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Structures of an RNA Polymerase Promoter Melting Intermediate Elucidate DNA Unwinding

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Abstract

A key regulated step of transcription in all cellular organisms is promoter melting by the RNA polymerase (RNAP) to form the open promoter complex^{1–3}. To generate the open complex, the conserved catalytic core of the RNAP combines with initiation factors to locate promoter DNA, unwind 12–14 base pairs (bps) of the DNA duplex, and load the template-strand (t-strand) DNA into the RNAP active site. Formation of the open complex is a multi-step process with transient intermediates of unknown structures^{4–6}. Here, we present cryo-electron microscopy structures of bacterial RNAP-promoter DNA complexes, including the first structures of partially melted intermediates. The structures unequivocally show that late steps of promoter melting occur within the RNAP cleft, delineate key roles for fork-loop 2 (FL2) and switch 2 (Sw2), universal structural features of RNAP, in restricting access of DNA to the RNAP active site, and explain why clamp opening is required to allow entry of single-stranded template DNA into the active site. The key roles of the FL2 and Sw2 suggest a common mechanism for late steps in promoter DNA opening to enable gene expression in all domains of life.

> Structures of promoter melting intermediates are essential to understand the mechanism of open promoter complex (RPo) formation. We took advantage of corallopyronin A (Cor; Extended Data Fig. 1A), an antibiotic closely related in structure and function to myxopyronin (Myx)^{7,8}, which was previously shown to inhibit RPo formation by trapping a partially melted intermediate⁷. A structure of bacterial RNAP with Cor is not available. As a target for Cor inhibition, we used *Mycobacterium tuberculosis (Mtb)* RNAP, which was inhibited by Cor in abortive initiation assays using the well-characterized Mtb rrnAP3 promoter as a template(Fig. 1A, Extended Data Fig. $1B$)⁹. To assemble the complex for structural studies, we incubated the components of the *Mtb* RNAP housekeeping initiation

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Author contributions H.B expressed and purified proteins, performed biochemical assays, and prepared cryo-EM grids. H.B. and J.C. collected and processed cryo-EM data. R.J. prepared and validated Cor. H.B, S.A.D., and E.A.C. built, refined, and validated the structures. S.A.D. and E.A.C. conceived the project. H.B., J.C., S.A.D., and E.A.C. prepared and revised the manuscript.

Data availability

The cryo-EM density maps have been deposited in the EMDataBank under accession codes EMD-9041 (Cor-RP2), EMD-9037 (RPo), EMD-9039 (RP2), and EMD-9047 (Cor-holo). The atomic coordinates have been deposited in the Protein Data Bank under accession codes 6EEC (Cor-RP2), 6EDT (RPo), 6EE8 (RP2), and 6M7J (Cor-holo).

complex (*Mtb* core RNAP and σ^A , along with essential *Mtb* general transcription factors RbpA and CarD ⁹ with Cor, then added duplex AP3 promoter template. After incubation, cryo-electron microscopy (cryo-EM) grids were prepared. A parallel sample was prepared without Cor.

Cryo-EM data from the sample with Cor yielded a single class at a nominal resolution of 3.6 Å, comprising the *Mtb* initiation complex (CarD-RbpA- σ ^A holoenzyme) bound to Cor (Extended Data Figs. 1C, 2) and also bound to the promoter DNA in a configuration distinct from RPo (Fig. 1B, Extended Data Table 1)¹⁰. The dataset without Cor was processed in a similar fashion and gave rise to two distinct classes (Fig. 1C, Extended Data Fig. 3A). The first class contained approximately 60% of the particles at a nominal resolution of 3.6 Å (Extended Data Fig. 3B–E) and comprised the Mtb initiation complex engaged with the promoter DNA melted into a complete 13 nucleotide transcription bubble (−11 to +2) with the t-strand DNA base at the transcription start site (TSS) (+1) near the RNAP active site Mg^{2+} (11 Å away) ready to template the incoming initiating NTP (Figs. 1C, 2A). The overall path of the DNA backbone matched previously determined RPo structures (Extended Data Fig. 4), hence we call this structure RPo. The second class contained approximately 40% of the particles at a nominal resolution of 3.9 Å (Extended Data Fig. 5) and comprised the *Mtb* initiation complex bound to the promoter DNA in a configuration very similar to the Cor-bound structure (Figs. 1B–D). We call this structure RP2 (and the Cor-bound structure Cor-RP2) for reasons explained below.

