in methanol and further processed as described for experiments in Figure 2D.

### Checkerboard interaction assays

Antibacterial activities of combinations of Lpm and Rif were assessed in checkerboard interaction assays (White et al., 1996; Meletiadiis et al., 2010). Broth-microdilution assays (procedures as described in the preceding section for determination of resistance levels of chromosomal Lpm-resistant mutants) were performed in checkerboard format, using *E. coli* D21f2toC (Fralick and Burns-Keliher, 1994) and using LB broth (Sambrook and Russell, 2001) containing all pairwise combinations of: (i) Lpm at 2.0x, 1.75x, 1.5x, 1.3125x, 1.25x, 1.125x, 0.9375x, 0.75x, 0.5625x, 0.5x, 0.375x, 0.25x, and 0.1875x MIC<sub>Lpm</sub> and (ii) Rif at 2.0x, 1.75x, 1.5x, 1.3125x, 1.25x, 1.125x, 0.9375x, 0.75x, 0.5625x, 0.5x, 0.375x, 0.25x, and 0.1875x MIC<sub>Rif</sub>. Fractional inhibitory concentrations (FICs), FIC indices (FICIs), and minimum and maximum FICs (FIC<sub>min</sub> and FIC<sub>max</sub>) were calculated as in Meletiadiis et al. (2010). FIC<sub>min</sub>  $\leq$  0.5 was deemed indicative of super-additivity (synergism), FIC<sub>min</sub>  $>$  0.5 and FIC<sub>max</sub>  $\leq$  4.0 was deemed indicative of additivity, and FIC<sub>max</sub>  $>$  4.0 was deemed indicative of sub-additivity (antagonism) (White et al., 1996; Meletiadiis et al., 2010).

### Cryo-EM structure determination (*M. tuberculosis* RNAP-Lpm): sample preparation

Lacey carbon grids (LC300-CU-100; Electron Microscopy Sciences) were glow-discharged for 30 s using a glow-discharge cleaning system (PELCO easiGlow; Ted Pella) and mounted in the sample chamber of an EM GP grid plunger (Leica) at 18°C and relative humidity = 95%. Grids were spotted with 3.5  $\mu$ L 1  $\mu$ M *Mtb* RNAP-Lpm and 50  $\mu$ M Lpm in 20 mM Tris-HCl, pH 8.0, 75 mM NaCl, 5 mM MgCl<sub>2</sub>, 5 mM dithiothreitol, and 0.1% n-octyl- $\beta$ -D-glucopyranoside [prepared by pre-equilibrating 150  $\mu$ L samples containing components other than n-octyl- $\beta$ -D-glucopyranoside (Biosynth) 30 min at 25°C, and adding n-octyl- $\beta$ -D-glucopyranoside immediately before spotting], incubated 10 s, blotted with filter paper (Whatman Grade 541; Sigma-Aldrich) for 2.3 s, flash-frozen by plunging in liquid ethane cooled with liquid nitrogen, and stored in liquid nitrogen.

### Cryo-EM structure determination (*M. tuberculosis* RNAP-Lpm): data collection and data reduction

Data were collected at the National Resource for Automated Molecular Microscopy of the Simons Electron Microscopy Center using a 300 keV Titan Krios (FEI/ThermoFisher) electron microscope equipped with a K2 Summit direct electron detector (Gatan) operating in counting mode and a GIF Quantum imaging filter (Gatan) with slit width of 20 eV. Data were collected semi-automatically using the software package Legion (Suloway et al., 2005), a nominal magnification of 130,000x, a calibrated pixel size of 1.061 Å, and a dose rate of 8 electrons/pixel/s. Movies were recorded at 200 ms/frame for 10 s (50 frames total), resulting in a total radiation dose of 72.05 electrons/Å<sup>2</sup> per movie. Defocus range was varied between 1.0  $\mu$ m and 2.0  $\mu$ m. A total of 2,458 micrographs were recorded from two grids over 2 days.

