Tetracenomycin X sequesters peptidyl-tRNA during translation of QK motifs

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SUMMARY

With antibiotic-resistant bacteria threatening our ability to treat common infections, new lead compounds with distinct target binding sites and limited cross-resistance are urgently needed. Natural products that inhibit the bacterial ribosome – a target for more than half of the antibiotics in use today – are a promising source of such leads and have the potential to be developed into potent drugs through structure-guided design. However, because the mechanisms of action of many of these compounds are not well understood, they are often poor candidates for a structure-based approach. Here, we use inverse toeprinting coupled to next-generation sequencing to show that the aromatic polyketide tetracenomycin X (TcmX) primarily inhibits the formation of a peptide bond between an incoming aminoacyl-tRNA and a terminal Gln-Lys (QK) motif in the nascent polypeptide. Using cryogenic electron microscopy, we reveal that translation inhibition at QK motifs occurs via an unusual mechanism involving sequestration of the 3' adenosine of peptidyl-tRNA^{Lys} in the drug-occupied nascent polypeptide exit tunnel of the ribosome. Our study provides mechanistic insights into the mode of action of TcmX on the bacterial ribosome and paves the path for the development of novel antibiotics based on a common aromatic polyketide scaffold.

INTRODUCTION

Aromatic polyketides are a large group of secondary metabolites that include various antimicrobial or antitumor agents, such as tetracycline or doxorubicin, respectively, or the lesser known tetracenomycins¹⁻⁴. Of these, TcmX (Fig. 1a) has been shown to have moderate antimicrobial activity against drug-resistant pathogenic bacteria, including methicillin-resistant Staphylococcus aureus and vancomycin-resistant enterococci⁵, and some degree of cytotoxic activity against human cell lines⁵⁻⁷. Unlike doxorubicin, with whom it shares a planar tetracyclic core, TcmX does not trigger the SOS response and does not inhibit DNA synthesis in living cells⁶. Like tetracycline⁸, however, TcmX inhibits protein synthesis in bacteria and in human lysates⁶, suggesting that the ribosome and protein synthesis are the physiological targets for this drug. Accordingly, mutating residues U2586, U2609 and U1782 of the Escherichia coli (E. coli) 23S ribosomal RNA (rRNA) confers resistance to TcmX⁶. Moreover, high-resolution cryogenic electron microscopy (cryo-EM) structures of TcmX bound to the prokaryotic 70S and eukaryotic 80S ribosomes show that the tetracyclic chromophore of TcmX stacks onto the non-canonical base pair formed by 23S rRNA residues U2586 and U1782 (E. coli numbering) within the nascent polypeptide exit tunnel⁶. The proximity of this binding site to that of macrolide antibiotics and the observation that, like these sequence-dependent translation inhibitors^{9,10}, TcmX only partially obstructs the exit tunnel, suggest that this drug is also likely to block protein synthesis in a manner that depends on the nascent amino acid context⁶. Some evidence in support of this hypothesis comes from the observation that ribosomes translating $ermBL^{11}$ or $trpL^{12}$ leader sequences in the presence of TcmX undergo sequence-dependent stalling on Leu-Lys (LK) motifs⁶. However, a comprehensive analysis is needed to assess the generality of these findings, determine the precise

mechanism by which TcmX inhibits translation, and provide a molecular basis for the development of novel ribosome-targeting antimicrobials.

RESULTS

TcmX inhibits translation at QK motifs

To determine how TcmX alters the translational pausing landscape of E. coli ribosomes in vitro, we used inverse toeprinting coupled to next-generation sequencing (iTP-seq)¹³, a profiling method that locates ribosomes on a library of mRNAs with codon resolution (Fig. 1b). Since iTPseq requires no prior knowledge of the sequences expressed, we could simultaneously monitor the translation of ~10¹² distinct short open reading frames, each featuring a stretch of 15 random codons (Extended Data Fig. 1). Translation was carried out using a reconstituted PURE translation system¹⁴ in the absence or presence of 100 µM TcmX. Following iTP-seq, we measured the frequency of amino acid motifs of various lengths in samples treated with antibiotic and in untreated samples and calculated the fold change in frequency of each motif after antibiotic treatment. For consistency, we shall define the position of all residues/codons in this work relative to the ribosomal P site (position 0), and motifs will be defined according to the range of positions they cover, for example [-1,+1] for a three-amino acid motif covering the E, P and A sites of the ribosome. When individual positions were considered, we did not observe a significant change in the frequency of amino acids between positions -2 and +1, indicating that TcmX has no detectable impact on the translation of individual codons (Extended Data Fig. 2). In contrast, we could observe a TcmX-dependent 1.9-fold change in frequency for a Gln-Lys (QK) motif at the [-1,0] position, corresponding to stalled ribosomes with a Gln codon in the E site and a Lys codon in the P site (Fig. 1c). To a lesser extent, we also observed a 1.4-fold enrichment in Pro-Gln (PO) at the [-2,-1] position, and a 1.7-fold enrichment in Lys-Cys (KC) at the [0,+1] position following drug treatment (Extended Data Fig. 2). To our surprise, sequences containing an LK motif at the [0,+1] position did not cause ribosomes to undergo substantial TcmX-dependent stalling in our iTP-seq experiments, as reported in an earlier study⁶ (Extended Data Fig. 2, Extended Data Fig. 3). This may indicate that LK must form part of a longer arrest motif, with the surrounding sequence context providing additional elements to promote ribosome stalling, as observed previously for polyproline motifs¹⁵. Thus, our iTP-seq results show that TcmX is a contextdependent inhibitor of bacterial translation that causes ribosomes to stall primarily at QK motifs.