The cryo-EM maps revealed clear details of the interactions between RNAP and the DNA (Extended Data Fig. 6). In RPo, the promoter DNA from −12 to +3 (the transcription bubble from −11 to +2 plus the upstream and downstream double-strand/single-strand, ds/ss, junctions) interacts with conserved elements of the σ-factor as well as RNAP structural elements conserved in all kingdoms of life (Fig. 2A) 11 . We also observed protein/DNA interactions between the general transcription factors CarD and RbpA, both essential in mycobacteria but absent in E. coli (Eco)^{9,12} (Fig. 2A).

RP2 is a newly observed intermediate that contains a partially melted eight nucleotide bubble (−11 to −4) (Figs. 1B, 1C, 2B). While the upstream duplex portion of the promoter DNA in both the Cor-RP2 and RP2 structures is nearly identical to RPo (Fig. 1D), the protein/DNA interactions downstream differ substantially (Fig. 2). Belogurov et al.⁷ showed that Myx blocked bubble propagation downstream of −3, consistent with our Mtb RNAP RP2 structures. In the RP2 structures, the TSS $(+1)$ is base-paired with the nontemplate strand (nt-strand) and located more than 30 Å from the RNAP active site Mg^{2+} (Figs. 1B, 1C, 2B). The base-paired DNA in RP2 that is ultimately melted in RPo (−3 to +2) as well as the duplex DNA further downstream to about +12 is enclosed in the RNAP cleft (Figs. 1B, 1C), but the helical axis is tilted about 35° compared to the downstream duplex of RPo (Extended Data Fig. 7).

Clamp dynamics play an important role in promoter melting for all cellular RNAPs^{13–15}. In bacteria, early steps of promoter melting (transcription bubble nucleation) require clamp closure, while later steps (transcription bubble propagation to $+1$ and loading of the t-strand

DNA into the RNAP active site) require clamp opening¹³. Finally, clamp closure stabilizes RPo^{16} .

Like Myx, Cor closes the Eco RNAP clamp in solution, as shown by FRET¹⁶. We determined a cryo-EM structure of Mtb RNAP with Cor in the absence of downstream DNA (Cor-holo-us-fork) to a nominal resolution of 4.4 Å (Extended Data Figs. 8A–C) and compared it with a cryo-EM structure of holo without Cor $(6C05)^{17}$, confirming that Cor closes the Mtb RNAP clamp in solution as well (Extended Data Fig. 8D).

When the cryo-EM structures of promoter complexes (RPo, RP2, Cor-RP2) were superimposed (Extended Data Table 2), clamp conformational changes could be characterized as rigid body rotations about a common axis (Fig. 3A). Assigning a clamp rotation angle of 0° (closed clamp) to RPo, the Cor-RP2 and RP2 clamps are rotated open by 1.8° and 5.3°, respectively. By comparison, the clamp of an open clamp structure $(6BZO)^{17}$ is opened by 14°.

Analyses of the RP2 and Cor-RP2 structures delineate key roles for FL2 and Sw2, as well as clamp dynamics¹³, in the late stages of promoter melting. In the RP2 structures, the duplex DNA that needs to be melted to form RPo (from −3 to +2) is blocked from approaching the active site Mg^{2+} by interactions with FL2 and the narrow gap between FL2 and Sw2 due to the relatively closed clamp conformation of RP2 (Figs. 3A–C). The Cor-RP2 clamp presumably cannot open due to the bound Cor (Fig. 3B, Extended Data Fig. 8D) 8 , but opening of the RP2 clamp would break the FL2-duplex DNA interactions and widen the FL2-Sw2 gap from 15.4 Å to 20.3 Å (determined by measuring the minimal FL2-Sw2 α carbon to α-carbon distance; βE466 to β'K409), allowing passage of the ss t-strand DNA as it unwinds from the duplex by rotation of the downstream DNA^{13} . With the t-strand DNA in place, clamp closure would again restrict the FL2-Sw2 gap (Fig. 3D), helping to enforce the separation of the two DNA strands at the downstream ss/ds junction and stabilizing RPo (Fig. 3D). Single amino acid substitutions at multiple positions in or near FL2 destabilize RPo and reduce transcription from promoters limited by the RPo lifetime, consistent with a role for FL2 in RPo formation and/or dissociation^{18,19}.

Like Myx^7 , Cor refolds Sw2, resulting in a configuration that is sterically incompatible with the ss t-strand DNA in RPo (Fig. 3B). While Mukhopadhyay et al.⁸ proposed that Myx functions by interfering with clamp opening, Belogurov et al.⁷ proposed that the steric clash of the reconfigured Sw2 with the t-strand DNA explained Myx action. Our analysis suggests that both mechanisms likely contribute to Myx/Cor action.

