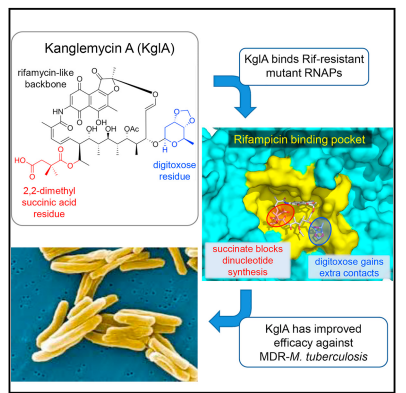
Molecular Cell

Mode of Action of Kanglemycin A, an Ansamycin Natural Product that Is Active against Rifampicin-Resistant *Mycobacterium tuberculosis*

Graphical Abstract



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In Brief

Resistance to rifamycins, inhibitors of bacterial RNA polymerase used for treatment of tuberculosis, is increasing. Mosaei et al. report an analog of the rifamycins, kanglemycin A, that inhibits rifampicin-resistant RNA polymerases and is effective against multidrugresistant *M. tuberculosis*, and they describe its mechanism of action.

Highlights

- Rifamycin kanglemycin A (KgIA) inhibits rifampicin-resistant RNA polymerases (RNAPs)
- KgIA's extra groups improve binding to RNAP and block first dinucleotide formation
- KgIA is effective against multidrug-resistant M. tuberculosis





Mode of Action of Kanglemycin A, an Ansamycin Natural Product that Is Active against Rifampicin-Resistant *Mycobacterium tuberculosis*

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SUMMARY

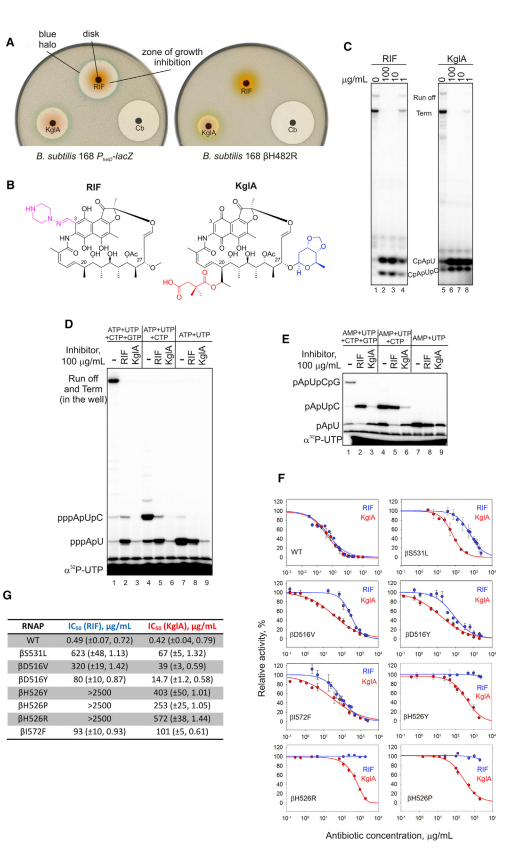
Antibiotic-resistant bacterial pathogens pose an urgent healthcare threat, prompting a demand for new medicines. We report the mode of action of the natural ansamycin antibiotic kanglemycin A (KgIA). KgIA binds bacterial RNA polymerase at the rifampicin-binding pocket but maintains potency against RNA polymerases containing rifampicin-resistant mutations. KgIA has antibiotic activity against rifampicin-resistant Gram-positive bacteria and multidrug-resistant Mycobacterium tuberculosis (MDR-M. tuberculosis). The X-ray crystal structures of KgIA with the Escherichia coli RNA polymerase holoenzyme and Thermus thermophilus RNA polymerase-promoter complex reveal an altered-compared with rifampicin-conformation of KgIA within the rifampicin-binding pocket. Unique deoxysugar and succinate ansa bridge substituents make additional contacts with a separate, hydrophobic pocket of RNA polymerase and preclude the formation of initial dinucleotides, respectively. Previous ansa-chain modifications in the rifamycin series have proven unsuccessful. Thus, KgIA represents a key starting point for the development of a new class of ansachain derivatized ansamycins to tackle rifampicin resistance.

INTRODUCTION

Antibiotic resistance is on the increase, exacerbated by the fact that, in recent years, only a few new compounds have reached the clinic. RNA polymerase (RNAP) is an essential enzyme accomplishing transcription in all living organisms. At the same time, the sequence divergence between bacterial and human RNAPs makes bacterial RNAP an excellent target for antibiotic development (Ho et al., 2009). In the last few years, details of the mode of action of a number of inhibitors of RNAP have been described (Bae et al., 2015; Degen et al., 2014; Maffioli et al., 2017; Molodtsov et al., 2015; Mukhopadhyay et al., 2008; Zhang et al., 2014a). However, most of these molecules have limited clinical utility, mainly because of their poor penetration of the bacterial cell envelope or strong binding to human plasma protein. Nevertheless, the search for new molecules that target RNAP and analysis of their mode of action is a promising approach for the development of new antibiotics, as exemplified by the now clinically used fidaxomicin (trade name Dificid) for Clostridium difficile-associated diarrhea (Tupin et al., 2010).

The most studied class of antibiotics known to inhibit bacterial RNA transcription are the rifamycins, a family of mainly semisynthetic ansamycins derived from the natural product rifamycin S, of which rifampicin (RIF) is the most widely used (Aristoff et al., 2010). RIF (and the other rifamycins) binds in a pocket (the RIF pocket) on the β subunit of RNAP in the vicinity of the catalytic site and sterically blocks propagation of the growing RNA chain through the main channel of RNAP (Campbell et al., 2001; McClure and Cech, 1978). The collision of RNA and RIF happens when the RNA reaches a length of 3-4 nucleotides, after which the RNA is released from the promoter complex as an abortive transcript. The length of the abortive RNA product can be influenced by the nature of the C3 substituent of the individual semisynthetic rifamycin. For example, rifabutin even inhibits synthesis of a trinucleotide (when reaction is started with a 5'-nonphosphorylated dinucleotide primer), whereas RIF still allows it (Artsimovitch et al., 2005).

The rifamycins are used, as part of the current combination therapy, for the first-line treatment of tuberculosis (TB), the



(legend on next page)

leading cause of death by infectious disease worldwide (World Health Organization, 2017), as well as treatment for non-tuberculous mycobacterial infections (Ryu et al., 2016). One of the major problems with known rifamycins is the rapid selection for mutations in the RIF-binding pocket of RNAP that lead to resistance. Multidrug-resistant strains of Mycobacterium tuberculosis (MDR-M. tuberculosis) present a major clinical problem because these strains are resistant to RIF in addition to isoniazid. In 2016, there were 600,000 new cases with resistance to RIF, of which 490,000 had MDR-TB. With a treatment success rate of only 54%, this highlights the requirement for improved treatment options. There are ~ 100 different amino acid substitutions in the RIF pocket that have been found in RIF-resistant clinical isolates of *M. tuberculosis* (Sandgren et al., 2009), with mutations at three positions of the RIF pocket that account for \sim 70% of the total substitutions (D516, 7.4%; H526, 20%; S531, 42%; E. coli numbering).

Previous efforts to probe the structure-activity relationships of the rifamycins and, thus, to maximize their binding affinity for RNAP have shown that the minimum structural requirements essential for activity include two free hydroxyl groups at the C21 and C23 positions of the ansa chain and two polar (free hydroxyl or carbonyl) groups at the C1 and C8 positions of the napthoquinone ring. Furthermore the molecule must be able to adopt a specific conformational geometry to position these key functional groups in the correct spatial arrangement required for RNAP binding (Bacchi et al., 1998; Lancini and Zanichelli, 1977). Thus, modification of the ansa bridge (e.g., hydrogenation or epoxidation of the C16,17 or C18,19 carbon-carbon double bonds) results in decreased binding to RNAP because of conformational changes in the molecule altering the spatial arrangement of the C1, C8, C21, and C23 functional groups, thus disrupting their critical hydrogen bonds with the RIF pocket (Aristoff et al., 2010; Bacchi et al., 2008). Hence, the production of clinically used semisynthetic rifamycins (RIF, rifabutin, rifapentine, rifalazil, and rifaximin) has focused on the introduction of functionality at the C3 position to stabilize the oxidation state of the naphthoquinone core and to improve bioavailability (Floss and Yu, 2005; Perlman, 1977).

Here we isolated and, by using microbiological, biochemical, and structural biology techniques, studied an uncharacterized naturally produced antibiotic, kanglemycin A (KgIA). KgIA is an ansamycin antibiotic from the rifamycin family that displays two important and unusual *ansa* bridge modifications. We show that, despite having a mode of action similar to that of RIF, KgIA has a larger binding surface with the RNAP provided by interactions of the *ansa*-bridge substituents and areas of RNAP that have not been previously accessible by rifamycin derivatives. As a result, KgIA has an altered binding conformation compared with RIF, which makes it maintain higher activity against both RIF-resistant RNAPs and RIF-resistant bacteria.

RESULTS

Isolation of KgIA

Actinomycete bacteria have historically been an excellent source of bioactive compounds, particularly antibiotics. To identify new antibiotics targeting transcription, we screened a subset of strains from the Demuris collection of actinomycetes for their ability to inhibit cell growth and to inhibit cellular transcription. As a reporter strain, we used a previously described B. subtilis strain that has a lacZ gene fused to the promoter of the bacterial helicase helD. This promoter was selected as the one that is upregulated during the stress caused by partial inhibition of transcription in sub-inhibitory concentrations of an antibiotic (Urban et al., 2007a). In the disk assay shown in Figure 1A, when a paper disk soaked with antibiotic is placed on the X-gal-infused agar plate with a lawn of growing cells, this results in a colorimetric response at the edge of the zone of growth inhibition, observed as a blue halo (Figure 1A, left; compare the halos produced by RIF and carbenicillin, an inhibitor of cell wall synthesis). The isolate Amycolatopsis DEM30355 was of particular interest because it produced a compound (later identified as KgIA, see below) that gave a blue halo at the edge of the zone of bacterial growth inhibition, indicative of transcription inhibition (Figure 1A, left). Importantly, in contrast to RIF, this new compound inhibited the RIF-resistant B. subtilis strain (BH482R; BH526R in E. coli numbering) (Figure 1A, right). We cultivated Amycolatopsis DEM30355 in a stirred tank bioreactor for 103.5 hr and extracted the secondary metabolites from the supernatant using Amberlite XAD-16 beads. Subsequent elution with methanol. extraction with ethyl acetate, and purification by multistep chromatography (STAR Methods) produced an orange crystalline compound that retained the observed biological activity against both the reporter and RIF-resistant strains (Figure 1A). Electrospray ionization-mass spectrometry (ESI-MS) revealed the mass, $[M+Na]^+$ = 1004.3866 m/z (Figure S1A), which corresponded to the mass of the previously described KgIA. ¹H and ¹³C nuclear magnetic resonance (NMR), extensive 2D NMR analysis, and, finally, single-crystal X-ray analysis (Figures S1B-S1D;

Figure 1. Identification and Mode of Action of KgIA

(E) The experiment was performed as in (D) but in the presence of AMP instead of ATP.