To assess whether the sequence surrounding the QK motif affects its ability to induce TcmXdependent ribosome stalling, we calculated the fold change in the frequencies of all measured four-amino acid motifs following addition of the drug. As expected, nearly all the motifs that caused ribosomes to undergo strong translational arrest contained a OK motif at the [-1,0] position (Fig. 1d). Since PQKC was the most enriched four-amino acid motif with a low p-value $(>20-fold change; p=4.3x10^{-7})$ (Fig. 1d), we estimated the contribution of each of its constituent amino acids to stalling by analyzing all point variants of the motif present in our iTP-seq data set. While most substitutions strongly decreased stalling efficiency, positively charged residues (H,R,K) were tolerated at the -2 position, cysteine at the -1 position, and medium-sized hydrophobic (A,V,I,L) or, to a lesser extent, polar (S,T) residues at the +1 position (Fig. 1d, Fig.1e, Extended Data Fig. 4). Importantly, synonymous codons were enriched to similar extents, suggesting that the nascent amino acid sequence, rather than the mRNA or tRNA sequence, was responsible for the context-dependent response to TcmX (Extended Data Fig. 5). Using classical toeprinting^{16,17}, we then confirmed that TcmX induces ribosome stalling during translation of an mRNA encoding a MAAAPQKCAAA peptide when tRNA^{Lys} is in the P site of the ribosome, but does not prevent translation of a consecutive stretch of alanine codons (Fig. 2a). Overall, these observations reveal the importance of the nascent amino acid sequence surrounding the core QK motif for TcmX-dependent ribosome stalling.



Fig. 1 | **iTP-seq analysis of the context dependence of ribosome inhibition by TcmX. a,** Chemical structure of TcmX. **b**, Overview of iTP-seq, with a schematic showing the range of inverse toeprints retained after gel electrophoresis. **c**, Heatmap showing the enrichment of motifs at position [-1,0] (E and P sites) after treatment with TcmX. Enrichment is defined as $log_2(F_{TcmX}/F_{untreated})$, where F_{TcmX} is the mean frequency of occurrence of a given motif in the samples treated with TcmX and $F_{untreated}$ is its mean frequency in the untreated sample. **d**, Volcano plot of statistical significance (-log₁₀(p-value)) against enrichment (log₂(fold change)) for four-amino acid motifs at position [-2,+1]. The majority of highly enriched and statistically significant motifs feature a QK motif at position [-1,0] (yellow points), though a few other motifs were also found (blue points). Thresholds of log₂(fold change) and -log₁₀(p-value) of 1 and 5, respectively, were used to highlight such motifs. **e**, Analysis of single amino acid variants of the PQKC motif, where $log_2(fold change)$ in motif frequencies relative to the original motif are shown using a gray scale. Red crosses denote variants that are not sufficiently represented in the iTP-seq data. The values shown in c, d and e are the mean from three independent experiments.

TcmX hinders QK translation in vivo

To determine whether TcmX-dependent translational arrest at PQKC motifs occurs in vivo, we performed a β -galactosidase complementation assay that monitors $lacZ\alpha$ expression as a proxy for ribosome stalling (Fig. 2b). In the original form of this assay¹⁸, expression of $lacZ\alpha$ is controlled by a well-characterized translation attenuation mechanism involving the antibiotic-dependent stalling of a ribosome translating the leader open reading frame (ORF) ermCL¹⁹⁻²¹. In the absence of added antibiotic, cells express *ermCL* constitutively, but $lacZ\alpha$ is translationally repressed due to sequestration of its ribosome binding site within an mRNA hairpin. Addition of subinhibitory concentrations of the macrolide antibiotic erythromycin (ERY) causes the ribosome translating ermCL to stall on the ninth codon of this leader ORF. This, in turn, leads to a rearrangement of the mRNA secondary structure within the ermCL-lacZ α intergenic region, which frees the lacZ α ribosome binding site, enabling the production of α -fragment. As previously shown²², ERYinduced ribosome stalling on ermCL was manifested by the appearance of a blue halo surrounding the filter disk soaked with this drug, whereas no such halo was observed for untreated cells (Fig. 2b). When, instead, TcmX was added to the disc, only a very weak halo was observed, indicating that TcmX treatment resulted in marginal ribosome stalling on ermCL (Fig. 2b). Next, we replaced the first nine codons of ermCL with a nucleotide sequence coding for a MAAAPQKC peptide, thus preserving the ermCL-ermC intergenic region necessary for $lacZ\alpha$ induction, as well as the expected position of the TcmX-dependent stalling site observed by toeprinting. When cells carrying this construct were left untreated or were treated with subinhibitory concentrations of ERY, no $lacZ\alpha$ induction was observed (Fig. 2b). However, treatment with TcmX resulted in the appearance of a strong blue halo in the disc-diffusion assay, indicating

that this antibiotic causes ribosomes that reach the PQKC motif to stall, resulting in $lacZ\alpha$ expression (Fig. 2b). Together, these results demonstrate that the context-dependent action of TcmX on QK motifs observed *in vitro* also occurs *in vivo*.



Fig. 2 | *In vitro* and *in vivo* validation of TcmX-dependent ribosome stalling at a PQKC motif. **a**, Toeprinting assay showing the TcmX-dependent stalling of ribosomes translating a MAAAPQKCAAA* sequence. Stalling occurred when the K codon was positioned in the P site (black arrow). The antibiotic retapamulin was used as a control to indicate the position of ribosomes stalled on the start codon (white arrow). Ribosomes that reached the stop codon also gave rise to a toeprint corresponding to a termination complex (gray arrow). This experiment was repeated three times independently with similar results. **b**, β -galactosidase complementation assay to monitor *lacZa* expression as a proxy for ribosome stalling. Blue halos result from *lacZa* expression induced by drug-dependent ribosome stalling on the *ErmCL*₁₋₁₀ or MAAAPQKC leader ORFs. This experiment was repeated three times independently with similar results.