Our finding that the partially melted state trapped by Cor (Fig. 1B) is very similar to a state observed in the absence of Cor (Fig. 1C) suggests that this structure corresponds to an 'onpathway' promoter melting intermediate. In RP2, the TSS is base-paired as a part of the duplex DNA enclosed within the RNAP cleft (Figs. 1C, 2B, 3C). In RPo, the start site is fully melted to template the incoming initiating NTP (Figs. 1B, 2A, 3D). Thus, our RP2 structures show that promoter melting occurs in multiple steps and that opening of the TSS occurs within the RNAP active site cleft.

Seminal work on the kinetics of RPo formation by Eco RNAP, as well as an analysis of RPo formation by mycobacterial RNAP on the AP3 promoter used here, established a minimal, three-step sequential mechanism^{$4-69$}. After formation of the initial encounter complex between RNAP and duplex promoter DNA (the closed complex, which we term RP1; Fig. 4), two major energetic barriers exist on the pathway to RPo formation, corresponding to the $RPI \rightarrow RP2$ and $RP2 \rightarrow RP0$ transitions (Fig. 4). Strikingly, consideration of the trajectory the DNA must traverse on its way from outside the active site cleft (in the RP1 model) to the final position near the RNAP active site in RPo reveals two major physical barriers. First, in the closed clamp RNAP conformation required for transcription bubble nucleation¹³, the gate loop (GL) of the RNAP β pincer²⁰ interacts with the clamp, sealing off DNA access to the RNAP cleft (Fig. 4). An RNAP conformational change, either opening of the βlobe/GL and/or clamp opening, is required to allow DNA access to a vestibule in the RNAP cleft between the GL and FL2, giving rise to RP2 (Fig. 4). As outlined above, clamp opening is then required to open the narrow FL2/Sw2 gap (Figs. 3C, 4), allowing the ss t-strand DNA to pass and position itself near the RNAP active site in RPo (Figs. 3D, 4). This second step, which corresponds to propagation of the transcription bubble from -3 to $+2$ and positioning of the ss t-strand into the RNAP active site, is the highest energy barrier (rate-limiting step) at the AP3 promoter⁹.

Thus, we propose that the two final major energetic barriers to RPo formation correspond to steric obstacles along the DNA trajectory that must be relieved by RNAP conformational changes. These barriers could serve as regulation checkpoints to control RNAP active site access. Indeed, the RP1 \leftrightarrow RP2 and RP2 \leftrightarrow RPo transitions were modulated by the mycobacterial transcription factors CarD and RbpA⁹, and the transition from RP1 \rightarrow RPo is a regulated step at some eukaryotic promoters 21 .

The RNAP structural features, namely FL2 and Sw2, highlighted in this study as key players in the final promoter melting transition, as well as clamp dynamics, are universally conserved features of all cellular $RNAPs^{11,13–15}$. In this regard, single molecule observations of the yeast RNAP II transcription initiation system support a two-step promoter melting process with an intermediate containing a 6 base-pair bubble²². We thus suggest that the RP2 intermediate observed here, and the RNAP clamp opening that allows the RP2 \rightarrow RPo transition, are common features of promoter melting and regulation for RNAPs from all three kingdoms of life.

METHODS

Structural biology software was accessed through the SBGrid consortium²³.

Protein Expression and Purification.

Mycobacterium tuberculosis RbpA/ σ^A -holoenzyme was purified and assembled as described previously¹⁷. *M. tuberculosis (Mtb)* CarD was overexpressed and purified as previously described for *Thermus thermophilus* CarD^{24} .

In vitro Transcription Assays.

In vitro abortive initiation transcription assays were performed using a wild-type Mtb AP3 promoter²⁵ template (−87 to +71) at 37°C as described previously^{17,26}. Assays were performed in KCl assay buffer (10 mM Tris-HCl, pH 8.0, 50 mM KCl, 10 mM $MgCl₂$, 0.1 mM EDTA, 0.1 mM DTT, 50 μg/mL BSA). CarD (250 nM) was added at 5X molar excess over RbpA/holoenzyme (50 nM) and incubated for 10 min at 37°C prior to the addition of corallopyronin (Cor) at the indicated concentrations (Extended Data Fig. 1B). Template DNA (10 nM) was added and the samples were incubated for 15 min at 37°C to allow RPo formation. Transcription was initiated by the addition of GpU initiating dinucleotide (250 $μ$ M, TriLink Biotechnologies) and $α-32P$ -UTP (15 nM). After 10 min., the reaction was quenched by the addition of $2X$ stop buffer (8 M urea, $0.5X$ TBE, 0.05% bromophenol blue, 0.05% xylene cyanol), and transcription products were visualized by polyacrylamide gel electrophoresis using phosphorimagery and quantified using Image J^{27} . The transcription assay (Extended Data Fig. 1B) was performed once to calculate the concentration of Cor to use to obtain the structure. Since the experiment was performed with two batches of enzyme (one prepared with and one prepared without the transcription CarD, which would not affect the inhibition) with a range of concentrations and showed similar IC50 (1 and 2 μ M) we considered this duplicated and cautiously used 100 times above the IC50 (100 μM) for cryo-EM sample preparation.