Data were processed as summarized in Figures S1C–S1E. Data processing was performed on an Ubuntu 16.04 Linux GPU workstation (Titan Computers) containing four GeForce GTX 1080 Ti graphic cards (Nvidia; Kimanius et al., 2016). Frames in individual movies were aligned using MotionCor2 (Zheng et al., 2017), and the first 35 frames per movie were merged to calculate individual micrographs. Contrast-transfer-function estimations were performed using Gctf (Zhang, 2016), yielding defocus-range estimates ranging from 0.4  $\mu$ m to 4.0  $\mu$ m for individual micrographs. Initial particle picking was performed on 25 selected micrographs from the first grid (8,609 particles) using the Xmipp routine of the software package Scipion v1.1 (de la Rosa-Trevín et al., 2016), and particles were used for two-dimensional class averaging in Relion v2.0.5 (Fernandez-Leiro and Scheres, 2017). Eight distinct two-dimensional classes were selected as templates for picking 452,912 particles from 1,236 selected micrographs from the first grid and 366,594 particles from 1,025 selected micrographs from the second grid, using the Autopick routine of Relion. Two- and three-dimensional classifications were performed on the 452,912 particles from the first grid and 366,594 particles from the second grid, using Relion and using a 60 Å low-pass-filtered map calculated from the crystal structure of *Mtb* RPo (PDB: 5UH5; Lin et al., 2017; protein residues only) as the starting reference model for three-dimensional classification. Following identification and removal of heterogeneous particles in the two- and three-dimensional classifications, independent but similar density maps were obtained from 126,977 particles from the first grid and 97,412 particles from the second grid. Following merging of the 224,389 particles from the first and second grids, three-dimensional auto-refinement using Relion, and local angular sampling using Relion, a final density map was obtained from a subset of 68,895 particles. Gold-standard Fourier-shell-correlation analysis (FSC; Henderson et al., 2012) indicated a mean map resolution of 3.52 Å, and ResMap (Kucukelbir et al., 2014) indicated a median map resolution of 3.5 Å (Figures S1F and S1G; Table 1).

The initial atomic model for protein residues of *Mtb* RNAP-Lpm was built by manual rigid-body fitting of RNAP  $\beta'$ , RNAP  $\beta$ , RNAP  $\alpha^I$ , RNAP  $\alpha^{II}$ , RNAP  $\omega$ , and  $\sigma$  segments from the crystal structure of *Mtb* RPo (PDB: 5UH5; Lin et al., 2017; protein residues only) into the cryo-EM density map in Coot (Emsley et al., 2010), followed by adjustment of backbone and sidechain conformations in Coot. For  $\sigma$ R4 (residues 464–528), density was weak, suggesting high segmental flexibility;  $\sigma$ R4 was fitted as a rigid-body segment and was not further adjusted. For the RNAP  $\beta'$  N- and C-termini (residues 1–2 and 1282–1316), the central part of the RNAP  $\beta'$  trigger loop (residues 1014–1022), the RNAP  $\beta$  N- and C-termini (residues 1–27 and 1145–1172), the RNAP  $\alpha^I$  N-terminus and C-terminal domain (residues 1–2 and 227–347), the RNAP  $\alpha^{II}$  N-terminus and C-terminal domain (residues 1–2 and 233–347), the RNAP  $\omega$  N-terminus

(residues 1–27),  $\sigma$ R1.1 (residues 1–224), and a loop and short extended segment of the  $\sigma$ R3- $\sigma$ R4 linker (residues 426–433 and 443–445), density was absent, suggesting very high segmental flexibility; these segments were not fitted. The initial atomic model for Lpm atoms of *Mtb* RNAP-Lpm was built by manual rigid-body fitting of a crystal structure of Lpm (CCDC 114782; [Ihle et al., 2000](#)) into the cryo-EM density map using Coot, followed by torsion-angle adjustments using Coot. Iterative cycles were performed of real-space model building in Coot followed by reciprocal-space fitting to structure-factor amplitudes and phases calculated from the cryo-EM density map in Phenix ([Adams et al., 2010](#)). The final atomic model, with map-to-model correlation of 0.83, was deposited in the Electron Microscopy Data Bank (EMDB) and Protein Data Bank (PDB) with accession codes EMDB: 4230 and PDB: 6FBV ([Table 1](#)).