TcmX sequesters the 3' end of P-tRNA

To understand the structural basis of TcmX-dependent translation inhibition at QK motifs, we determined the cryo-EM structure of an E. coli 70S ribosome stalled during translation of the PQKC-containing toeprinting template in the presence of 100 µM TcmX, at an overall resolution of 2.7 Å (Fig. 3a, Extended Data Fig. 6). The structure, which we will refer to as 70S-MAAAPQKC-TcmX, was obtained from a major class of particles (48.1%) containing tRNA^{Cys} and tRNA^{Lys} in the ribosomal A and P sites, respectively, as confirmed by the presence of tRNA-specific modifications (Extended Data Fig. 7). A less clearly defined and seemingly aminoacylated tRNA (modeled as tRNA^{Gln2}) was also present in the E site, indicating that deacylated tRNA^{Gln2} must first have dissociated from and subsequently been replaced with aminoacyl-tRNA in the ribosomal E site during sample preparation (Extended Data Fig. 7). Local resolution extending to 2.67 Å in the core of the particle enabled us to accurately model all the major ligands including TcmX, peptidyl-tRNA^{Lys} and aminoacyl-tRNA^{Cys} (Fig. 3b, Extended Data Fig. 7). As previously reported⁶, TcmX was bound to the nascent polypeptide exit tunnel (Fig. 3a,b), where it stacks against the noncanonical U2586:U1782 base pair. In the A site, we could observe clear density for a cysteinyl moiety attached to tRNA^{Cys} (residue C⁺¹) (Fig. 3b). The bases of 23S rRNA residues U2506, G2583 and U2584 were in their induced conformation, but 23S rRNA residue U2585 and residue A76 of the P-site tRNA remained in the uninduced conformation²³, indicating only partial aminoacyl-tRNA accommodation into the peptidyl transferase center (Extended Data Fig. 8). In the P site, clear density was also visible for all but the first two residues of the nascent peptide (A⁻⁴A⁻³P⁻²Q⁻¹K⁰) (Fig. 3b), confirming that the last amino acid incorporated into the nascent peptide was a lysine residue, in agreement with our iTP-seq and toeprinting data.



Fig. 3 | **Cryo-EM structure of a 70S-MAAAPQKC-TcmX complex. a,** Transverse section of the cryo-EM map for the 70S-MAAAPQKC-TcmX complex with isolated densities showing the ribosome (light gray), peptidyl-tRNA (blue) and TcmX (orange). **b**, Molecular model and surrounding map densities for the MAAAPQK nascent chain (red), P-site tRNA (dark blue), A-site tRNA (light blue), incoming residue C⁺¹ (salmon) and TcmX (orange). **c**, Residue A76 of the P-site tRNA forms a non-canonical *trans* Hoogsteen–Hoogsteen base pair with residue A2439 of the 23S rRNA. **d**, Residue K⁰ from the nascent peptide occupies the A-site crevice, where it is likely to interfere with the accommodation of some but not all aminoacyl-tRNAs. **e**, The side chain of residue Q⁻¹ from the nascent peptide stacks against the base of 23S rRNA residue A2062. It also forms a hydrogen bond with the backbone carbonyl oxygen of residue -3, which is likely to be facilitated by the kink induced in the nascent peptide by residue P⁻².

Unexpectedly, residue A76 of the P-site tRNA was rotated and shifted by nearly 12 Å compared to the pre-attack state, in which peptidyl-tRNA and aminoacyl-tRNA are poised to undergo peptide bond formation²⁴. This atypical conformation of residue A76 is distinct from those observed previously in structures of the ribosome in complex with antibiotics that perturb the position of the 3' end of peptidyl-tRNA, such as blasticidin S^{25,26} or hygromycin A²⁷. Several elements stabilize the highly unusual peptidyl-tRNA conformation induced by TcmX, most notably a trans Hoogsteen-Hoogsteen base pair between residue A76 and 23S rRNA residue A2439 (Fig. 3c). On the peptide side, residue K⁰ occupies the A-site crevice, where its backbone carbonyl oxygen forms a hydrogen bond with the amine of 23S rRNA residue G2061 (Fig. 3d). The side chain of residue Q⁻¹ stacks against the base of 23S rRNA residue A2062 (Fig. 3e). Mutating the residue at position -1 to the shorter asparagine would likely result in suboptimal stacking with A2062, explaining why this conservative substitution is not enriched in our iTP-seq dataset. The side chain and backbone amide groups of residue Q⁻¹ also form intramolecular hydrogen bonds with the backbone carbonyl of residue A⁻³ (Fig. 3d) and with TcmX (Fig. 3b), respectively. Finally, residue P⁻² forms a kink within the nascent peptide that may favor its interaction with the drug (Fig. 3d). However, our iTP-seq data show that proline is not strictly required at this position, even though it may contribute to stalling efficiency. Thus, the structure of the 70S-MAAAPQKC-TcmX complex observed by cryo-EM features a highly disrupted peptidyl transferase center stabilized by a network of interactions between the drug, the QK-containing peptidyl-tRNA and the ribosome.

Mechanism of ribosome inhibition by TcmX

Under physiological conditions, the α -amino group of an incoming aminoacyl-tRNA is deprotonated to yield an $-NH_2$ nucleophile capable of attacking the ester bond connecting the

C-terminal amino acid of the nascent peptide to the P-site tRNA. For peptide bond formation to occur, the attacking nucleophile must be located within a relatively short distance (~3 Å) of the carbonyl ester carbon (Fig. 4a)²³. In our structure, the distance between the α -amine of the incoming cysteine (C⁺¹) and the carbonyl ester carbon of the terminal lysine (K⁰) of the nascent peptide is increased to 5.2 Å, and the geometry of the reactants is highly unfavorable for peptidyl transfer (Fig. 4b). In addition, the α -amine of residue C⁺¹ appears to form a salt bridge with the backbone phosphate of tRNA^{Lys} residue A76 in the P site, suggesting that its protonation state (– NH₃⁺) renders it unable to perform a nucleophilic attack (Fig. 4b). The poor placement of the reactants and the lack of a –NH₂ nucleophile resulting from the likely protonation of the α -amino group are thus likely to account for the inability of the ribosome to form a peptide bond after QK motifs in the presence of TcmX.