Preparation of Mtb RNAP/σ **^A/CarD/RbpA/Cor Complex for Cryo-EM.**

 Mtb RNAP/ σ^A /RbpA (0.5 ml of 5 mg/ml) was injected into a Superose 6 Increase column (GE Healthcare) equilibrated with 20 mM Tris-HCl, pH 8.0, 150 mM K-Glutamate, 5 mM $MgCl₂$, and 2.5 mM DTT. The peak fractions of the eluted protein were concentrated by centrifugal filtration (Millipore) to 6 mg/mL protein concentration. CarD was added to 5X molar excess and incubated for 37°C for 10 min. Cor (10 mM stock solution in DMSO) was added to a final concentration of 0.1 mM [final 1% (v/v) DMSO], and DMSO to 1% (v/v) was added to the sample with no Cor, then incubated for 10 min at 37°C. Next, duplex AP3 promoter fragment (-60 to +30, Fig. 1A) was added to a final concentration of 20 μ M and the sample incubated for 15 min at 37°C. CHAPSO (3-([3 cholamidopropyl]dimethylammonio)-2-hydroxy-1-propanesulfonate) was then added to a final concentration of 8 mM and sample was kept at room temperature prior to grid preparation.

Cryo-EM grid preparation.

C-flat holey carbon grids (CF-1.2/1.3–4Au) were glow-discharged for 20 s prior to the application of 3.5 μl of the sample (4.0– 6.0 mg/ml protein concentration). After blotting for (3–4.5 s) the grids were plunge-frozen in liquid ethane using an FEI Vitrobot Mark IV (FEI, Hillsboro, OR) with 100% chamber humidity at 22 °C.

Cryo-EM data acquisition and processing

Mtb RNAP/σ **^A/CarD/RbpA/AP3.—**The grids were imaged using a 300 keV Titan Krios (FEI) equipped with a K2 Summit direct electron detector (Gatan). Images were recorded with Leginon²⁸ in counting mode with a pixel size of 1.07 Å and a defocus range of 0.8 μ m

to 1.8 μm. Data were collected with a dose of 8 electrons/px/s. Images were recorded over a 10 second exposure with 0.2 second frames (50 total frames) to give a total dose of 70 electrons/ \AA^2 . Dose-fractionated subframes were aligned and summed using MotionCor2²⁹ and subsequent dose-weighting was applied to each image. The contrast transfer function was estimated for each summed image using Gct^{30} . From the summed images, Gautomatch (developed by K. Zhang, MRC Laboratory of Molecular Biology, Cambridge, UK, [http://](http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch) [www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch\)](http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch) was used to pick particles with an autogenerated template. A subset of the dataset was used to generate an initial model of the complex in cryoSPARC (*ab initio* reconstruction)³¹. Many classification schemes were tested that converged on the conclusion that two major, high-resolution classes were present in the particle dataset. For the final structures, the *ab initio* model (low-pass filtered to 30 \AA resolution) was used as a template to 3D classify the particles into two classes using cryoSPARC heterogeneous refinement. Two more rounds of classification (into 2 classes) were performed to remove bad particles (low resolution reconstructions that did not resemble RNAP). Then, cryoSPARC homogenous refinement was performed for each resulting class using the class map and corresponding particles, yielding two different structures: RPo and RP2 (Extended Data Fig. 3A). The RPoclass contained 211,381 particles with a nominal resolution of 3.6 Å (Extended Data Fig. 3) while the RP2 class contained 140,333 particles with a nominal resolution of 3.9 Å (Extended Data Figs. 3A, 5).