(F and G) Inhibition of wild-type and RIF-resistant E. coli RNAPs by RIF and KgIA. Error bars (F) are ± SD. The brackets (G) contain ± SE and the Hill slope.

⁽A) Disk diffusion assay with rifampicin (RIF), carbenicillin (Cb), and kanglemycin A (KgIA). Paper disks soaked with antibiotics were placed on the lawns of wildtype *B. subtilis* carrying the *lacZ* gene under the *P*_{*helD*} promoter, inducible during partial inhibition of transcription (left), and RIF-resistant *B. subtilis* (right). Note the blue halo around the growth inhibition zones in the case of transcription inhibitors.

⁽B) Chemical structure of KglA in comparison with RIF. The *ansa* chain modifications in KglA are highlighted. The semisynthetic group of RIF is shown in purple. See also Figure S1 and Table S1.

⁽C) Transcription *in vitro* by *E. coli* RNAP started with CpA on a linear template containing the T7A1 promoter. Runoff, termination, and abortive tri- and tetra-nucleotides are marked. Note that tetra-nucleotides migrate faster than tri-nucleotides under these electrophoretic conditions.

⁽D) Different ratio of the di-nucleotide-long (pppApU) and the tri-nucleotide-long (pppApUpC) abortive products in the presence of RIF and KglA. Transcription was performed in the presence of the nucleotides depicted (in the absence of the CpA primer). Note that, in the 33% gel, runoff and termination products remain in the well.

Table S1) unambiguously confirmed the structure of the isolated compound to be KgIA (Figure 1B).

KglA is an antibiotic originally isolated from the fermentation broth of *Nocardia mediterranei* var. *kanglensis* 1741-64 (Wang et al., 1988), for which no mode of action has been previously proposed, and only limited information regarding its biological activity has been reported. It is an ansamycin antibiotic that shares similarities with rifamycins, albeit with a number of unusual structural features. Most notably, KglA differs from known rifamycins and their semisynthetic derivatives in that it contains two large *ansa* bridge substituents, a pendant 2,2-dimethyl succinic acid side chain at C20 and a unique sugar moiety (β -O-3,4-*O*,O'-methylene digitoxose) at C27 (Figure 1B). Furthermore, these modifications are unusual for bioactive natural and semisynthetic rifamycins, whose modifications are typically concentrated at the naphthoquinone core.

Mode of Action and Specificity of KgIA

Activation of the *helD* promoter and its structural similarity to the rifamycins suggested that KglA targets bacterial RNAP. Therefore, we analyzed the effects of KglA on *in vitro* transcription by wild-type *E. coli* RNAP. KglA efficiently inhibited transcription on a linear DNA template containing the strong T7A1 promoter (Figure 1C). Inhibition of full-sized RNA products (run off or terminated) coincided with increased accumulation of short abortive tri- and tetra-nucleotide products (Figure 1C; the reactions were started with the dinucleotide primer CpA). Such a mode of action is characteristic of the rifamycins (Campbell et al., 2005; McClure and Cech, 1978), as can be seen from the pattern of inhibition of transcription by RIF (Figure 1C). In this mode of action, the antibiotic binds in the pathway of the nascent RNA, physically blocking its extension and inducing the release of short RNAs as abortive products.

Dinucleotide primer CpA that was used to start transcription does not carry a negatively charged triphosphate moiety at the 5' end, which is present in NTPs used by RNAP in vivo. Therefore, we analyzed the effects of RIF and KgIA in a native situation when transcription was started with ATP (in the absence of the CpA primer). As can be seen in Figure 1D, KgIA completely blocked the formation of the triphosphorylated trinucleotide, pppApUpC, whereas RIF allowed some of its production (compare lanes 4, 5, and 6). Furthermore, KglA strongly diminished (70-fold inhibition) the formation of the first dinucleotide, pppApU, whereas RIF had only a moderate effect (3-fold inhibition) (compare lanes 7, 8, and 9). Next, to start the reaction, we used AMP, which lacks γ and β 5' phosphates, present in ATP (Figure 1E). KgIA only moderately inhibited formation of pApU (4-fold), suggesting that it is γ and, possibly, β phosphates at the 5' end of the initiating nucleotide that clash with the additional groups of KgIA (compare lanes 7, 8, and 9). However, the formation of trinucleotide pApUpC was still strongly inhibited by KgIA compared with RIF (Figure 1E, lanes 4, 5, and 6). These results further suggest that extra groups and/or the mode of binding of KgIA provide an additional steric and/ or electrostatic (KgIA has a negatively charged C20 2,2dimethyl succinic acid side chain) hindrance for translocation of the 5' phosphorylated dinucleotide RNA. Furthermore, KglA interference with the 5' triphosphate moiety of the initiating nucleotide leads to inhibition even of the first phosphodiester bond formation (see structural support for these findings below).

The half maximal inhibitory concentration (IC₅₀) values of KgIA and RIF were similar against wild-type RNAP (Figures 1G and 1F). However, despite having a similar mode of action as RIF, we observed that KgIA killed RIF-resistant bacteria (compare the inhibition zones in Figure 1A, right). To investigate this phenomenon, we prepared E. coli RNAPs carrying a series of RIFresistant mutations, including BS531L, BD516V, BD516Y, βH526Y, βH526P, βH526R, and βI572F. These mutations correspond to the RIF resistance mutations in M. tuberculosis most frequently observed in clinical isolates, with *β*S531L, *β*D516V, and BH526Y accounting for 73% of all clinical RIF-resistant M. tuberculosis isolates (Gill and Garcia, 2011). Compared with RIF, KgIA was much more potent against six of seven RIF-resistant RNAPs (Figures 1G and 1F); in particular, KglA was 10 times more active than RIF against the β S531L and β D516V mutants. Note that, although mutations at BH526 made RNAP insensitive to RIF (no inhibition at 2.5 mg/mL), KgIA was still able to inhibit RNAPs with these mutations.

These results and *in vivo* data (Figure 1A) suggest that KgIA binds to RNAP in the RIF-binding pocket of the β subunit but in a manner different from known rifamycins, resulting in an increased resilience to the RIF-resistant amino acid substitutions in the RIF-binding pocket of RNAP. Furthermore, the mode of action of KgIA deviates from that of RIF because KgIA more efficiently inhibits the early stages of RNA synthesis.

Structural Basis of RNAP Inhibition by KgIA

To reveal the basis of the KgIA efficacy enhancement as induced by the modifications of the ansa bridge in comparison with rifamycin, we performed a crystallographic analysis of the complexes of KgIA with RNAPs. To analyze the RNAP and KgIA interaction on the atomic scale, a crystal of the T. thermophilus RNAP σ^{A} holoenzyme-*pyrG* promoter DNA complex (Murakami et al., 2017) was used to determine the X-ray crystal structure of the RNAP-KgIA complex at 3.0-Å resolution (Table 1). Additionally, to understand the mechanism of inhibition of RIF-resistant RNAP by KgIA, the crystal of the *E. coli* β S531L RNAP σ^{70} holoenzyme (Molodtsov et al., 2017) was used to determine the structure of the RNAP-KgIA complex at 4.1-Å resolution (Table 1). The electron density maps of both RNAPs show unambiguous densities for the compound located in the RIF-binding pockets in the β subunits (Figures 2A and 2B). The binding modes of KgIA to the RNAPs are almost identical; therefore, the atomic resolution structure of T. thermophilus RNAP-KgIA was used for the precise modeling of KgIA in complex with E. coli RNAP.

The binding mode of KgIA is similar to that of RIF and involves the formation of hydrogen bonds with the same amino acid residues of the RIF-binding pocket, including β Q393, β S411, and β R420 (side chain) and β F394 (main chain) (Figures 2C and S2A; the numbering is for *T. thermophilus*; see Figures S2C and S2D for the corresponding amino acid residues in *E. coli* and *M. tuberculosis* RNAPs). Because of a different topology of the ansa bridge of KgIA (Figure S2B), the C23 hydroxyl group does not form an additional hydrogen bond with β F394, as observed in the *T. thermophilus* RNAP-RIF complex. However,

Table 1. Collection of Crystallographic Data and Refineme	nt
Statistics	

Statistics		
Complex	<i>Eco</i> RNAP – KgIA	<i>Tth</i> RNAP – KgIA
PDB code	6CUX	6CUU
Data Collection		
Space group	P2 ₁ 2 ₁ 2 ₁	C2
Cell dimensions		
a (Å)	188.167	185.221
b (Å)	204.685	101.315
c (Å)	311.577	294.279
β (°)	90	98.886
Resolution (Å)	50.0-4.1	50.0-3.0
Total reflections	608,898	358,319
Unique reflections	93,684	107,473
Redundancy	6.5 (6.2)	3.3 (3.0)
Completeness (%) ^a	99.9 (100.0)	99.7 (97.1)
l/σ ^a	16.2 (1.95)	10.2 (1.14)
R _{sym} (%) ^a	7.7 (85.1)	9.6 (61.4)
CC ^{1/2a}	(0.633)	(0.629)
Refinement		
Resolution (Å)	45.03–4.1	46.3–3.0
R _{work}	0.2100	0.2012
R _{free}	0.2547	0.2483
No. of atoms	54,867	28,254
RMSDs		
Bond length (Å)	0.013	0.006
Bond angles (°)	1.776	1.089
Clashscore	11.47	5.84
Ramachandran favored, %	91.57	97.64
Ramachandran outliers, %	0.94	0.35
RMSD, root-mean-squ	are deviation.	

niviod, root-mean-square deviation.