Fig. 4 | **Mechanism of TcmX-dependent translation inhibition at QK motifs. a,** Pre-attack complex (PDB 1VY4) where phenylalanyl-tRNA^{Phe} in the A site is poised to attack formyl-methionyl-tRNA^{Met} in the P site. This complex, which was obtained using a non-reactive form of formyl-methionyl-tRNA^{Met} containing a non-hydrolyzable amide bond instead of the usual ester bond shows that an attacking nucleophile positioned 2.4 Å away from the ester carbonyl carbon. **b**, In the 70S-MAAAPQKC-TcmX complex, sequestration of residue A76 of the P-site tRNA into the nascent polypeptide exit tunnel of the ribosome increases the distance between the attacking amine and the ester bond to 5.2 Å, making peptide bond formation less likely. Moreover, a salt bridge between the attacking amine and the backbone phosphate of P-site tRNA residue A76 renders protonation of the α -amino group highly likely.

CONCLUSION

In this work, we reveal the principal mechanism by which the recently discovered antibiotic TcmX inhibits bacterial translation upon binding to the nascent polypeptide exit tunnel of the ribosome. By using iTP-seq to characterize the stalling landscape of TcmX-bound *E. coli* ribosomes, we showed that TcmX inhibits translation at QK motifs, in a manner that is dependent on the nascent amino acid sequence. Using complementary biochemical and structural approaches, we identified how a nascent peptide featuring a C-terminal QK motif interacts with the bacterial ribosome and TcmX to inhibit peptide bond formation *in vitro* and *in vivo*.

The mechanism of action of TcmX is reminiscent of that by which macrolide antibiotics inhibit the ribosome²⁸. Indeed, both macrolides and TcmX inhibit translation in a peptide sequence-dependent manner²⁸. In the presence of macrolides, translational arrest occurs primarily at +X+ motifs, where + represents a positively charged amino acid like arginine or lysine²⁹. Both TcmX and macrolides bind near the peptidyl transferase center, albeit on opposite faces of the nascent

polypeptide exit tunnel^{6,28}, explaining the lack of cross-resistance to TcmX and macrolides⁶. These different binding sites and sequence specificities ultimately result in substantially distinct mechanisms of action. On a macrolide-bound ribosome, nascent peptides containing a +X+ motif adopt a conformation where the first positively charged residue within the motif (typically an arginine at position -1) occupies the A-site crevice and prevents the accommodation of an incoming positively charged amino acid into the A site, leading to ribosome stalling²⁹. In contrast, translation of QK motifs in the presence of TcmX appears to only mildly impair aminoacyl-tRNA accommodation, but instead results in the sequestration of the peptidyl-tRNA 3' end inside the nascent polypeptide exit tunnel, and subsequent inhibition of peptide bond formation. Other tetracenomycins, such as elloramycin or tetracenomycin C, exert antimicrobial activity against a variety of *Actinomyces* and *Streptomyces* species³⁰⁻³⁴ and may operate through similar context-dependent mechanisms. Elucidation of the modes of action of this very promising family of natural products could open new avenues for the development of highly specific narrow-spectrum antibiotics to treat multidrug-resistant infections.

METHODS

General experimental procedures for iTP-seq

DNA and RNA products at various points of the iTP-seq protocol were analyzed on 9% acrylamide (19:1) TBE (90 mM Tris, 90 mM boric acid, and 2 mM EDTA) gels and stained with SyBR Gold (Invitrogen). Inverse toeprints were excised from 12% acrylamide TBE gels using a clean scalpel. RNA gel electrophoresis was performed under denaturing conditions (8 M urea in the gel). All reactions were performed using molecular biology grade water (Millipore). Reagents and oligonucleotides used in this study are listed in Supplementary Table 1.

Generation of an (NNN)₁₅ template library

A DNA library composed of short ORFs with a random N-terminal stretch of 15 NNN (aNy aNy, aNy) codons was purchased from Eurogentec as a single stranded oligonucleotide (*iTP_NNN15_random-library*). An expression cassette for *in vitro* transcription and translation was obtained by polymerase chain reaction (PCR) with Phusion DNA polymerase (6 cycles [98°C, 10 s; 62°C, 5 s; 72°C, 10 s]), using the *iTP_NNN15_random-library* oligonucleotide in combination with oligonucleotides *iTP/TP_T7_RBS_ATG_f* and *iTP_Stop_EcoRV_r* as templates (0.1 pmol of each oligonucleotide per 50 µl reaction) and oligonucleotides *iTP/TP_T7_f* and *iTP_EcoRV_r* for amplification (1 pmol of each oligonucleotide per 50 µl reaction). The PCR product was purified using a PCR purification kit (Qiagen) according to the manufacturer's instructions and quantified with a 2100 Agilent Bioanalyzer. The sequence of the final (NNN)₁₅ template was:

(the T7 promoter is underlined, the translated region is in bold; the EcoRV site for iTP-seq is in italics).

In vitro transcription

In vitro transcription was performed at 20°C for 3 h, in a 200 µl reaction volume containing 8 pmol of $(NNN)_{15}$ library, 3.7 µl of 20 mg/ml stock of homemade T7 RNA polymerase (Addgene plasmid p6XHis-T7-P266L³⁵), 80 mM Tris–HCl pH 7.6, 24 mM MgCl₂, 2 mM spermidine, 40 mM DTT, 7.5 mM ATP (Sigma Aldrich), CTP and UTP, 0.3 mM GTP (CTP, UTP, and GTP from Jena Bioscience), and 4.5 mM Thio-Phosphate-GMP (Genaxxon Bioscience). Transcripts were purified using an RNA Clean & Concentrator-5 purification kit (Zymo Research) according to the

manufacturer's instructions. The concentration of mRNA was determined using a NanoDrop spectrophotometer (Thermo Fisher Scientific).