Mtb RNAP/σ **^A/CarD/RbpA/CarD/AP3/Cor.—**The grids were imaged using a 300 keV Titan Krios (FEI) equipped with a K2 Summit direct electron detector (Gatan). Images were recorded with Serial EM^{32} in super-resolution counting mode with a super-resolution pixel size of 0.65 Å and a defocus range of 0.8 μ m to 2.0 μ m. Data were collected with a dose of 8 electrons/px/s. Images were recorded over a 15 second exposure using 0.3 second subframes (50 total frames) to give a total dose of 71 electrons/ \AA ². Dose-fractionated subframes were 2 \times 2 binned (giving a pixel size of 1.3 Å), aligned and summed using MotionCor2²⁹. The contrast transfer function was estimated for each summed image using $Gctf³⁰$. From the summed images, Gautomatch (developed by K. Zhang, MRC Laboratory of Molecular Biology, Cambridge, UK,<http://www.mrc-lmb.cam.ac.uk/kzhang/Gautomatch>) was used to pick particles with an auto-generated template. Autopicked particles were manually inspected. A subset of the dataset was used to generate an initial model of the complex in cryoSPARC (*ab initio* reconstruction)³¹. Many classification schemes were tested that converged on the conclusion that one major, high-resolution class was present in the particle dataset. For the final structure, the *ab initio* model (low-pass filtered to 30 Å-resolution) was used as a template to 3D classify the particles into two classes using cryoSPARC heterogeneous refinement. Two more rounds of classification (into 2 classes) were performed to remove bad particles (low resolution reconstructions that did not resemble RNAP). Then, cryoSPARC homogenous refinement was performed for the best class using the class map and corresponding particles, yielding the Cor-RP2 structure containing 246,409 particles with a nominal resolution of 3.6 Å (Extended Data Figs. 1C, 2).

Mtb RNAP/σ^A/RbpA/us-fork DNA/Cor.—Grids were imaged on two separate 300 keV Krios microscopes (FEI), both equipped with K2 Summit direct electron detectors (Gatan). Two datasets were recorded with Serial EM^{32} in super-resolution mode over a defocus range

of 0.8 μm to 2.5 μm (Extended Data Table 1). The first dataset was collected at 8 electrons/ physical pixel/second with a super-resolution pixel size of 0.65 Å. Images in the first dataset were recorded in dose-fractionation mode with subframes of 0.3 s over a 15 s exposure (50 frames) to give a total dose of 71 electrons/ \AA^2 . Dose-fractionated movies were gainnormalized, Fourier binned by 2 (giving a pixel size of 1.3 Å), drifted-corrected, summed, and dose-weighted using MotionCor229. The second dataset was collected at 5 electrons/ physical pixel/s with a superresolution pixel size of 0.515 Å. Images in the second dataset were recorded in dose-fractionation mode with subframes of 0.3 s over an exposure of 15 s (50 total frames) to give a total dose of 71 electrons/ \AA ². Dose-fractionated movies were gain-normalized, scaled to the pixel size of the first dataset using a Fourier binning factor of 2.52 (giving a pixel size of 1.30 Å), drifted-corrected, summed, and dose-weighted using MotionCor 2^{29} . CTF estimations were calculated for each dataset using Gctf 30 . Particles were picked using Gautomatch (K. Zhang, [http://www.mrc-lmb.cam.ac.uk/kzhang/\)](http://www.mrc-lmb.cam.ac.uk/kzhang/) without using a 2D template. Picked particles were extracted from the dose-weighted images in RELION³⁴ using a box size of 256 pixels. The first dataset consisting of 2,059 images with 504,577 particles was combined with the second dataset consisting of 2,839 images with 420,432 particles. Particles were subjected to multiple rounds of cryoSPARC³¹ 3D classifications using a cryo-EM map of Mtb RNAP/ σ^A /RbpA (EMD-7322) as a 3D template (Extended Data Fig. 8A). The best class consisted of 222,962 particles with a nominal resolution of 4.38 Å (Extended Data Figs. 8B–C) after homogenous refinement in cryoSPARC³¹.

The distribution of particle orientations for each class was plotted using cryoSPARC (Extended Data Figs. 2A, 3B, 5A, 8B). FSC calculations were performed in cryoSPARC (Extended Data Figs. 2B, 3C, 5B, 8C) and half-map FSCs (Extended Data Figs. 2C, 3D, 5C) were calculated using EMAN235. Local resolution calculations (Extended Data Figs. 2D, $3E$, 5D) were performed using blocres³⁶.

Model building and refinement.