^aHighest-resolution shells are shown in parentheses.

the C27 β -O-3,4-O,O'-methylene digitoxose side chain establishes a hydrogen bond with the side chain of β R134 (Figure 2C). This KgIA *ansa* bridge side chain inserts into a shallow pocket in the RIF-binding site and establishes additional non-polar interactions with the surrounding residues, significantly increasing the binding surface of KgIA compared with RIF (Figure 2C). One of the structural consequences of the sugar side chain accommodation in the RIF-binding pocket is a rotational shift of KgIA that increases the distance between the naphthalene core and the binding surface of the pocket compared with RIF (Figure 2D).

To validate the new interaction of the C27 β -O-3,4-O,O'-methylene digitoxose side chain with β R134, we constructed an *E. coli* RNAP carrying a mutation at the corresponding amino acid residue, β R143A. The mutation had little effect on RIF sensitivity (Figure S2E, left). However, KgIA was at least an order of magnitude less effective against β R143A RNAP than against wild-type RNAP, indicating the importance of this interaction for KglA binding (Figure S2E, center). The IC₅₀ of KglA for β R143A RNAP was still lower than for any RIF-resistant mutant RNAP, indicating that interactions of KglA within the rest of the RIF-binding pocket were still critical for its binding. Only a double mutation, β R143A/S531L, led to KglA resistance (Figure S2E, right).

The presence of additional bulky side chains on the ansa bridge of KgIA has an effect on the mechanism of de novo transcription initiation. Ansamycin binding in the RIF-binding pocket inhibits transcription by clashing with the 5' end of a nascent RNA. Modeling shows that, in the T. thermophilus RNAP-RIF complex, the triphosphate moiety of the initiating nucleoside triphosphate (NTP) guided by hydrogen bonds with side chains of two essential residues, *β*Q567 and *β*H999 (Basu et al., 2014), moderately clashes with the C3 side chain of RIF (Figure 2E, right). This clash does not strongly affect the formation of the first phosphodiester bond, likely because of flexibility of the C3 side chain. Even in the presence of RIF, there is an empty space (triphosphate retreat zone) that allows for accommodation of the 5' triphosphate group followed by an extension of the nascent RNA to 3-mer (McClure and Cech, 1978) before a severe clash with the +1 NTP inhibits further primer extension (Figure 2E, right). In contrast, the positional shift of KgIA and the orientation of the negatively charged C20 2,2-dimethyl succinic acid side chain not only do not leave space for accommodation of the 5' triphosphate group but also interfere with the correct placement of the initiating NTPs (Figure 2E, left), which explains the strong inhibition of the initial dinucleotide synthesis (Figure 1D). The electrostatic repulsion by the C20 2,2-dimethyl succinic acid side chain of KgIA can also explain the inhibition of monophosphorylated pApUpC synthesis (which is allowed by RIF; Figure 1E).

A comparative analysis of the structures of the E. coli ßS531L RNAP σ^{70} holoenzyme in apo form and in complex with RIF and KglA reveals the mechanism of KglA's efficiency against many RIF-resistant forms of mutant RNAPs (Figure 2F). The amino acid substitution BS531L is the most clinically important RIFresistant mutation in bacterial RNAP because of its low structural and functional effects on the enzyme in the absence of RIF. As shown in a recent study (Molodtsov et al., 2017), RIF binding to the mutant RNAP results in a steric clash of the naphthalene core with the side chain of β 531L, which triggers disordering of a constituent part of the RIF-binding pocket, the fork loop2 (Figure 2F, left versus right). This partial disordering of the RIF-binding site efficiently prevents RIF binding. The naphthalene core of KgIA is almost identical to RIF; however, it is rotated in the binding site compared with RIF (Figure 2D). This positional shift allows KgIA to maintain a distance to §531L, avoiding a severe clash with the residue and minimizing structural rearrangements of the fork loop2 and the binding pocket (Figure 2F, center). This fine-tuning of the core binding mode and the increased interaction surface provided by the C27 sugar side chain enable high affinity of KgIA in a broad context of RIF resistance.

Kanglemycin A Has Antimicrobial Activity against MDR-*M. tuberculosis* and Other Gram-Positive Pathogens

The above results suggested that KgIA may be effective against RIF-resistant bacteria, in particular against clinically

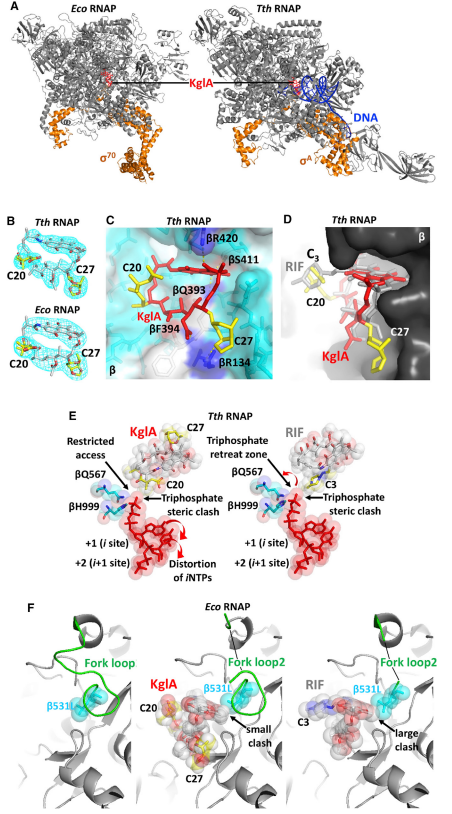


Figure 2. KgIA Binding to Free and Promoter-Bound RNAP Holoenzyme

(A) Overall views of *E. coli* (left) and *T. thermophilus* (right) RNAPs with KgIA bound in the RIF-binding pocket. RNAPs (gray core and orange σ) and DNA (blue) are shown as ribbon models, and KgIA (red) is shown as a stick model.

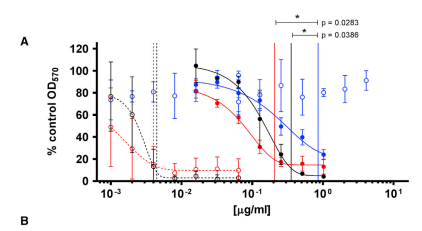
(B) Electron density maps of KglA in complexes with *T. thermophilus* (top) and *E. coli* (bottom) RNAPs. Electron densities are shown as a light blue mesh, and KglA molecules are shown as stick models colored by element. The *ansa* bridge side chains of KglA extending from C20 and C27 are colored in yellow.

(C) A close-up view of the RIF-binding pocket of the *T. thermophilus* RNAP containing KgIA (in red, with the side chains in yellow). RNAP is shown as a transparent surface model (light blue), KgIA is shown as a stick model with the *ansa* bridge side chains labeled, and the β subunit residues forming the RIF-binding pocket are shown as stick models. The hydrogen bonds between KgIA and β amino acid residues (colored by element) are depicted by yellow dotted lines; the β R134 involved in polar interaction with the C20 sugar of KgIA is shown in blue.

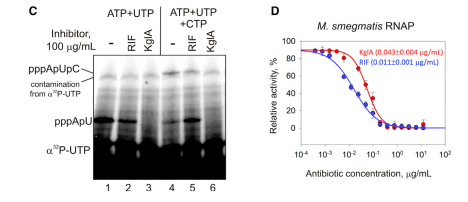
(D) A side view of the RIF-binding pocket shown in (C). KgIA is overlaid on RIF (gray); the side chains of the two molecules are labeled. KgIA maintains a larger distance to the wall in the RIF-binding pocket than RIF.

(E) Modeling of the initiating nucleotides in the active site of the *T. thermophilus* RNAP (from PDB: 4Q4Z) in the presence of KgIA (left) and RIF (right). The ribonucleotides (red), the β Q567 and β H999 residues coordinating the 5' end triphosphate (blue), and KgIA and RIF (gray) are shown as stick and sphere models. The side chains of the inhibitors and the positions of the initiating NTPs are indicated. Red arrows indicate the translocation directions for the corresponding substrates.

(F) Structural rearrangements in the RIF-binding pocket of *E. coli* β S531L RNAP triggered by KgIA and RIF binding. RNAP is shown as a ribbon model in gray, the structured regions of the fork loop2 are shown in green, and the β 531L residue (blue), KgIA (colored by element with yellow side chains), and RIF (colored by element) are shown as stick and sphere models. The side chains of KgIA and RIF are indicated. Arrows mark the clash points between the naphthalene cores of the corresponding compounds with the β 531L side chain. See also Figure S2.



Organism/isolate	Strain Reference	RIF- Resistance	RIF, μg/mL	KglA, μg/mL
M. tuberculosis	H37Rv	susceptible	0.005 ± 0.001	0.33 ± 0.09
M. tuberculosis	1192/015	susceptible	0.004 ± 0.001	0.21 ± 0.07
Multi-drug-resistant M. tuberculosis	08/00483E	resistant	>4.096	0.85 ± 0.02



problematic MDR-M. tuberculosis strains. Thus, we tested KgIA against a clinical MDR-M. tuberculosis isolate (Beijing 08/00483E) that is fully resistant to RIF (Jamieson et al., 2014; Zhang et al., 2014b) and the other first-line antibiotics (isoniazid, ethambutol, and pyrazinamide). Beijing 08/00483E carries the most frequent RNAP amino acid change found in RIF-resistant M. tuberculosis (BS450L, M. tuberculosis numbering; ßS531L, E. coli numbering; Figures S2C and S2D). As positive controls, we used the RIF-susceptible reference strain of M. tuberculosis H37Rv and a RIF-susceptible clinical isolate, Beijing 1192/015. The clinical strain Beijing 08/00483E was resistant to RIF at 1,024 times the minimum inhibitory concentration (MIC) for RIF-sensitive strains (Figures 3A and 3B). In contrast, all three strains displayed a dosedependent response to KgIA exposure (Figure 3A), demonstrating its antimicrobial activity against MDR-M. tuberculosis. KgIA was statistically significantly more active against RIFsensitive Beijing 1192/015 than RIF-resistant Beijing 08/ 00483E (Figures 3A and 3B).