### Crystal structure determination (*Tth* RNAP core enzyme): sample preparation

Robotic crystallization trials were performed for *Tth* RNAP core enzyme using a Gryphon liquid handling system (Art Robbins Instruments), commercial screening solutions (Emerald Biosystems, Hampton Research, and QIAGEN), and the sitting-drop vapor-diffusion technique (drop: 0.2  $\mu$ L RNAP plus 0.2  $\mu$ L screening solution; reservoir: 60  $\mu$ L screening solution; 22°C). 900 conditions were screened. Rod-like crystals appeared under the identified crystallization conditions [0.1 M HEPES-NaOH, pH 7.5, 20 mM MgCl<sub>2</sub>, and 22% poly(acrylic acid sodium salt), 5,100 Da (Hampton Research); 22°C] within two weeks. Crystals were transferred from the sitting drop to a reservoir solution containing 20% (v/v) ethylene glycol (Sigma-Aldrich) and flash-cooled by immersing in liquid nitrogen.

### Crystal structure determination (*Tth* RNAP core enzyme): data collection and data reduction

Diffraction data and selenium single-wavelength anomalous dispersion data were collected from cryo-cooled crystals at Argonne Photon Source beamline 19ID-D. Data were processed using HKL2000 ([Otwinowski and Minor, 1997](#)). The resolution cut-off criteria were  $I/\sigma > 1.1$  and  $R_{\text{merge}} < 1$ .

The structure of *Tth* RNAP core was solved by molecular replacement with Molrep ([Vagin and Teplyakov, 1997](#)) using PDB: 4GZY ([Weixlbaumer et al., 2013](#)) as the search model. One RNAP molecule was present in the asymmetric unit. Early-stage refinement included rigid-body refinement of RNAP, followed by rigid-body refinement of RNAP subunits, followed by rigid-body refinement of 44 RNAP domains (methods as in [Zhang et al., 2012](#)). Cycles of iterative model building with Coot ([Emsley et al., 2010](#)) and refinement with Phenix ([Adams et al., 2010](#)) were performed. Improvement of the coordinate model resulted from improvement of phasing. The final model was generated by XYZ-coordinate refinement with secondary-structure restraints in Phenix, followed by group B-factor and individual B-factor refinement in Phenix. The final model, refined to  $R_{\text{work}}$  and  $R_{\text{free}}$  of 0.22 and 0.27, respectively, was deposited in the PDB with accession code PDB: 6ASG ([Table 2](#)).