To build initial models of the protein components of the complexes, a model of *Mtb* RNAP σ ^A/RbpA/us-double-fork structure with the DNA removed (PDB ID 6C04)17 was manually fit into the cryo-EM density maps using Chimera³⁷ and real-space refined using Phenix³⁸. The DNAs were mostly built *de novo* based on the density map. For real-space refinement, rigid body refinement with fourteen manually-defined mobile domains was followed by allatom and B-factor refinement with Ramachandran and secondary structure restraints. A model of Cor was generated from a SMILES string and edited in Phenix REEL, and refined into the cryo-EM density. Refined models were inspected and modified in Coot³⁹.

Generation of cartoons in figure 4.

Cartoons were drawn by superimposition unto the pdbs of the RP1 model⁹, and Rp2 (6EE8) and RPo (6EDT) structures. In the side views, the main body of the RNAP has been cut away at the level of the RNAP active site Mg^{2+} (yellow sphere) but the full clamp (pink) and nucleic acids are shown. In the cross-sections (viewed from the top), most of the RNAP β subunit has been removed (except for FL2 and the GL in green) to reveal the inside of the RNAP active site cleft.

Extended Data Figure 1 |.

Cor inhibits Mtb RNAP transcription initiation but not promoter DNA binding and data processing pipeline for the cryo-EM movies of Mtb RNAP/σ^A/CarD/RbpA/Cor/AP3 promoter

A. Chemical structure of corallopyronin A⁴⁰

B. Abortive transcription initiation assays measuring GpUpU production in the presence of increasing concentrations of Cor. $\lceil 32P \rceil$ -labeled abortive transcript production was monitored by polyacrylamide gel electrophoresis and autoradiography. Full gel is shown in supplementary Fig. 1.

C. Flowchart showing the image processing pipeline for the cryo-EM data of Mtb RNAP/ σ ^A/CarD/RbpA/Cor/AP3 promoter complexes starting with 3,718 dose-fractionated movies collected on a 300 keV Titan Krios (FEI) equipped with a K2 Summit direct electron detector (Gatan). Movies were frame aligned and summed using MotionCor2²⁹. CTF estimation for each micrograph was calculated with $Gct³⁰$. A representative micrograph is shown following processing by MotionCor2. Particles were autopicked from each micrograph with Gautomatch and then sorted by 2D classification using RELION³⁴ to assess quality. The twelve highest populated classes from the 2D classification are shown. After picking, the dataset contained 1,026,386 particles. A subset of particles was used to generate an *ab-initio* map in cryoSPARC³¹. Using the low-pass filtered (30 Å) *ab initio* map as a template, three rounds of 3D heterogeneous refinement were performed using cryoSPARC in a binomial-like fashion. One major, high-resolution class emerged, which was refined using cryoSPARC homogenous refinement and then sharpened for model building.

Extended Data Figure 2 |.

Cryo-EM of Cor-RP2.

A. Angular distribution calculated in cryoSPARC for particle projections. Heat map shows number of particles for each viewing angle (less=blue, more=red).

B. Gold-standard FSC⁴¹, calculated by comparing the two independently determined halfmaps from cryoSPARC. The dotted line represents the 0.143 FSC cutoff which indicates a nominal resolution of 3.6 Å.

C. FSC calculated between the refined structure and the half map used for refinement (work, red), the other half map (free, blue), and the full map (black).

D. (top) The 3.6 Å-resolution cryo-EM density map of Cor-RP2 is colored according to the key on the right. The right view is a cross-section of the left view revealing the DNA inside the RNAP cleft.

(bottom) Same views as (top) but colored by local resolution³⁶. The boxed region in the right view is magnified on the far right and sliced at the level of the Cor binding pocket. Density for the Cor molecule is outlined in red.

E. (left) Overview of the Cor-RP2 structure, shown as a molecular surface. The boxed region is magnified on the right.

(right) Magnified view of the Cor binding pocket at the same orientation as the boxed region on the left. Proteins are shown as α-carbon backbone worms. Residues that interact with Cor are shown in stick format. Cor is shown in stick format with green carbon atoms. Hydrogenbonds are indicated by dashed gray lines. The cryo-EM difference density for the Cor is shown (green mesh).

Extended Data Figure 3 |.

Data processing pipeline for the cryo-EM movies of Mtb RNAP/σ^A/CarD/RbpA/AP3 promoter (RPo, RP2) and cryo-EM of RPo.