Figure 3. KgIA Inhibits Mycobacterial RNAP and Retains Antimicrobial Activity against MDR-*M. tuberculosis*

(A) Dose-dependent growth inhibition of the RIF-susceptible *M. tuberculosis* strains H36Rv (black) and 1192/015 (red) and RIF-resistant *M. tuberculosis* 08/00483E (blue) by KgIA (solid line, solid symbols) and RIF (dashed line, open symbols). Data presented are the mean of four independent experiments \pm SD and analyzed using a Student's t test.

(B) MIC values \pm SD, as determined using the modified Gompertz function for *M. tuberculosis* strains H37Rv, 1192/015, and 08/00483E, which were exposed to RIF or KgIA.

(C) Mode of action of KgIA and RIF against *M. smegmatis* RNAP. The experiment was performed exactly as in Figure 1D. Note that the high radioactive background compared with Figure 1D is due to much lower activity of *M. smegmatis* RNAP.

(D) Inhibition of *M. smegmatis* RNAP by RIF and KgIA.

Error bars are \pm SD. Shown in brackets are the IC_{50} values \pm SE.

As seen in Figures 3A and 3B, RIF was more active against RIF-susceptible *M. tuberculosis* cells than KgIA. This apparently contradicted our *in vitro* results of very similar inhibition of *E. coli* wild-type RNAP by both inhibitors. It is possible that the mode of inhibition of *M. tuberculosis* RNAP by KgIA is different from that of *E. coli* RNAP. Therefore, we analyzed inhibition of transcription by *Mycobacterium smegmatis* RNAP, which is commonly used as an experimental model for *M. tuberculosis* RNAP. Both RNAPs have an identical amino acid composition of

the RIF-binding pocket (Figure S2C). We found that RIF and KgIA inhibited RNA synthesis by *M. smegmatis* RNAP and *E. coli* RNAP in a very similar way (compare Figure 1D with Figure 3C); i.e., KgIA fully blocked synthesis of the first dinucleotide (pppApU), whereas RIF partly allowed synthesis even of the trinucleotide (pppApUpC). The IC₅₀ of KgIA was nearly the same as that of RIF. Therefore, the affinity of RNAP binding cannot account for the observed difference in the activities of RIF and KgIA against RIF-sensitive *M. tuberculosis* cells. The likely explanation of differences in activities of KgIA and RIF against RIF-sensitive *M. tuberculosis* is the decreased ability of KgIA to pass through the cell envelope.

We further tested KgIA against a range of other species, including a number of clinically relevant pathogens. These included 8 Gram-positive bacteria, 10 Gram-negative bacteria, and 2 eukaryotes (Table 2). Of 10 Gram-negative pathogens, KgIA was active against only *Providencia rettgeri*. No activity was observed toward the two strains of eukaryotic microorganisms (yeast): *Candida albicans* and *Candida glabrata*. Among the

	Organism/Isolate	Strain Reference	KglA (μg/mL)	RIF (μg/mL)
aram-Negative	Acinetobacter baumannii	ATCC 19606	>128	–
ian nogatio	Burkholderia cepacia	ATCC 25416	>128	-
	Enterobacter cloacae	NCTC 11936	>128	_
	Escherichia coli	NCTC 10418	>128	16ª
	Klebsiella pneumoniae	NCTC 9528	>128	-
	Providencia rettgeri	NCTC 7475	64	_
	Pseudomonas aeruginosa	NCTC 10662	>128	_
	Salmonella enterica Typhimurium	NCTC 74	>128	_
	Serratia marcescens	NCTC 10211	>128	_
	Yersinia enterocolitica	NCTC 11176	>128	_
ram-Positive	Bacillus subtilis	NCTC 9372	1	_
	Enterococcus faecalis	NCTC 775	64	_
	Enterococcus faecium	NCTC 7171	128	_
	Listeria monocytogenes	NCTC 11994	8	_
	Staphylococcus epidermidis	NCTC 11047	1	_
	Staphylococcus aureus	NCTC 6571	0.25	0.004 ^ª
	Staphylococcus aureus (MRSA)	NCTC 11939	1	-
	Streptococcus pyogenes	NCTC 8306	0.031	_
ast	Candida albicans	ATCC 90028	>128	_
	Candida glabrata	NCPF 9725	>128	_
aureus	Bel	97597	2	2
aureus	Bel	97598	64	>128
	Fin	37481	0.5	≤0.008
	Fin	54511	32	>128
	Fin	54518	128	>128
	Fra	462	0.5	≤0.008
	Fra	920	1	≤0.008 ≤0.008
	Fra	95035	1	<u>≤</u> 0.000
	Ger	131/98 (Sger 11d2)	0.5	0.016
	Ger	1966/97 (Han 111c)	0.5	
	Ger	2594-1/97 (Sger 11a)	1	≤0.008 0.016
	Ger			
		2594-2/97 (Sger 11b) 1729/98	0.25	≤0.008
	EMRSA 15 EMRSA 15		0.5	≤0.008 <0.008
		1758/98	0.5	≤0.008
	EMRSA 15	14956	0.5	≤0.008
	EMRSA 15	12484/98	0.5	≤0.008
	EMRSA 15	14185/98	0.5	≤0.008
	EMRSA 15	16822/98	0.5	≤0.008
	EMRSA 15	19972/98	0.5	≤0.008
	EMRSA 15	20460/98	0.5	0.016
	EMRSA 15	21268/98	0.5	≤0.008
	EMRSA 15	21698/98	0.5	≤0.008
	EMRSA 15	2501/98	0.5	≤0.008
	EMRSA 15	6323/98	0.5	≤0.008
	EMRSA 16	00036/95	1	0.016
	EMRSA 16	00998/95	1	≤0.008
	EMRSA 16	03732/95	1	0.016
	EMRSA 16	07121/95	1	\leq 0.008

(Continued on next page)

Organism/Isolate	Strain Reference	KglA (μg/mL)	RIF (μg/mL)
EMRSA 16	07924/95	0.5	≤0.008
EMRSA 16	18200/95	0.5	\leq 0.008
EMRSA 16	18205/95	0.5	\leq 0.008
EMRSA 16	21354/95	1	0.016
EMRSA 16	23728/98	1	\leq 0.008
EMRSA 16	23729/98	1	\leq 0.008
EMRSA 16	32010/96	1	0.016
EMRSA 16	34256/96	1	\leq 0.008
MRSA	NCTC 11939	1	\leq 0.008
Staphylococcus aureus	NCTC 6571	0.25	≤0.008

8 Gram-positive strains, KgIA was highly active (MIC, $\sim 1 \mu g/mL$) against staphylococci and Streptococcus pyogenes but less active against enterococci (MIC, 64-128 µg/mL). The relatively high activity of KgIA against staphylococci prompted evaluation of its activity against a large collection of methicillin-resistant Staphylococcus aureus (MRSA) and a comparison of its activity with RIF. S. aureus NCTC 6571 was re-tested as a control strain with a known MIC of RIF (EUCAST, 2018). Thirty-seven MRSA were tested in total, and 33 of these proved to be susceptible to RIF, as defined by a MIC of \leq 0.06 µg/mL (EUCAST, 2018). KgIA was substantially less active than RIF against RIF-susceptible MRSA, with MICs of KgIA between 0.25 and 1 μ g/mL. Conversely, for 3 of 4 RIF-resistant MRSA, MICs of KgIA were 1-3 double dilutions lower than those of RIF, although still relatively high (32–128 µg/mL). In the light of our in vitro data (Figures 1C–1F and 3C), no activity against most of Gram-negative bacteria, and lower-than-RIF activity against some Gram-positive bacteria support the idea of a diminished ability of KgIA to enter bacterial cells. This, however, requires further investigation coupled with further in vivo analyses to develop KgIA as a future antibiotic.

DISCUSSION

Easily acquired RIF resistance is a major problem in the clinical usage of rifamycins. The principal finding of our study is that KgIA is much more effective against RIF-resistant RNAPs and bacteria than its commonly used relative, RIF. The structural analysis suggests that this is explained by both additional interactions and the binding mode of KgIA with RNAP unique to KgIA. The sugar (β -O-3,4-O,O'-methylene digitoxose) and acid (2,2dimethyl succinic acid) moieties at C27 and C20 of KgIA, respectively, establish these additional contacts and influence the overall binding mode of KgIA in the pocket (Figures 2C and 2D). Compared with RIF, KgIA rotates to reposition slightly away (by 1.2 Å) from the wall of the binding pocket, which may decrease the effect of resistance mutations arising in RNAP on its affinity. Thus, the increased distance between the naphthalene core along with the increased binding surface provided by the ansa-bridge side chains likely represent the mechanisms that enable KgIA to maintain its inhibitory activity against many

RIF-resistant bacterial RNAPs. As the result of additional interactions by the sugar moiety with the protein, only an unlikely event of two simultaneous mutations in the RIF-binding pocket can lead to KgIA resistance.

KgIA inhibits RNA synthesis at the stage of the first phosphodiester bond (2-mer RNA) formation, which is different from RIF, which inhibits formation of the second phosphodiester bond (3-mer RNA) (Figures 1D and 2E). Our biochemical and structural data suggest that this may happen because of the steric and/or electrostatic clash of the C20 2,2-dimethyl succinic acid side chain of KgIA with γ and/or β phosphates of the +1 NTP that may disturb the conformation of initiating nucleotides and, thus, catalysis. We cannot directly compare these results with the previous study that found that some rifamycins may inhibit RNA synthesis one step earlier than RIF (Artsimovitch et al., 2005) because they used a 5' non-phosphorylated dinucleotide to start transcription. We, however, did not observe inhibition of 5' non-phosphorylated trinucleotide synthesis by KglA (Figure 1C), as observed for some of the rifamycins in that study (Artsimovitch et al., 2005), suggesting that inhibition of 5' triphosphorylated dinucleotide synthesis by KglA takes place via a different mechanism.