### Single-molecule FRET: sample preparation

Observation wells were prepared essentially as described ([Duchi et al., 2016, 2018](#); [Figure 4B](#)): Borosilicate glass coverslips (#1.5; Menzel/ThermoFisher) were incubated in 40 mL acetone 5 min at 22°C, incubated in 40 mL 2% (v/v) Vectabond aminosilane reagent (Vector Labs) in acetone 5 min at 22°C, washed with 100 mL water at 22°C, dried under nitrogen, and bonded to CultureWell 6 mm silicone gaskets (GBL103280; Grace Bio-Labs), yielding 30  $\mu$ L wells containing aminosilane-functionalized glass floors. Aliquots (20  $\mu$ L) of 30 mM methoxy-PEG succinimidyl valerate (mPEG-SVA, MW 5,000; Laysan Bio) and 0.75 mM biotinyl-PEG succinimidyl valerate (Biotin-PEG-SVA, MW 5,000; Laysan Bio) in 50 mM MOPS-NaOH, pH 7.5, were added to wells and incubated 90 min at 22°C, supernatants were removed, and wells were washed with 5x200  $\mu$ L PBS (0.01 M sodium phosphate, pH. 7.4, 137 mM NaCl, and 2.7 mM KCl), yielding wells with biotin-PEG/mPEG-functionalized borosilicate glass floors. Aliquots (30  $\mu$ L) of 10  $\mu$ M NeutrAvidin (ThermoFisher) in 0.5xPBS then were added to wells and incubated 10 min at 22°C, supernatants were removed, and wells were washed with 3x100  $\mu$ L PBS, yielding wells with NeutrAvidin-biotin-PEG/PEG-functionalized glass floors. Aliquots (30  $\mu$ L) of 40 nM biotinylated anti-hexahistidine monoclonal antibody (Penta-His Biotin Conjugate; QIAGEN) in buffer KG7 (40 mM HEPES-NaOH, pH 7.0, 100 mM potassium glutamate, 10 mM MgCl<sub>2</sub>, 1 mM dithiothreitol, 100  $\mu$ g/mL bovine serum albumin, and 5% glycerol) then were added to wells and incubated 10 min at 22°C, supernatants were removed, and wells were washed with 3x100  $\mu$ L KG7, yielding wells with (biotinylated anti-hexahistidine monoclonal antibody)-NeutrAvidin-biotin-PEG/mPEG-functionalized glass floors.

For experiments in [Figures 4C–4F](#), fluorescent-probe-labeled, hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme was immobilized in observation wells containing (biotinylated anti-hexahistidine monoclonal antibody)-Neutravidin-biotin-PEG/mPEG-functionalized glass floors ([Figure 4B](#)) as follows: Aliquots (30  $\mu$ L) of 0.1 nM fluorescent-probe-labeled, hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme and 0 or 20  $\mu$ M (2.5xIC<sub>50</sub>) Lpm in KG7 (prepared by pre-equilibrating 50 nM fluorescent-probe-labeled, hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme and 0 or 20  $\mu$ M Lpm in KG7 10 min at 37°C, and then diluting 1:500 with 0 or 20  $\mu$ M Lpm in KG7 at 37°C) were added and incubated 2–4 min at 22°C, supernatants were removed, wells were washed with 2x30  $\mu$ L KG7, and 30  $\mu$ L KG7 containing 2 mM Trolox (Sigma-Aldrich) and an oxygen scavenging system [12.5  $\mu$ M glucose oxidase (Sigma-Aldrich), 16 nM catalase (C30; Sigma-Aldrich), and 8 mM D-glucose] at 22°C was added. Immobilization densities typically were ~30 molecules per 10  $\mu$ m x 12  $\mu$ m field of view. Immobilization specificities typically were > 98% (assessed in control experiments omitting biotinylated anti-hexahistidine monoclonal antibody).

For experiments in [Figure 4G](#), fluorescent-probe-labeled, hexahistidine-tagged *E. coli* RNAP  $\sigma^{70}$  holoenzyme was immobilized in observation wells containing (biotinylated anti-hexahistidine monoclonal antibody)-Neutravidin-biotin-PEG/mPEG-functionalized

glass floors in the absence of Lpm as described above, and 3  $\mu$ L 200  $\mu$ M Lpm in the same buffer was added during data acquisition [Lpm final concentration = 20  $\mu$ M (2.5 $\times$ I $C_{50}$ )].