A. Flowchart showing the image processing pipeline for the cryo-EM data of Mtb RNAP/ σ ^A/CarD/RbpA/AP3 promoter complexes starting with 8,577 dose-fractionated movies collected on a 300 keV Titan Krios (FEI) equipped with a K2 Summit direct electron detector (Gatan). Movies were frame aligned and summed using MotionCor2²⁹. CTF estimation for each micrograph was calculated with $Gct³⁰$. A representative micrograph is

shown following processing by MotionCor2. Particles were auto picked from each micrograph with Gautomatch and then sorted by 2D classification using $RELION³⁴$ to assess quality. The twelve highest populated classes from the 2D classification are shown. The dataset contained 931,461 particles. A subset of particles was used to generate an ab*initio* map in cryoSPARC³¹. Using the low-pass filtered (30 Å) *ab initio* map as a template, two rounds of 3D heterogeneous refinement were performed using cryoSPARC in a binomial-like fashion. Two major classes emerged, which were refined using cryoSPARC homogenous refinement and then sharpened for model building.

B. Angular distribution calculated in cryoSPARC for particle projections. Heat map shows number of particles for each viewing angle (less=blue, more=red).

C. Gold-standard FSC^{41} , calculated by comparing the two independently determined halfmaps from cryoSPARC. The dotted line represents the 0.143 FSC cutoff which indicates a nominal resolution of 3.6 Å.

D. FSC calculated between the refined structure and the half map used for refinement (work, red), the other half map (free, blue), and the full map (black).

E. (top) The 3.6 Å-resolution cryo-EM density map of RPo is colored according to the key on the right. The right view is a cross-section of the left view revealing the DNA inside the RNAP cleft.

(bottom) Same views as (top) but colored by local resolution³⁶.

Extended Data Figure 4 |.

Overall DNA path of RPo matches previously determined RPo structures. Selected RPo structures containing a completely intact transcription bubble $42,43$ were superimposed with the cryo-EM RPo structure by α-carbons of the structural core module

(Extended Data Table 2). The resulting superposition of the nucleic acids is shown. The nucleic acids are shown as phosphate backbone worms, color-coded as shown in the key.

Extended Data Figure 5 |.

Cryo-EM of RP2 class.

(**A**) Angular distribution calculated in cryoSPARC for particle projections. Heat map shows number of particles for each viewing angle (less=blue, more=red).

 (B) Gold-standard FSC⁴¹, calculated by comparing the two independently determined halfmaps from cryoSPARC. The dotted line represents the 0.143 FSC cutoff which indicates a nominal resolution of 3.9 Å.

(**C**) FSC calculated between the refined structure and the half map used for refinement (work, red), the other half map (free, blue), and the full map (black).

(**D**) (top) The 3.9 Å-resolution cryo-EM density map of RP2 is colored according to the key on the right. The right view is a cross-section of the left view revealing the DNA inside the RNAP cleft.

(bottom) Same views as (top) but colored by local resolution³⁶.

Extended Data Figure 6 |.

Sample cryo-EM density.

Stereo views of cryo-EM density (blue mesh) and superimposed models.

A. Protein/DNA interactions for t-strand DNA of RPo.

B. Downstream ss/ds fork junction of RPo. The t-strand +1 position (transcription start site, T+1, colored lemon-green) is unpaired and positioned near the RNAP active site Mg^{2+} (not visible in this view).

C. Same view as (B) but showing cryo-EM density and model for RP2. The T+1 nucleotide (lemon-green) is base paired with the nt-strand and more than 30 Å away from the RNAP active site Mg^{2+} .

Extended Data Figure 7 |.

Downstream duplex DNA helical axes.

The RPo, RP2, and Cor-RP2 structures were superimposed and the nucleic acid backbones are shown (RPo, blue; RP2, green; Cor-RP2, magenta). The helical axes of the downstream duplex DNAs, determined using curves⁴⁴, are shown as thick colored lines. The RP2 and Cor-RP2 downstream duplexes are tilted about 35° with respect to RPo.

Extended Data Figure 8 |.

Cryo-EM of *Mtb* RNAP/σ^A/RbpA/Cor/us-fork.

A. Flowchart showing the image processing pipeline for the cryo-EM data of Mtb RNAP/ σ ^A/RbpA/Cor/us-fork complexes starting with 4,897 dose-fractionated movies collected on a 300 keV Titan Krios (FEI) equipped with a K2 Summit direct electron detector (Gatan). Movies were frame aligned and summed using MotionCor2²⁹. CTF estimation for each micrograph was calculated with Gctf³⁰. A representative micrograph is shown following processing by MotionCor2. Particles were autopicked from each micrograph with

Gautomatch and then sorted by 2D classification using $RELION³⁴$ to assess quality. The twelve highest populated classes from the 2D classification are shown.

The dataset contained 925,009 particles. A subset of particles was used to generate an ab*initio* map in cryoSPARC³¹. Using the low-pass filtered (30 Å) *ab initio* map as a template, two rounds of 3D heterogeneous refinement were performed using cryoSPARC in a binomial-like fashion. One major class emerged, which was refined using cryoSPARC homogenous refinement and then sharpened for model building.