KgIA, compared with RIF, appears to show reduced penetration through the bacterial cell envelope, likely because of the presence of the polar carboxyl group at C20. For example, KgIA has minimal activity against Gram-negative bacteria in comparison with RIF and displays higher MICs than RIF against most of the Gram-positive species tested. However, KgIA is more effective than RIF against RIF-resistant Gram-positive bacteria. Notably, it is effective against MDR-M. tuberculosis. KglA inhibited the growth of MDR-M. tuberculosis (strain 08/00483E at a MIC of 0.854 µg/mL [850.3 nM]), which was completely resistant to RIF because of a *β*S450L substitution (*β*S531L in E. coli numbering). Finding new antibiotics for the treatment of MDR-TB is a challenge, and KgIA demonstrates potential as a candidate for further development, particularly in light of the fact that the MIC of KgIA is lower than the MIC values displayed for both isoniazid and pyrazinamide against drug-susceptible M. tuberculosis, and there is a wealth of information about drug toxicity and patient responses to RIF. These considerations are likely to accelerate the progression of KgIA or its derivatives to the clinic for the treatment of both drug-susceptible and drug-resistant TB.

A new mode of KgIA binding to RNAP may enrich the strategy of structure-guided antibiotic design against RIF-resistant pathogens such as MDR-M. tuberculosis. RIF resistance caused by a point mutation of the β subunit of RNAP is associated with either loss of important contact between a target and a compound (in the case of BS531L) or steric exclusion of a compound from its binding site (in the case of β H526Y) (Molodtsov et al., 2017). In many cases, research efforts focusing on developing new antibiotics targeting the RIF-resistant bacterial RNAP are searching for new interaction surfaces away from the RIF-binding pocket (Lin et al., 2017; Maffioli et al., 2017). The enhanced affinity of KgIA for certain RIF-resistant RNAPs along with the moderate changes in its mode of binding compared with RIF (Figure 2D) indicate that the RIF-binding pocket still possesses significant potential for the development of potent antibiotics against certain RIF-resistant pathogens. Targeting the RIF-binding pocket has some advantages compared with targeting a new site. For example, development of resistance to KgIA is possible via additional mutations in the RIF-binding pocket (e.g., βR143A; Figure S2E) in addition to the first RIF-resistant mutation (e.g., β S531L); however, such mutations may compromise the transcriptional integrity and further diminish the fitness of the RNAP, rendering the resulting pathogens non-viable (Song et al., 2014). Another important advantage of developing new rifamycin derivatives is that this approach exploits the well-known rifamycin scaffold and may likely overcome some obstacles common in the development of new antibiotics, such as the problems of drug permeability through bacterial membranes, human toxicity, and specificity (Aristoff et al., 2010).

The modifications in the ansa bridge are highly unusual in bioactive rifamycins, with even minor semisynthetic modifications in an ansa bridge often leading to loss of RNAP inhibition (see Introduction). Our study reveals the significant potential of increasing the affinity of the rifamycins to RNAP via ansa bridge modifications, in particular by demonstrating that C27 substitution can allow access to a previously unexplored hydrophobic pocket adjacent to the RIF-binding pocket while maintaining the critical interactions between the rifamycin backbone and the RIF-binding pocket of RNAP. Furthermore, substitution at C20 with a flexible carboxylic acid residue has revealed the potential for new drug-initiating NTP interactions in the rifamycins. Altogether, KgIA provides an example of an evolved solution to RIF resistance in which these unusual ansa bridge modifications lead to improved potency against RIF-resistant RNAPs and represents an important starting point for the development of new therapeutic agents.

STAR * METHODS

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

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- METHOD DETAILS
 - Disk diffusion assay
 - Isolation of KgIA from Amycolatopsis sp. DEM30355

- Small molecule X-ray crystallography including absolute configuration of KgIA
- NMR analysis of KglA
- \odot Purification of *E. coli* RNAP core enzyme and σ^{70} , and *M. smegmatis* RNAP holoenzyme for transcription assays
- In vitro transcription
- Crystallization of RNAP KglA complexes
- Crystallographic data collection and X-ray crystal structure determinations
- Determination of minimum inhibitory concentrations (MICs) for *M. tuberculosis*
- Determination of minimum inhibitory concentrations for other bacteria
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes two figures and two tables and can be found with this article online at https://doi.org/10.1016/j.molcel.2018.08.028.

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AUTHOR CONTRIBUTIONS

H.M., L.C., J.H., and Y.Y. performed biological experiments and analyzed the data. B.K., S.M.L., and N.E.E.A. purified KgIA. V.M., Y.S., and K.S.M. performed crystallographic experiments and analyzed the structures. M.J.H., S.M.L., and C.W. performed the NMR analysis and structural assignment of KgIA. W.C. performed small-molecule X-ray crystallography. C.W.M., R.E.J., and J.B. measured the MICs for the *M. tuberculosis* strains. J.D.P. and E.C.L.M. measured KgIA MICs for other species. J.E., N.E.E.A., and N.Z. conducted the study. J.B., C.W.M., V.M., M.J.H., K.S.M., and N.Z. wrote the paper.

DECLARATION OF INTERESTS

N.E.E.A. is an employee of and J.E. is a scientific founder of and shareholder in Demuris Ltd. Demuris and Newcastle University have filed UK patent 1812078.2.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
E. coli BL21(DE3)	Novagen	Cat# 69450
<i>M. smegmatis</i> mc ² 155 (SM07)	Mukherjee and Chatterji, 2008	N/A
<i>Mycobacterium tuberculosis</i> H37Rv, Beijing 1192/015 and Beijing 08/00483E	PHE National Mycobacterial Reference Laboratory	N/A
E. coli T7 Express	New England BioLabs	Cat# C2566H
B. subtilis 168 amyE::PheID-lacZ cat	Urban et al., 2007a	N/A
B. subtilis 168 amyE::PhelD-lacZ cat rpoB _{H482R}	This paper	N/A
Enterobacter cloacae	NCTC	NCTC 11936
Escherichia coli	NCTC	NCTC 10418
Klebsiella pneumoniae	NCTC	NCTC 9528
Providencia rettgeri	NCTC	NCTC 7475
Pseudomonas aeruginosa	NCTC	NCTC 10662
Salmonella typhimurium	NCTC	NCTC 74
Serratia marcescens	NCTC	NCTC 10211
Yersinia enterocolitica	NCTC	NCTC 11176
Bacillus subtilis	NCTC	NCTC 9372
Enterococcus faecalis	NCTC	NCTC 775
Enterococcus faecium	NCTC	NCTC 7171
Listeria monocytogenes	NCTC	NCTC 11994
Staphylococcus epidermidis	NCTC	NCTC 11047
Staphylococcus aureus	NCTC	NCTC 6571
Staphylococcus aureus (MRSA)	NCTC	NCTC 11939
Streptococcus pyogenes	NCTC	NCTC 8306
Candida glabrata	NCPF	NCPF 9725
Acinetobacter baumannii	ATCC	ATCC 19606
Burkholderia cepacia	ATCC	ATCC 25416
Candida albicans	ATCC	ATCC 90028
Clinical isolates of methicillin-resistant <i>Staphylococcus aureus</i> (MRSA) X 36.	Public Heath England, Colindale, UK	N/A
T. thermophilus HB8	ATCC	ATCC 27634
Chemicals, Peptides, and Recombinant Proteins		
Ni-NTA agarose	QIAGEN	Cat# 30230
HiTrap Q HP, 1 mL	GE Healthcare Life Sciences	Cat# 29051325
HiLoad 26/600 Superdex 200	GE Healthcare Life Sciences	Cat# 28989336
HiTrap Heparin HP, 5 mL	GE Healthcare Life Sciences	Cat# 17040601
HisTrap HP, 5 mL	GE Healthcare Life Sciences	Cat# 17524801
Bio-Rex 70 cation exchange resin	Bio-Rad	Cat# 1425842
Biotage® SNAP Ultra 100 g	Biotage	Cat# FSUL-0442-0100
[alpha-P32]Uridine 5'-triphosphate (UTP)	Hartmann Analytic	Cat# SRP-810
rNTP Set, 100 mM Solutions (ATP, CTP, GTP, UTP)	GE Healthcare	Cat# 27202501
QuikChange II kit (Stratagene)	Agilent	Cat# 200523
CpA	IBA Lifesciences	Cat# 5-0710-186
RIF	Sigma	Cat# R3501

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Resazurin sodium salt	Sigma	Cat# R7017
CMM MOD2	James et al., 2000	N/A
Middlebrook 7H9 medium	BD Difco	Cat# 271310
Phosphate buffered saline, pH 7.4	Severn Biotech	Cat# 20-74-10
soSensitest agar	Oxoid	Cat# CM0471
Amberlite XAD-16	Sigma	Cat# XAD16
Deposited Data		
KgIA X-ray diffraction data	This paper	CCDC: 1827601 and 1827602
E. coli RNAP – KgIA complex	This paper	PDB: 6CUX
. thermophilus RNAP – KgIA complex	This paper	PDB: 6CUU
E. coli RNAP	Murakami 2013	PDB: 4YG2
T. thermophilus RNAP – pyrG promoter complex	Murakami et al., 2017	PDB: 5VOI
Oligonucleotides		
See Table S2 for list of oligonucleotides		
Recombinant DNA		
oVS10(S531L)	Gill and Garcia, 2011	N/A
DGEMD	Murakami, 2013	N/A
bET21a-SigA	Basu et al., 2014	N/A
DETLrpoB	Smith and Savery, 2005	N/A
Software and Algorithms		
COOT	Emsley and Cowtan, 2004	https://www2.mrc-lmb.cam.ac.uk/ personal/pemsley/coot/
HKL2000	Otwinowski and Minor, 1997	http://www.hkl-xray.com
PHENIX	Adams et al., 2010	https://www.phenix-online.org
Pymol	PyMOL	https://pymol.org/2/
Graphpad 7.00	Prism	https://www.graphpad.com/
SigmaPlot 11.0	Systat Software	https://systatsoftware.com/
Other		
/DX48 Plate with sealant	Hampton Research	Cat# HR3-275
12 mm x 0.22 mm Siliconized circle cover slides	Hampton Research	Cat# HR3-279
18 mm Mounted CryoLoop - 20 micron	Hampton Research	Cat# HR4-973
CrystalCap ALS with vial	Hampton Research	Cat# HR4-779
/ivaspin sample concentrator (MWCO 10 kDa)	GE Healthcare Life Sciences	Cat# 28-9323-60
Fecan Sunrise microplate absorbance reader	Tecan instruments	https://www.tecan.com/
nfinite M1000 multimode plate reader	Tecan instruments	https://www.tecan.com/
solaera ONE	Biotage	Cat# ISO-1SW
Jeol Lambda 500 MHz spectrometer	JEOL Ltd	N/A
Jeol ECS 400 MHz spectrometer	JEOL Ltd	N/A
Gemini X-ray Diffractometer/Atlas detector	Oxford Diffraction	N/A
Beam Station I19	National Synchrotron Facility Diamond Light Source	N/A

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to Lead Contact (n.zenkin@ncl.ac.uk).