### Single-molecule FRET: data collection and data analysis

FRET experiments were performed on an objective-type total-internal-reflection (TIRF) microscope (Holden et al., 2010; Duchi et al., 2016, 2018). Light from a green laser (532 nm; Samba; Cobolt) and a red laser (635 nm; CUBE 635-30E; Coherent) was combined using a dichroic mirror, coupled into a fiber-optic cable, focused onto the rear focal plane of a 100 $\times$  oil-immersion objective (numerical aperture 1.4; Olympus), and displaced off the optical axis such that the incident angle at the oil-glass interface of a stage-mounted observation chamber was greater than the critical angle, thereby creating an exponentially decaying evanescent wave. Alternating-laser excitation (ALEX; Kapanidis et al., 2004) was implemented by directly modulating the two lasers using an acousto-optical modulator (1205C; Isomet). Fluorescence emission was collected from the objective, separated from excitation light by a dichroic mirror (545 nm/650 nm; Semrock) and emission filters (545 nm LP, Chroma; and 633/25 nm notch filter, Semrock), focused on a slit to crop the image, and then spectrally separated using a dichroic mirror (630 nm DRLP; Omega) into donor and emission channels focused side-by-side onto an electron-multiplying charge-coupled device camera (EMCCD; iXon 897; Andor Technology). A motorized x/y-scanning stage with continuous reflective-interface feedback focus (MS-2000; ASI) was used to control the sample position relative to the objective.

For experiments in Figure 4C-F, laser powers were 3.5 mW (532 nm laser) and 0.7 mW (635 nm laser), and data were collected for 20 s using a frame rate of 1 frame per 20 ms. For experiments in Figure 4G, laser powers were 0.5 mW (532 nm laser) and 0.15 mW (635 nm laser), and data were collected for 50 s using a frame rate of 200 ms.

Fluorescence emission intensities in donor (green) and acceptor (red) emission channels were detected using the peak-finding algorithm of the MATLAB (MathWorks) software package TwoTone-ALEX (Holden et al., 2010), as in Holden et al. (2010). Peaks detected in both emission channels (i.e., peaks for molecules containing both donor and acceptor probes) and meeting ellipticity and distance-to-nearest-neighbor thresholds (i.e., ellipticity  $\leq$  0.6 and distance-to-nearest-neighbor  $\geq$  6 pixels) were fitted with two-dimensional Gaussian functions to extract background-corrected intensity-versus-time trajectories for donor emission intensity upon donor excitation ( $I_{DD}$ ), acceptor emission intensity upon donor excitation ( $I_{DA}$ ), and acceptor emission intensity upon acceptor excitation ( $I_{AA}$ ) (Figure 4C, top), as described (Holden et al., 2010). Intensity-versus-time trajectories were curated to exclude trajectories exhibiting  $I_{DD} < 300$  or  $> 2,000$  counts or  $I_{AA} < 200$  or  $> 2,000$  counts, trajectories exhibiting multiple-step donor or acceptor photobleaching, trajectories exhibiting donor or acceptor photobleaching in frames 1-50, and trajectories exhibiting donor or acceptor photoblinking, and to exclude portions of traces following donor or acceptor photobleaching.

Intensity-versus-time trajectories were used to calculate trajectories of apparent donor-acceptor FRET efficiency ( $E^*$ ) and donor-acceptor stoichiometry (S) (Figure 4C, bottom), as described (Kapanidis et al., 2004 and Lee et al., 2005):

$$E^* = I_{DA} / (I_{DD} + I_{DA})$$

$$S = (I_{DA} + I_{DD}) / (I_{DD} + I_{DA} + I_{AA})$$

$E^*$ -versus-S plots were prepared, S values were used to distinguish species containing only donor, only acceptor, and both donor and acceptor, and  $E^*$  histograms were prepared for species containing both donor and acceptor, as described (Kapanidis et al., 2004; Lee et al., 2005).