Biotinylation

Biotin-maleimide (Vectorlabs) was dissolved in dimethylformamide according to the manufacturer's instructions. 800 pmol mRNA were mixed with 800 nmol biotin-maleimide in 100 mM Bis-Tris-acetate buffer pH 6.7 in a total volume of 50 µl and incubated at room temperature for 2.5 h. Unincorporated biotin was removed by washing the mRNA three times with H₂O (molecular biology grade, Millipore) in an Amicon membrane centrifugal concentrator with a molecular weight cutoff (MWCO) of 30 kDa (Millipore). mRNA was recovered and the biotinylation efficiency was determined by means of a dot blot, as follows. H+ bond membrane (GE Healthcare) was treated with 6Å~ SSC buffer (900 mM NaCl, 90 mM Na₃-citrate, pH 7.0) for 10 min and dried briefly between two pieces of Whatman paper. Biotinylated (NNN)₁₅ mRNA and a 5'-biotinylated oligonucleotide standard (Biotin standard) were diluted in 6x SSC buffer to 0.5, 1.0, 2.5, and 5.0 μ M, and 1 μ l of each dilution was pipetted onto the prepared membrane. The membrane was baked for 2 h at 80°C to adsorb nucleic acids, and subsequently blocked in 2.5% dry milk solution in TBS-T (50 mM Tris-HCl, 150 mM NaCl, and 0.05% [vol/vol] Tween-20, pH 7.5) for 1 h at room temperature. The milk solution was removed and the membrane was incubated with a 1:1,000 dilution of streptavidin-alkaline phosphatase antibody (Promega) in TBS-T for 1 h at room temperature. Unbound antibody was removed by washing three times with TBS-T buffer. Colorimetric detection was performed using an NBT/BCIP detection kit (Promega) according to the manufacturer's instructions. The membrane was imaged immediately on a Bio-Rad Imager. The biotinylation efficiency was estimated by comparing the intensity of the sample dots with the intensity of the standard dots.

Polyadenylation of the mRNA

Polyadenylation of the biotinylated mRNA was performed using Poly-A polymerase (New England Biolabs) using the buffer supplied. The ratio of mRNA to ATP molecules was chosen to be 1:100. The reaction was incubated at 37°C for 2 h and the efficiency of the polyadenylation reaction was assessed by denaturing polyacrylamide gel electrophoresis (PAGE) (9%). Polyadenylated mRNA was purified using the RNA Clean & Concentrator-5 kit (Zymo Research) according to the manufacturer's instructions.

Inverse toeprinting

Inverse toeprinting was performed as described previously, with modifications¹³. Briefly, *in vitro* translation was carried out with a PURExpress Δ RF-123 Δ Ribosomes kit (New England Biolabs), using ~5 pmol of 5'-biotinylated and 3'-polyadenylated mRNA as a template. TcmX dissolved in dimethyl sulfoxide (DMSO) was supplemented at a final concentration of 100 µM in 5 µl reactions. Release factors 1 and 3 were added to the translation reaction according to the manufacturer's instructions. Translation was performed at 37°C for 30 min, after which the samples were placed on ice and 5 µl ice-cold Mg²⁺ buffer (50 mM Hepes-KOH, 100 mM K-glutamate, 87 mM Mg-acetate, and 1 mM DTT, pH 7.5) was added to the reactions, thereby increasing the Mg²⁺ concentration to 50 mM. 1 µl of RNase R (1 mg/ml) was added, followed by an additional incubation for 30 min at 37°C to ensure complete mRNA degradation. 139 µl of 1x BWT buffer was added to stop the reaction (5 mM Tris–HCl, 0.5 mM EDTA, 1 M NaCl, and 0.05% [vol/vol] Tween-20, pH 7.5).

mRNA purification and linker ligation

For each sample, 5 μ l of M-280 streptavidin Dynabead (Thermo Fisher Scientific Scientific) were washed three times with 1x BWT buffer in DNA loBind tubes (Eppendorf) and resuspended in 50

 μ I of the same buffer. Dynabeads and RNA from the previous step were combined into these tubes and incubated on a tube rotator for 15 min at room temperature to allow binding of the biotinylated mRNA to the streptavidin beads. After incubation, beads were collected using a magnet and the supernatant was discarded. The beads were washed one time with 1x BWT buffer to remove unbound RNA, followed by two washes with H₂O to remove the 1x BWT buffer. Beads were resuspended in 9.5 μ I of linker ligation reaction mixture containing 4 μ I of water, 1 μ I of 10x T4 RNA ligase 2 (truncated) buffer (New England Biolabs), 3 μ I of 50% PEG 8,000 (New England Biolabs), 1 μ I of *iTP_3'_linker_ApoI* (10 μ M) and 0.5 μ I of ligase (T4 RNA ligase 2, truncated - 200 000 U/mI - New England Biolabs) per reaction. Linker ligation was allowed to proceed on a tube rotator for 2.5 h at room temperature.

Reverse transcription

Following ligation of the linker, beads were washed once with H₂O to remove unincorporated linker oligonucleotide and were resuspended in 18.5 μ l of reverse transcription reaction mixture containing 11.5 μ l of water, 1 μ l of dNTPs (10 mM of each – New England Biolabs), 1 μ l of *iTP_Linker_r* oligonucleotide (2 μ M), 4 μ l of first strand synthesis buffer (5X - Thermo Fisher Scientific) and 1 μ l of DTT (0.1 M - Thermo Fisher Scientific) per reaction. The samples were incubated for 5 min at 65°C to anneal the primer to the complementary sequence and then placed on ice. 1 μ l of reverse transcriptase (Superscript III – 200,000 U/ml – Thermo Fisher Scientific) was added to each tube and the samples were incubated for 30 min at 55°C in a Thermomixer at 500 rpm to allow reverse transcription of the Dynabead-bound mRNA.

PCR on cDNA, restriction digestion

Reverse transcribed cDNA was used without further purification as a template for PCR. To generate double stranded DNA for restriction digestion, a fill-up reaction was performed using *iTP_cDNA_f* oligonucleotide and the reverse transcribed cDNA (10 s denaturation, 10 s annealing at 42°C, and 30 s elongation at 72°C). The resulting double stranded DNA was combined with 1 µl of EcoRV-HF restriction enzyme and the sample was incubated at 37°C for 1 h. To amplify undigested DNA, *iTP_Linker_r* oligonucleotide was added, and 10-16 cycles of PCR were performed (denaturation at 98°C for 10 s, annealing at 60°C for 10 s, and elongation at 72°C for 10 s). The number of PCR cycles was adjusted to give a visible band on the gel while minimizing non-specific byproducts.