METHOD DETAILS

Disk diffusion assay

Standard disc diffusion assay (Kirby-Bauer) was performed with the *B. subtilis* 168 *amyE*:: P_{helD} -*lacZ cat* reporter strain or RIF-resistant *B. subtilis* 168 *rpoB_{H482R}* strain as described (Bauer et al., 1966). Briefly, disks were loaded with to 10 µg of each drug, air-dried and placed on nutrient agar plates with embedded lawn of bacteria and x-gal (100 µg/mL). Plates were incubated overnight at 37°C and scanned.

Isolation of KgIA from Amycolatopsis sp. DEM30355

The fermentation of the *Amycolatiopsis* sp. DEM303555 was conducted as previously described. (Kepplinger et al., 2018). One kilogram of Amberlite XAD-16 resin was washed with deionized water and then applied to 50 L of filtered culture supernatant in batch absorption. The material was eluted with 32 L of 100% methanol and concentrated under reduced pressure (2.5 L). The concentrate was adjusted to pH 6.5 and extracted eight times with an equal amount of ethyl acetate. The organic extracts were combined and evaporated to dryness to yield 22.5 g of crude extract. A sample of the crude extract (3.91 g) was subjected to silica gel chromatography (eluent = 100% ethyl acetate, Biotage SNAP 100 g cartridge). KgIA-containing fractions were combined and the solvent was removed under reduced pressure to give 1.24 g of a dark colored solid. This material was subjected to a second silica gel chromatography step (eluent = linear gradient from diethyl ether to ethyl acetate. Evaporation of the KgIA containing fractions yielded 806.2 mg as an orange solid.

Small molecule X-ray crystallography including absolute configuration of KgIA

Slow evaporation of a methanol solution of KgIA gave crystals suitable for single-crystal X-ray analysis. The molecular structure of KgIA was obtained to high precision using synchrotron radiation with a very small crystal of the methanol solvate at 120 K; all H atoms were located and freely refined. The short X-ray wavelength of 0.6889 Angstroms did not permit the determination of the absolute configuration, which was subsequently established unambiguously with CuKalpha radiation and a larger crystal, giving a molecular structure with otherwise slightly lower precision. CCDC 1827601 and 1827602 contain the supplementary crystallographic data for this paper, these data can be obtained free of charge from The Cambridge Crystallographic Data Centre (https://www.ccdc. cam.ac.uk/).

NMR analysis of KgIA

All NMR spectra were recorded in CDCl₃ at 10° C. ¹H and ¹³C{¹H} NMR spectra were recorded directly with a Jeol Lambda 500 MHz spectrometer (Figure S1), while ¹H COSY; 45, 90 and 135 DEPT; HMQC and HMBC were recorded with a Jeol ECS 400 MHz spectrometer (not shown). Proton and carbon assignment was performed *de novo* from both 1D and 2D NMR experiments and was in close agreement with the literature (Wang et al., 1988, Table S1).

Purification of *E. coli* RNAP core enzyme and σ^{70} , and *M. smegmatis* RNAP holoenzyme for transcription assays

RIF-resistant mutations were introduced in pETLRpoB plasmid (6xHis-tag on N terminus of β subunit of *E. coli* RNAP) (Smith and Savery, 2005) by site-directed mutagenesis using QuikChange II kit (Stratagene). Wild-type or mutant plasmids were transformed along with pACYCDuet-1_Ec_rpoZ (encoding for ω subunit) in T7 Express strain (New England Biolabs). Expression and purification of core RNAP was performed as described (Kashlev et al., 1996) (with heparin chromatography used instead of gel filtration as suggested). N-terminal 6xHis-tagged *E. coli* σ^{70} subunit was purified as described (Zenkin et al., 2007).

Cellular *M. smegmatis* RNAP holoenzyme with 6xHis-tag on C terminus of β ' subunit was purified as *E. coli* RNAP above from *M. smegmatis* mc²155 (SM07) (Mukherjee and Chatterji, 2008), with exception that cells were grown in MB-7H9 media supplemented with 0.05 Tween 80 and 2% glucose to late exponential phase.

In vitro transcription

One pmol of wild-type or mutant *E. coli* RNAP core and 3 pmols of σ^{70} (or one pmol of *M. smegmatis* holoenzyme) and 0.1 pmols of linear template containing T7A1 promoter (Julius and Yuzenkova, 2017; Roghanian et al., 2015) were mixed in 7 µL of transcription buffer (20 mM Tris HCl pH 7.9, 40 mM KCl, 10 mM MgCl₂). 1 µL of DMSO containing or not containing antibiotics (concentrations specified in Figures) was added to the reaction. Transcription was initiated by the addition of 2 µL of the mixture of nucleotides in transcription buffer, containing (final concentrations): 25 µM CpA or 0.5 mM ATP or 5 mM AMP with different combinations or all of 100 µM ATP, CTP and GTP and 10 µM α -[³²P]UTP (20 Ci/mmol) (Hartmann Analytic). Reactions were stopped after 10-min incubation at 37°C by the addition of formamide-containing loading buffer. Products were separated on 20% or 33% denaturing (8M urea) polyacrylamide gels, revealed by PhosphorImaging (GE Healthcare), and analyzed using ImageQuant software (GE Healthcare). Activities (a sum of run off and termination products) were normalized to the activity without antibiotic, which was taken as 100%. The average activity of at least three replicates for each concentration of antibiotic was plotted (with their standard deviations as error bars) against concentration and fitted in four parameter logistic equation by nonlinear regression using SigmaPlot. The averages and standard deviations of the three individual IC₅₀s are reported in Figure 1E.

Crystallization of RNAP – KgIA complexes

The β S531L RIF-resistant mutant *E. coli* RNAP σ^{70} holoenzyme was prepared and crystallized as previously described (Molodtsov et al., 2017). The complex of the *T. thermophilus* RNAP σ^{A} holoenzyme bound to *pyrG* promoter was prepared and crystallized as previously described (Murakami et al., 2017). To prepare complexes of RNAPs with KgIA, the RNAP crystals were soaked overnight at 22°C in the respective cryoprotection solutions supplemented with 1 mM KgIA. After the soaking, crystals were frozen by plunging to liquid N₂.

Crystallographic data collection and X-ray crystal structure determinations

The X-ray diffraction datasets were collected at the Macromolecular Diffraction at the Cornell High Energy Synchrotron Source (MacCHESS) F1 beamline (Cornell University, Ithaca, NY) and the data were processed by HKL2000 (Otwinowski and Minor, 1997). Structures were determined by molecular replacement using the structures of the *E. coli* RNAP (Murakami, 2013) and the *T. thermophilus* RNAP – *pyrG* promoter complexes (Murakami et al., 2017) as corresponding initial search models. The structures were refined as the rigid body and the positional refinement with non-crystallographic symmetry and reference model structure restraints using the suite of programs Phenix (Afonine et al., 2010). The resulting electron density maps revealed clear maps corresponding to KgIA that was not present in the initial search models, and the KgIA model was fitted into the density maps using Coot (Emsley and Cowtan, 2004). The Table 1 contains data collection and refinement statistics for the structures of the determined RNAP – KgIA complexes. The final coordinates and structure factors were deposited to the PDB with ID codes listed in Table 1.

Determination of minimum inhibitory concentrations (MICs) for M. tuberculosis

Bacterial strains and reagents

Mycobacterium tuberculosis strains: H37Rv, Beijing 1192/015 and Beijing 08/00483E (multi-drug resistant, MDR) were used in the modified resazurin microtiter assay (REMA) plate method. Beijing strains 1192/015 and 08/00483E were obtained from the PHE National Mycobacterial Reference Laboratory. Beijing 08/00483E contains a single nucleotide polymorphism (SNP) at position 761155 of $C \rightarrow T$, which has resulted in an amino acid change, S450L (corresponding to S531L with *E. coli* numbering), within the RIF-binding pocket. Beijing 08/00483E is also resistant to isoniazid, ethambutol, and pyrazinamide, through mutations in known resistance genes. Other reagents used were: Middlebrook 7H9 medium (BD Difco), OADC supplemented with 0.5% glycerol (24388.295, VWR Chemicals) and 0.2% Tween 80 (P1754, Sigma), CAMR Mycobacterium Medium MOD2 (CMM MOD2) (James et al., 2000), RIF (PanReac AppliChem, cat no.A2220) and 0.02% resazurin solution (10 mg resazurin sodium salt (R7017, Sigma) in 50 mL phosphate buffered saline, pH 7.4 (Severn Biotech) containing 5% Tween 80) (Gold et al., 2016). All the antibiotics were dissolved in methanol (322415, Sigma) to a stock concentration of 10 mg/mL.

Preparation of bacterial inocula

Bacterial strains were grown in 50 mL of OADC-supplemented, Middlebrook 7H9 medium with 0.5% glycerol and 0.2% Tween 80 in vented Erlenmeyer flasks, shaken at 37°C for 5 days. Inocula for the REMA method were prepared to an OD_{540} of 0.5, from Middlebrook 7H9 cultures by diluting cells in CMM MOD2 medium.