$E^*$ -versus time trajectories that on visual inspection exhibited transitions between distinct  $E^*$  states (dynamic  $E^*$ -versus-time trajectories;  $\sim$ 34%; N = 207) were analyzed globally to identify  $E^*$  states by use of Hidden Markov Modeling (HMM) with an empirical Bayesian as implemented in the MATLAB (MathWorks) software package ebFRET (van de Meent et al., 2014), essentially as described (van de Meent et al., 2014; Duchi et al., 2016, 2018).  $E^*$ -versus-time trajectories were fitted to HMM models with two, three, four, five, or six distinct  $E^*$  states; the mean scoring parameter L (lower bound per trajectory) was extracted for each model. and the model with three distinct  $E^*$  states was found to best describe the data (Figure 4D, right; L = 536, 538, 537, 537 and 537 for models with two, three, four, five, and six states).  $E^*$  values from the three-state HMM model were extracted, plotted using Origin (OriginLab), and fitted to Gaussian distributions using Origin (Figure 4D, left, colored curves). The resulting histograms provide equilibrium population distributions of  $E^*$  states and, for each  $E^*$  state, define mean  $E^*$  (Figure 4D, left, colored curves and inset).

Dwell times for  $E^*$  states were extracted from  $E^*$ -versus-time trajectories exhibiting  $> 3$  transitions between  $E^*$  states and were plotted as histograms in Origin. The resulting dwell-time histograms were fit with single-exponential functions, and mean dwell times were extracted (Figure 4F).

$E^*$  values were corrected, and accurate donor-acceptor efficiencies (E) and donor-acceptor distances (R) were calculated, as follows (Lee et al., 2005).

$$C_1 = E_{DO}^* / (1 - E_{DO}^*)$$

$$C_2 = S_{AO} / (1 - S_{AO})$$

$$E = [1 - C_2\{(1 - S)/S\} - C_1\{(1 - E^*)/E^*\}] / [\{(1 - E^*)/E^*\} + 1 - C_2\{(1 - S)/S\}] /$$

$$\times [\gamma - (\gamma - 1)[1 - C_2\{(1 - S)/S\} - C_1\{(1 - E^*)/E^*\}] / [\{(1 - E^*)/E^*\} + 1 - C_2\{(1 - S)/S\}]$$

$$R = R_0[(1/E) - 1]^{1/6}$$

where,  $C_1$  is “cross-talk” from leakage of donor emission into the acceptor-emission channel,  $C_2$  is “cross-talk” from direct excitation of the acceptor by the green laser,  $\gamma$  is the detection factor [1 in this work; determined as  $\gamma = \Delta I_{AA}/\Delta I_{DD}$ , where  $\Delta I_{AA}$  and  $\Delta I_{DD}$  are changes in  $I_{AA}$  and  $I_{DD}$  upon acceptor photobleaching (Ha et al., 1999)],  $E^*_{DO}$  is  $E^*$  of the donor-only subpopulation,  $S_{AO}$  is  $S$  of the acceptor-only subpopulation, and  $R_0$  is the Förster parameter [60.1 Å in this work; calculated as:  $R_0 = 9780(n^{-4}\kappa^2Q_DJ)^{1/6}$  Å, where  $n$  is the refractive index of the medium,  $\kappa^2$  is the orientation factor relating donor emission dipole and acceptor excitation dipole (approximated as 2/3, noting that all mean  $E$  values are  $< 0.5$ ; Wu and Brand, 1992),  $Q_D$  is the quantum yield of the donor in the absence of acceptor, and  $J$  is the overlap integral of donor emission spectrum and acceptor excitation spectrum].

RNAP clamp angles were determined from  $E$  values using the plot of RNAP clamp angle versus  $E$  in Figure 1 of Chakraborty et al. (2012).

### QUANTIFICATION AND STATISTICAL ANALYSIS

Data for RNAP-inhibitory activities, resistance, and cross-resistance are means of at least two technical replicates

### DATA AND SOFTWARE AVAILABILITY

The accession numbers for the cryo-EM map and atomic coordinates for the cryo-EM structure of *Mtb* RNAP-Lpm reported in this paper are EMDB: 4230 and PDB: 6FBV. The accession number for the atomic coordinates and structure factors for the crystal structure of *Tth* RNAP core enzyme reported in this paper is PDB: 6ASG.