Purification of DNA fragments of interest after PCR

Bands containing inverse toeprints corresponding to stalled ribosomes from the initiation codon to the last NNN codon were excised from the gel with a clean scalpel. Gel pieces were crushed through a 5 ml syringe into 15 ml Falcon tubes and 10 ml of gel elution buffer (10 mM Tris–HCl, pH 8.0, 500 mM Na-Acetate, and 0.5 mM Na-EDTA) were added. The tubes were incubated on a tube rotator at room temperature overnight. Gel debris were separated from the extraction solution by filtering through 0.22 μ m centrifugal filters (Millipore). Each sample was then concentrated to ~1 ml using a SpeedVac. DNA was precipitated in 5 ml Eppendorf tubes using 1 ml of isopropanol with 3.7 μ l GlycoBlue (Thermo Fisher Scientific Scientific) and incubating at ~80°C overnight. After precipitation, DNA was recovered by centrifugation in a ThermoScientific Heraeus Multifuge X3R centrifuge at 20,000xg for 30 min at 4°C using a Fiberlite F15- 8x50cy rotor (ThermoScientific). The supernatant was removed, and DNA pellets were resuspended in 20 μ l H₂O (molecular biology grade, Milipore) for subsequent addition of the next-generation sequencing (NGS) adaptors.

Addition of NGS adaptors to amplified DNA

Long NGS adaptor oligonucleotides (NGS_adaptor_f and the reverse oligonucleotides

NGS_adaptor_index_number) contain Illumina TruSeq adapter sequences followed by 18 nucleotides complementary to the 5' or 3' region of the cDNA. The reverse oligonucleotides also contain barcode sequences for multiplexing according to the TruSeq v1/v2/LT protocol (Illumina). Sequencing libraries were obtained from 12–16 cycles of PCR using 0.02 μ M long NGS adapter oligonucleotides (forward and reverse) and 0.2 μ M short amplification oligonucleotides (*NGS_f* and *NGS_r*). PCR products were purified using a Qiagen PCR purification kit. The size and concentration of the fragments obtained were measured using a 2100 Agilent Bioanalyzer with the DNA 1000 kit.

Next generation sequencing

Next generation sequencing was performed by the BGI Facility in Hong Kong, on an Illumina Hiseq X Ten system in rapid run mode with 150 PE reads.

Analysis of iTP-seq data

Unless indicated otherwise, data analysis was carried out using a series of custom Python scripts. Read pairs were assembled using PEAR v0.9.10³⁶, with the maximal proportion of uncalled bases in a read set to 0 (–u option) and the upper bound for the resulting quality score set to 126 (–c option). The 5' flanking region was defined as GTATAAGGAAGGAAAAAAT and the 3' flanking region was GGTATCTCGGTGTGACTG. A maximum of two mismatches within each of these flanking regions was tolerated, whereas all other reads were discarded. Trimming of the retained reads resulted in sequences with a start codon directly at the 5' end and the site of RNase R cleavage at the 3' end (Extended Data Fig. 1). Reads were filtered using a custom script (parse_filter_fastq.py) to have a minimum quality score of 30 at all positions, contain no Ns, and be devoid of aberrations (e.g. known contaminants or abnormally long stretches of A) and inverse toeprints were aligned relative to the ribosomal P site. The data underlying the heatmaps and volcano plots were then analyzed using another custom script (make_report.py) to count the number of reads per motif for various combinations of positions, combined with DESeq2³⁷ to compute the enrichment statistics. The graphs were produced with Matplotlib 3.3.2³⁸. The logo of the top arrest motifs was generated with Logomaker³⁹.

Toeprinting assays

Toeprinting was performed as described previously¹⁷. Briefly, a DNA template containing a T7 promoter, a ribosome binding site, a sequence coding for the MAAAPQKCAAA peptide and the NV1 sequence²¹ was generated by PCR using as templates oligonucleotides iTP/TP_frag1_T7_RBS_ATG_f, TP_frag2_r, TP_frag2_NV1_r and TP_MAAAPQKCAAA*_f (0.1 pmol of each oligonucleotide per 50 µl reaction), and the short primers *iTP/TP_frag1_T7_f* and TP_short_r for amplification (1 pmol of each oligonucleotide per 50 µl reaction). The sequence of the TP_MAAAPQKCAAA* toeprinting mRNA template was:

CGATCGAATTC<u>TAATACGACTCACTATA</u>GGGCTTAAGTATAAGGAGGAAAAAAT**ATGGCAG CAGCACCGCAGAAGTGTGCAGCAGCATGA**AGCGAATAATAACTGACTCTGAACAACATCC *GTACTCTTCGTGCGCAGGCAAGG*TTAATAAGCAAAATTCATTATAACC

(the T7 promoter is underlined; the coding region is in bold; the NV1 sequence is in italics).

DNA templates were transcribed and translated *in vitro* using the PURExpress Δ RF-123 Δ Ribosomes Kit (New England Biolabs). Ligands were dissolved in water and added as needed at the beginning of the reaction. The Yakima Yellow-labelled probe complementary to the NV1 sequence (*TP_RT_yakimayellow_r*) was added to the 5 µl reaction after incubating for 15 min at 37°C (2 µM) and the sample was incubated for another 5 min at the same temperature. Reverse transcription was then performed with 50 U of Avian Myeloblastosis Virus reverse transcriptase (Promega Corporation) for 20 min at 37 °C. RNA was degraded by adding 0.5 µl of a 10 M NaOH