Modified resazurin microtiter assay (REMA) plate method

A modification of the REMA plate method previously described by (Nateche et al., 2006) was used. A 10 mg/mL stock solution of KglA was prepared in methanol (322415, Sigma). A two-fold serial dilution of the stock solution was performed in CMM MOD2 medium. 100 μ l of each KglA concentration was added to six replicate wells in a 96-well untreated flat bottomed 96-well plate (734-1554, VWR). Three of the wells remained as "drug-only" wells and the other three wells were "test wells," which were used to test the activity of KglA. 10 μ l of bacterial culture was added to the test wells to give a final OD₅₄₀ of 0.05. Three further wells contained 100 μ l of growth medium and 10 μ l of bacterial cells as "zero-drug" controls to confirm exponential growth of the bacteria in the absence of antibiotic. The drug-only controls allowed for confirmation that there was an absence of colorimetric interference from the candidate antibiotic. Bacterial cell samples were diluted to an optical density (OD₅₄₀) of 0.5 and 10 μ l of the inoculum was added to each of the test wells and the zero drug controls. 10 μ l of CMM MOD2 growth medium was added to the drug-only wells. The 96-well plates were incubated for 7 days at 37°C while shaking at 200 rpm. Drug activity was ascertained by adding 11 μ l resazurin to every well of the 96 well-plate and incubating the plate at room temperature for 6 hours. The optical density (OD₅₇₀) was measured for each well. The OD₅₇₀ measurements were converted to a percentage of the zero-drug control so that data were expressed as a percentage of the growth without antibiotic.

Statistical analyses

The average OD_{570} for each drug concentration was fitted to a sigmoidal curve using the modified Gompertz function ($y = A + Ce^{-eB(x - M)}$) and the minimum inhibitory concentration (MIC) was identified from the point of inflexion of the lower asymptote (Lambert and Pearson, 2000). This was applied to the mean of the OD_{570} of four biological repeats to produce a graphical representation for each condition. An unpaired, two tailed, Students t test was used to determine significance between biological repeats.

Determination of minimum inhibitory concentrations for other bacteria

MICs were determined using the agar dilution method specified by the British Society for Antimicrobial Chemotherapy against a collection of 56 isolates (Andrews, 2001). Briefly, KgIA was initially dissolved in a minimal volume of dimethyl sulfoxide and diluted in water before incorporation into IsoSensitest agar (Oxoid) at a final concentration range of 128 – 0.031 µg/mL. A final inoculum

of 10 000 CFU/spot was used for inoculation of plates. Cultures were incubated for 22 hours at $37 \pm 0.5^{\circ}$ C. The MIC was recorded as the lowest concentration of KgIA that achieved complete inhibition of visible growth. Inhibitor-free controls were also tested, including those containing equivalent solvent concentrations. All tests were performed at least twice on separate occasions to demonstrate reproducibility.

The collection included 16 isolates acquired from the National Collection of Type Cultures (NCTC; Colindale, UK), 3 isolates from the American Type Culture Collection (ATCC; Manassas, USA), and one from the National Collection of Pathogenic Fungi (NCPF; Colindale, UK). Also included were 36 strains of methicillin-resistant *Staphylococcus aureus* (MRSA) kindly supplied by Public Health England (Colindale, UK). These 36 strains were isolated from clinical samples in five European countries and included representatives of the major clones responsible for outbreaks of infection.

DATA AND SOFTWARE AVAILABILITY

The atomic coordinates and structure factors have been deposited in the Protein Data Bank, https://www.pdb.org (PDB ID codes 6CUX and 6CUU). CCDC 1827601 and 1827602 contain the supplementary crystallographic data for this paper, these data can be obtained free of charge from The Cambridge Crystallographic Data Centre (https://www.ccdc.cam.ac.uk/).

Molecular Cell, Volume 72

Supplemental Information

Mode of Action of Kanglemycin A, an Ansamycin

Natural Product that Is Active against

Rifampicin-Resistant *Mycobacterium tuberculosis*

Hamed Mosaei, Vadim Molodtsov, Bernhard Kepplinger, John Harbottle, Christopher William Moon, Rose Elizabeth Jeeves, Lucia Ceccaroni, Yeonoh Shin, Stephanie Morton-Laing, Emma Claire Louise Marrs, Corinne Wills, William Clegg, Yulia Yuzenkova, John David Perry, Joanna Bacon, Jeff Errington, Nicholas Edward Ellis Allenby, Michael John Hall, Katsuhiko S. Murakami, and Nikolay Zenkin

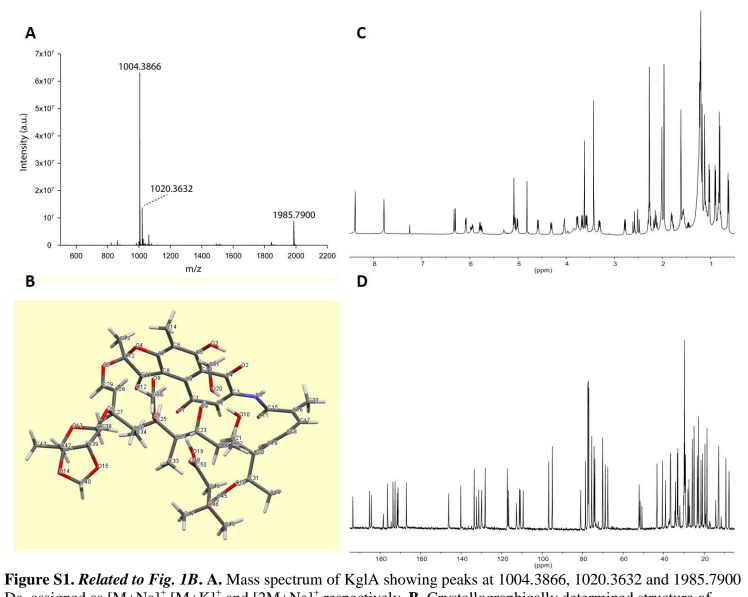


Figure S1. *Related to Fig. 1B.* **A.** Mass spectrum of KglA showing peaks at 1004.3866, 1020.3632 and 1985.7900 Da, assigned as $[M+Na]^+ [M+K]^+$ and $[2M+Na]^+$ respectively. **B.** Crystallographically determined structure of KglA (from synchrotron data). Non-H atoms are labelled. **C.** 500 MHz ¹H NMR spectra of KglA in CDCl₃. **D.** 125 MHz ¹³C{¹H} NMR spectra of KglA in CDCl₃.

Atom	δ _c				H	1 1	¹³ C- ¹ H (HMBC)
no.	- C	δ _H	Protons	Multiplicity	J (Hz)	¹ H- ¹ H (COSY)	
35	43.1	2.54	1	d	16.8	2.66 (35)	1.18 (37), 1.23 (38)
	.5.1	2.66	1	d	16.8	2.54 (35)	
36	172.0	-	-	-	-	-	2.54 (35), 2.66 (35)
34	40.4	-	-	-	-	-	2.54 (35), 2.66 (35), 1.18 (37), 1.23 (38)
38	24.6	1.23	3	s	-	-	2.54 (35), 2.66 (35)
37	25.7	1.18	3	s	-	-	2.54 (35), 2.66 (35)
33	176.0	-	-	-	-	-	2.54 (35), 2.66 (35), 1.18 (37), 1.23 (38), 5.07 (31)
31	67.7	5.07	1	"q"	6.4	1.09 (32)	3.70 (21), 1.09 (32), 5.83 (19)
32	19.3	1.09	3	d	6.4	5.07 (31)	5.07 (31), 2.17 (20)
20	52.4	2.17	1	"t"	9.7	5.83 (19), 3.70 (21)	5.97 (18), 5.83 (19), 5.07 (31), 3.70 (21), 3.52 (OH), 1.87 - 1.81 (22)
19	133.6	5.83	1	dd	15.6, 9.7	5.97 (18), 2.17 (20)	6.17 (17), 5.97 (18), 5.07 (31), 3.70 (21), 2.17 (20), 2.06 (30)
18	127.9	5.97	1	dd	15.9, 6.2	6.17 (17), 5.83 (19)	5.83 (19), 2.17 (20), 2.06 (30)
17	129.6	6.17	1	"dt"	6.2, 1.3	5.97 (18), 2.06 (30)	5.97 (18), 5.83 (19), 2.06 (30)
16	132.5	-	-	-	-	-	5.97 (18), 5.83 (19), 2.06 (30)
30	20.5	2.06	3	s	-	6.17 (17)	6.17 (17)
15	171.0	-	-	-	-	-	8.35 (NH), 6.17 (17), 2.06 (30)
21	68.5	3.70	1	d	10.1	2.17 (20)	5.83 (19), 5.07 (31), 3.52 (OH), 2.86 (23), 2.17 (20), 1.87 – 1.81 (22), 0.95 (39)
22	33.5	1.87-1.81 (overlap 45)	2 ^[a]	m	-	0.95 (39)	2.86 (23), 2.17 (20), 0.95 (39)
39	12.4	0.95	3	d	7.0	1.87 – 1.81 (22)	3.70 (21), 2.86 (23), 1.87 – 1.81 (22)
23	78.5	2.86	1	d	10.3	1.60 (24)	4.39 (25), 3.70 (21), 1.87 – 1.81 (22), 1.60 (24), 0.95 (39), 0.70 (40)
23	36.6	1.60	1	m	-	2.86 (23), 0.70 (40), 4.39 (25)	2.86 (23), 0.70 (40), 1.87 – 1.81 (22)
40	8.9	0.70	3	d	6.7	1.60 (24)	
40	0.9	0.70	5	u	0.7	1.60 (24)	4.39 (25), 2.86 (23), 1.60 (24) 3.86 (27), 2.86 (23), 2.06 (30), 0.70 (40), 0.39 (43)
25	73.7	4.39	1	d	9.9	2.13-2.08 (26), 1.60 (24)	
41	173.6	-	-	-	-	-	4.39 (25), 2.03 (42)
42	21.3	2.03	3	S	-	-	-
26	36.7	2.13-2.08	1	m	-	0.39 (43), 4.39 (25), 3.86 (27)	5.14 (28), 4.39 (25), 3.86 (27), 0.39 (43)
43	12.9	0.39	3	d	7.2	2.13-2.08 (26)	4.39 (25), 3.86 (27), 2.13-2.08 (26)
27	80.9	3.86	1	dd	9.1, 3.0	5.14 (28), 2.13-2.08 (26)	6.37 (29), 4. 66 (44), 4.39 (25), 2.13-2.08 (26), 0.39 (43)
28	112.8	5.14 (overlap 47)	2	dd	12.8, 9.1	3.86 (27), 6.37 (29)	6.37 (29), 3.86 (27),), 2.13-2.08 (26)
29	145.8	6.37	1	d	12.8	5.14 (28)	5.14 (28), 3.86 (27), 1.68 (13), 0.39 (43)
44	96.8	4.66	1	dd	8.9, 3.3	1.87 - 1.81 (45), 2.22 (45)	4.11 (46), 3.86 (27), 3.37 (49), 2.22 (45), 1.87 - 1.81 (45)
49	70.0	3.37	1	"dq"	9.0, 6.2	1.28 (50), 3.63 (48)	4.87 (47), 4.66 (44), 4.11 (46), 1.28 (50)
48	75.5	3.63	1	dd	9.0, 5.2	3.37 (49), 4.11 (46)	5.13 (47), 4.87 (47), 3.37 (49), 2.22 (45), 1.28 (50)
		5.13 (overlap 28)	2	"s"	-	-	()
47	94.9	4.87	1	"s"	-	-	3.63 (48)
46	74.3	4.11	1	"td"	4.8, 2.8	1.87 - 1.81 (45), 2.22 (45), 3.63 (48)	5.13 (47), 4.87 (47), 4.66 (44), 3.63 (48), 2.22 (45)
		2.22	1	"dt"	14.5, 2.4	1.87 – 1.81 (45), 4.11 (46), 4.66 (44)	
45	33.0	1.87-1.81 (overlap 22)	2 ^[a]	m	-	2.22 (45), 4.11 (46), 4.66 (44)	4.66 (44)
50	18.3	1.28	3	d	6.2	3.37 (49)	3.63 (48), 3.37 (49)
13	23.1	1.68	3	s		-	-
12	109.3			-		-	6.37 (29), 1.68 (13)
11	193.5	-	-	-	-	-	1.68 (13)
6	171.4	-		-		-	2.34 (14)
14	7.3	2.34	3	s	-	-	-
7	116.4		-				
	116.4	-		-	-		2.34 (14)
5 °		-	-	-	-	-	2.34 (14)
8	166.9	-	-	-	-	-	2.34 (14)
9	110.9	-	-	-	-	-	2.34 (14)
10	131.5	-	-	-	-	-	7.80 (3), 2.34 (14)
1/4	185.3	-	-	-	-	-	8.35 (NH), 7.80 (3)
1/4	184.3	-	-	-	-	-	8.34 (NH)
2	140.4	-	-	-	-	-	7.80 (3)
3	116.7	7.80	1	s	-	-	8.35 (NH)
NH	-	8.35	1	-	-	-	-
Ar-OH	-	12.60	1	-	-	-	-