B. Angular distribution calculated in cryoSPARC for particle projections. Heat map shows number of particles for each viewing angle (less=blue, more=red).

C. Gold-standard FSC⁴¹, calculated by comparing the two independently determined halfmaps from cryoSPARC. The dotted line represents the 0.143 FSC cutoff which indicates a nominal resolution of 4.4 Å.

D. RNAP clamp conformations. The RPo structure (Fig. 1C) was used as a reference to superimpose other structures via α-carbon atoms of the structural core module (table S2), revealing a common core RNAP structure (gray molecular surface) but with shifts in the clamp modules. The clamp modules are shown as backbone cartoons with cylindrical helices (RPo, blue; Cor-RbpA/holo, olive; RbpA/holo (6C05), cyan; open clamp (6BZO), red). The angles of clamp opening are shown (relative to RPo at 0°).

Extended Data Table 1 |

Cryo-EM data collection, refinement and validation statistics

 α Ramachandran plot parameters from PROCHECK 45

Extended Data Table 2 |

Structural superpositions.

 a The structural core module comprises the α subunits, the ω subunit, and conserved β and β' regions around the RNAP active center that have not been observed to undergo significant conformational changes.

 b_A Alignments performs using the PyMOL 'align' command.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Boyaci et al. Page 24

Figure 1 |. Structures of *Mtb* **transcription initiation complexes with AP3 promoter DNA.** A. Mtb AP3 promoter⁹ fragment used for cryo-EM. The core promoter elements and the TSS (+1) are denoted.

B. Overall structure of Cor-RP2. Proteins are shown as transparent surfaces and color-coded as shown in the key. The Cor and DNA (labeled with the −35, −10 elements and +1 colored as in (**A)**) are shown with transparent cryo-EM difference density and colored as labeled. **C.** Overall structures of RPo and RP2. Proteins and DNAs are shown as in (**B**) with DNA colors as labeled.

D. Superposed DNAs (colored as in **B** and **C**) from RPo, Cor-RP2, and RP2.

Figure 2 |. Protein/transcription bubble interactions.

DNAs are colored as in Fig. 1C. RNAP, CarD and RbpA residues that contact the DNA (< 4.5 Å) between −12 to +3 are shown as surfaces and labeled (FL2, fork loop 2; Sw2, switch 2; BH, bridge helix; AS, RNAP active site region). The double-arrowed black line denotes the shortest distance between the RNAP active site Mg^{2+} and an atom of the +1 t-strand DNA.

A. RPo.

B. RP2.

Figure 3 |. FL2, Sw2, and clamp dynamics.

A. RNAP clamp conformations. The RPo structure was used as a reference to superimpose the other structures by a common core RNAP structure (gray), revealing shifts in the clamp. Clamps are shown as cylindrical helices along with angles of clamp opening (relative to RPo at 0°). The open clamp structure (6BZO) is the *Mtb* RNAP bound to Fidaxomicin, which stabilizes the clamp in a fully open position without DNA in the active site cleft¹⁷. **B.** Overall structure of Cor-RP2 (upper left), with the boxed region magnified (lower right) showing the DNA and RNAP structural elements colored as labeled. Side chains of FL2

residues contacting the DNA (G462, S464, R467) are shown. Cor is also shown. The shortest distance between FL2 and Sw2 α-carbons is noted.

C. Same as **B** but showing RP2. Superimposed are the FL2, Sw2, and σ-fingers from the open clamp structure, aligned via the clamp, modelling a transient open-clamp intermediate. The shortest distance between FL2 and Sw2 α-carbons for each structure is noted. **D.** Same as **C** but showing RPo.

Reaction coordinate

Figure 4 |. Structural energetics for RPo formation.

Top; Three-step sequential kinetic scheme for RPo formation by bacterial RNAP⁹. Bottom; A schematic free energy profile for the last two steps of RPo formation is shown (red line) with cartoon structures of stable states and transitions, illustrating RNAP conformational changes. Details of the views are described in Methods. In RP1, the duplex DNA is outside of the RNAP cleft and must traverse through two physical barriers to reach the RNAP active site: 1) the βlobe/GL barrier, and 2) FL2/Sw2. In the first transition, the βlobe/GL and/or the clamp must swing open to allow the DNA to pass. In the second transition, the clamp must open to allow the ss t-strand DNA to pass through the FL2/Sw2 gap to access the RNAP active site.