stock at 37 °C for 15 min. Samples were neutralized with 0.7 μ l of a 7.5 M HCl stock and the remaining complementary DNA was purified using a nucleotide removal kit (QIAGEN). Sequencing reactions were performed using a commercial kit designed to be used with fluorescent dye-labeled primers (Thermo Fisher Scientific Scientific). ExoSAP-IT reagent (Thermo Fisher Scientific Scientific) was used to remove excess dNTPs and primers from the PCR product prior to the sequencing reaction. For the sequencing procedure, 4 μ l of purified PCR product (or approximatively 0.5 – 1 pmol of DNA) and 2 pmol of the 5'-labeled oligonucleotide were used to prepare a master reaction following the manufacturer's instructions. The following thermocycler program was used for the sequencing reaction: 30 s denaturation at 95°C; 15 s annealing at 50°C; 25 cycles of 60 s for elongation at 72°C. 2 μ L of formamide loading dye from the kit were then added to each Sanger reaction, samples were heated for 3 min at 75°C to denature the cDNA, while the toeprinting samples were denatured at 95 °C for 5 min. 3.5 μ l of the sequencing reactions and 3 μ l of the toeprinting reactions were separated by 7.5% sequencing PAGE (2,000 V, 40 W for 2–2.5 h) followed by detection on an Amersham Typhoon Gel and Blot Imaging System (GE Healthcare Life Sciences).

β-galactosidase complementation assay

To test for *in vivo* activity, a pERMZ α^{18} plasmid containing the *ermCL-ermC* operon in frame with the lacZ α reporter was used. A sequence coding for the MAAAPQKC peptide was inserted in place of the first ten codons of the ermCL sequence using the QuikChange Lightning Site-Directed Mutagenesis Kit (Agilent Technologies, Inc). Briefly, oligonucleotides (pZa_MAAAPQKC_fwd) and $(pZ\alpha_MAAAPQKC_rvs)$ were used to mutate the pERMZ α plasmid according to the manufacturer's instructions. Both the pERMZ α and pERMZ α -MAAAPQKC plasmids were transformed into E. coli JM109 from which the acrAB operon had been deleted to reduce drug efflux^{40,41}. Cells were grown in lysogeny broth (LB) at 37°C (180 rpm) with ampicillin (100 µg/ml) until they reached an optical density of 0.5 at 600 nm. 2 ml of the cell culture were added to 8 ml of 0.6% LB-agar at 50°C and plated onto 1.5% LB agar plates (120 mm diameter) supplemented with 0.5 mM ampicillin, 0.5 mM Isopropyl-β-D-1-thiogalactopyranoside (IPTG) and 0.5 mM 5-bromo-4-chloro-3-indolyl-beta D-galactopyranoside (X-gal). Once the soft agar had solidified, 6-mm-diameter Whatman paper discs (GE Healthcare) were placed on top of the plate and wetted with 5 µl of a solution at 10 mM of TcmX diluted in DMSO or 300 µg of erythromycin diluted in 10 µl of ethanol. The plates were then incubated at 30°C overnight and pictures were taken the next day.

Preparation of an E. coli 70S-MAAAPQKC-TcmX complex for cryo-EM

Stalled ribosomal complexes for cryo-EM were obtained by translating the PQKC-containing template used for toeprinting *in vitro*, in a reaction containing 2.2 μ M of reassociated *E. coli* 70S ribosomes, 100 μ M TcmX, 5 pmol of *TP_MAAAPQKCAAA** mRNA (see "Toeprinting assays" section) and components of the PURExpress Δ Ribosomes Kit (New England Biolabs). The reaction was incubated at 37°C for 20 min and then diluted in 50 mM Hepes KOH pH 7.5, 100 mM K-Acetate, 25 mM Mg-Acetate, 10 μ M TcmX to yield a final ribosome concentration of 300 nM. Quantifoil carbon grids (QF-R2/2-Cu) were coated with a thin carbon layer (~2 nm) using a Safematic ccu-010 HV carbon coater. Grids were glow-discharged for 30 s at 2 mA before application of 3.5 μ l of 70S-MAAAPQKC-TcmX complex. After blotting for 2.5 s and waiting for 30 s (blotting force 5), grids were plunge-frozen in liquid ethane using an FEI Vitrobot Mark IV (Thermo Fisher Scientific) set to 4 °C and 100% humidity.

Cryo-EM data acquisition

Cryo-EM images were collected in counting mode on a Talos Arctica (Thermo Fisher Scientific) operated at 200 kV and equipped with a K2 Summit direct electron detector (Gatan) in

Nanoprobe mode at the IECB in Bordeaux (France). Images were recorded with SerialEM with a magnified pixel size of 0.93 Å at a magnification of 45,000x to record 38 movies frames with an exposure time of 3.8 seconds using a dose rate of 0.94 e⁻/Å² per frame for a total accumulated dose of 35.72 e⁻/Å² (Supplementary Table 2). The final dataset was composed of 7,450 micrographs with defocus values ranging from -0.4 to -2.5 µm.

Cryo-EM image processing

Data were processed in Relion v4.0⁴² according to the scheme presented in Extended Data Fig. 6. Briefly, the raw movie frames were summed and corrected for drift and beam-induced motion at the micrograph level using the Relion v4.0⁴² implementation of MotionCor2⁴³. The resolution range of each micrograph and the contrast transfer function (CTF) were estimated with CTFFIND v4.1⁴⁴. Well-defined two-dimensional classes were selected by subsequent rounds of twodimensional classification of the particles obtained by automated picking in Relion v4.0⁴². Threedimensional classification was performed in Relion v4.042 in three steps: (1) unsupervised classification with particles downsized four times; (2) focused classification with background subtraction using a mask covering all three tRNA sites and the GTPase binding site on particles downsized twice; and (3) focused classification with background subtraction using a mask covering the A site on particles downsized twice. Classes containing tRNAs in the A, P and E sites were selected for automated 3D reconstruction and CTF refinement in Relion v4.042, followed by Bayesian polishing. The final reconstruction was sharpened by applying a negative sharpening B-factor of -10 in Relion v4.0⁴². The resolution for the electron density map was estimated using the gold standard criterion (FSC=0.143) resulting in a final reconstruction with a nominal resolution of 2.7 Å. Local-resolution estimation was performed using Relion v4.0⁴². The pixel size was optimized on the basis of model-to-map correlation in Chimera v1.16⁴⁵.