Table S1. *Related to Fig. 1B*. Assignment of the ¹H and ¹³C NMR data of KglA

^[a]The integration of the signals in the 1.87 - 1.81 ppm region corresponded to 3 protons. 2 of these protons are assigned as C⁴⁵-H and C²²-H. It is possible that the remaining signal corresponds to an OH.

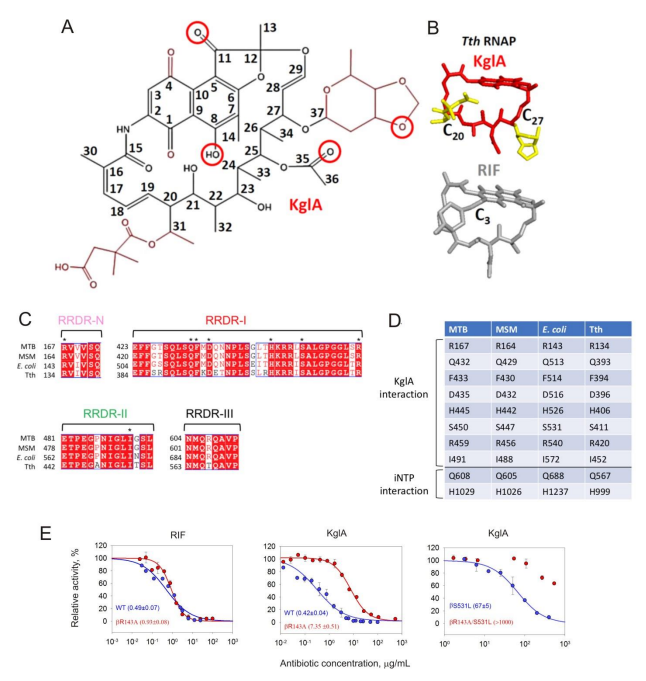


Figure S2. *Related to Fig.* **2. A**. The chemical structure of Kanglemycin A (KglA). The four oxygen atoms forming hydrogen bonds with the RIF-binding pocket of RNAP are marked by red circles. **B**. Conformations of KglA and RIF in the RIF-binding pocket of *T. thermophilus* RNAP. KglA (red) and RIF (grey) are shown as stick models; the KglA's *ansa*-bridge C20 and C27 side chains are shown in yellow and labeled; the C3 side chain of RIF is labeled. **C.** Sequence alignment spanning the RIF resistance-determining regions (RRDRs) in the β subunit (RpoB) of RNAPs from *M. tuberculosis* H37RV (MTB), *M. smegmatis* (MSM), *E. coli* and *T. thermophilus* (Tth). Amino acid residues involved in the KglA interaction (Fig. 2) and also having mutations to generate the RIF-resistant RNAPs (Figs. 1 and 3) investigated in this study are indicated by asterisks.

D. Amino acid residues of the β subunit of RNAPs involved in the KglA and initiating NTP (iNTP) interactions. **E.** Involvement of KglA-specific interaction with β R143 in KglA activity. Error bars are ±SD. The brackets contain calculated IC50±SE. IC₅₀ of KglA for double mutant was beyond our measurement limit.

Table S2. Oligonucleotides used in the study.

Digonucleotides		
0516V (GAC =>GTC)	This paper	N/A
poB_D516V_d: GCT GTC TCA GTT TAT GGT CCA GAA CAA CCC GCT G		
poB_D516V_r: CAG CGG GTT GTT CTG GAC CAT AAA CTG AGA CAG C		
5531L (TCC =>TTG)	This paper	N/A
poB S531 d: AAA CGT CGT ATC TTG GCA CTC GGC CC		
poB_S531_r: GGG CCG AGT GCC AAG ATA CGA CGT TT		
1526P (CAC =>CCG)	This paper	N/A
poB H526Y d: CTG AGA TTA CGC CGA AAC GTC GTA T	rine paper	
poB_H526Y_r: ATA CGA CGT TTC GGC GTA ATC TCA G		
1526Y (CAC =>TAC)	This paper	
poB_H526Y_d: CTG AGA TTA CGT ACA AAC GTC GTA T		
poB_H526Y_r: ATA CGA CGT TTG TAC GTA ATC TCA G		
572F (ATC =>TTC)	This paper	N/A
	This paper	IN/A
poB_I572_d: CAT CGG TCT GTT CAA CTC TCT GTC CG		
poB_I572_r: CGG ACA GAG AGT TGA ACA GAC CGA TG	This second	
Q513L (CAG =>CTG)	This paper	N/A
poB_Q513L_d: CCA GCC AGC TGT CTC TGT TTA TGG ACC		
DOB_Q513L_r: GGT CCA TAA ACA GAG ACA GCT GGC TGG		
9516Y (GAC =>TAC)	This paper	N/A
DOB_D516Y_d: GCT GTC TCA GTT TAT GTA TCA GAA CAA CCC GCT G		1.1/7
DoB_D516Y_r: CAG CGG GTT GTT CTG ATA CAT AAA CTG AGA CAG C		
1526R (CAC =>CGC)	This paper	N/A
poB_H526R_d: CTG AGA TTA CGC GCA AAC GTC GTC G	This paper	IN/A
poB_H526R_r: CGA CGA CGT TTG CGC GTA ATC TCA G	This man an	
0516V (GAC =>GTC)	This paper	N/A
poB_D516V_d: GCT GTC TCA GTT TAT GGT CCA GAA CAA CCC GCT G		
poB_D516V_r: CAG CGG GTT GTT CTG GAC CAT AAA CTG AGA CAG C		
531L (TCC =>TTG)	This paper	N/A
poB_S531_d: AAA CGT CGT ATC TTG GCA CTC GGC CC		
poB_S531_r: GGG CCG AGT GCC AAG ATA CGA CGT TT		
1526P (CAC =>CCG)	This paper	N/A
poB_H526Y_d: CTG AGA TTA CGC CGA AAC GTC GTA T		
poB_H526Y_r: ATA CGA CGT TTC GGC GTA ATC TCA G		
1526Y (CAC =>TAC)	This paper	N/A
DOB_H526Y_d: CTG AGA TTA CGT ACA AAC GTC GTA T		
DOB H526Y I: ATA CGA CGT TTG TAC GTA ATC TCA G		
572F (ATC =>TTC)	This paper	N/A
poB_I572F_d: CAT CGG TCT GTT CAA CTC TCT GTC CG	the baba	
DOB_I572F_r: CGG ACA GAG AGT TGA ACA GAC CGA TG		
(143A (CGT=>GCT)	This paper	N/A
poB_R143A_d: TGT TAT CAA CGG TAC TGA GGC TGT TAT CGT TTC CCA		,/
GCT G	`	
poB_R143A_r: CAG CTG GGA AAC GAT AAC AGC CTC AGT ACC GTT GAT	-	
AC A		
	Murakana	N1//
byrG non-template: TAT AAT GGG AGC TGG CTC TGA TGC AGG byrG template: CCT GCA TCA GAG CCC AAA ATA C	Murakami	N/A
	et al., 2017	1