Atomic model building and refinement

An initial model of the 70S-TcmX complex was obtained by placing the coordinates of an *E. coli* 70S ribosome (PDB: 6TBV) and of the TcmX antibiotic (PDB: 6Y69) into an auto-sharpened map obtained with Phenix v1.20.1⁴⁶. The nascent chain, mRNA and A-, P- and E-site tRNAs were modeled *de novo* into the corresponding density using Coot v0.9.7⁴⁷. The model was refined using Isolde v1.3⁴⁸ and the real space refinement procedure in Phenix v1.20.1⁴⁹, with reference model restraints.

Figure preparation

Plots were produced with Python3 and Matplotlib 3.3.2³⁸. An auto-sharpened map from Phenix v1.20.1⁴⁶ was used to prepare all figures showing cryo-EM density except Extended Data Fig. 6 and 7, for which maps obtained directly from Relion v4.0⁴² were used. Figures showing cryo-EM density or atomic models were prepared using ChimeraX v.1.3⁵⁰ or PyMOL v.2.5.0 (Schrödinger).

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AUTHOR CONTRIBUTIONS

C.A.I. designed the study. E.C.L. performed iTP-seq and toeprinting experiments. E.C.L. and T.T.R. processed and analyzed the iTP-seq data. T.N.P. and T.T.R. performed the blue ring assays. E.C.L. prepared complexes for cryo-EM; T.N.P. prepared grids and performed cryo-EM data collection. T.N.P. and C.A.I. processed the cryo-EM data, and E.C.L. and C.A.I. built the model. E.C.L., T.N.P., T.T.R. and C.A.I. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests. Correspondence and requests for materials should be addressed to T.T.R. (thibaud.renault@cnrs.fr) or C.A.I. (axel.innis@inserm.fr).

DATA AVAILABILITY

Cryo-EM maps of the 70S-MAAAPQKC-TcmX complex and the associated molecular models have been deposited in the Electron Microscopy Data Bank and Protein Data Bank, with the accession codes EMD-14956 and PDB 7ZTA, respectively. Sequencing data for the iTP-seq experiment have been deposited in the National Center for Biotechnology Information Short Read Archive with the accession code PRJNA854319. Source data are provided with this paper.

SUPPLEMENTARY INFORMATION

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, Peptides, and Recombinant Proteins		
Acrylamide/Bis-Acrylamide 37.5:1, 40%	Biosolve	Cat#014223
Ammonium persulfate	Sigma-Aldrich	Cat#A3678-25G
Bromophenol blue	Thermo Fisher Scientific	Cat#1010223280
Boric acid	Sigma-Aldrich	Cat#B7901-500G
Deoxynucleotide (dNTP) Solution Mix 10mM	New England Biolabs	Cat#N0447S
Diethyl Pyrocarbonate for DEPC water	Sigma-Aldrich	Cat#D5758-100ML
Merck™ Ultrapure Water for Molecular Biology	Thermo Fisher Scientific	Cat#15161735
Adenosine 5'-triphosphate disodium salt hydrate	Sigma-Aldrich	Cat#A2383-25G
СТР	Jena Bioscience	Cat#NU-1011-1G
GTP	Jena Bioscience	Cat#NU-1012-1G
UTP	Jena Bioscience	Cat#NU-1013-1G
Guanosine-5'-thiophosphatedisodium salt	Genaxxon Bioscience	Cat#\$5402.0025
SYBR Gold nucleic acid gel stain	Invitrogen	Cat#S11494
Trizma Base	Sigma-Aldrich	Cat#T1503-1KG
Acetic acid	Sigma-Aldrich	Cat#33209-1L
Trisodium citrate dihydrate	Sigma-Aldrich	Cat#S1804-500G
Ethanol	Sigma-Aldrich	Cat#32221-2.5L-M
N,N,N',N'-Tetramethylethylenediamine	Euromedex	Cat#50406-A
Urea	Sigma-Aldrich	Cat#U5378-1KG
Magnesium chloride hexahydrate	Sigma-Aldrich	Cat#M2670-1KG
Magnesium acetate tetrahydrate	Sigma-Aldrich	Cat#M5661-250G
Potassium glutamate	Sigma-Aldrich	Cat#49601-100G
Spermidine	Sigma-Aldrich	Cat#S2626-5G
NH4-acetate	Sigma-Aldrich	Cat#A1542-500G
Biotin (Long Arm) maleimide	Vectorlabs	Cat#SP-1501
Bis-Tris	Sigma-Aldrich	Cat#B7535-500G
N,N-Dimethylformamide	Sigma-Aldrich	Cat#D4551-250ML
Sodium chloride	VWR	Cat#27788.366
Tween 20	Sigma-Aldrich	Cat#P1379-100ML
Dynabeads M-280 Sreptavidin	Thermo Fisher Scientific	Cat#11205D
HEPES	Sigma-Aldrich	Cat#H4034-1KG
Potassium hydroxide	Sigma-Aldrich	Cat#P1767-250G
Hydrochloric acid	Sigma-Aldrich	Cat#30721-2.5L-M
Phusion polymerase	Recombinant	
Phusion HF buffer Pack 5X	New England Biolabs	Cat#B0518S
E. coli Poly(A) Polymerase supplied with 10X	New England	Cat#M0276L
Poly(A) Polymerase buffer	Biolabs	
T7 RNA polymerase (P266L) 20 mg/ml	Recombinant	Homemade
Ribonuclease R	Recombinant	Kind gift from Dr. Arun Malhotra (University of Miami)

Supplementary Table 1 | Reagents, oligonucleotides and software

T4 RNA ligase 2, truncated supplied with 10X T4 RNA ligase reaction buffer and 50 % PEC8000	New England Biolabs	Cat#M0242L
SuperScript TM III Reverse Transcriptise supplied with	Thermo Fisher	Cat#18080044
a vial (1 ml) of 5X first strand buffer and a vial (500	Scientific	
a viai (1 mil) of 5X mist-strand burlet and a viai (500	Scientific	
	Navy England	C-+#D210EC
ECOKV-TIF	New England	Cal#K31955
A silent I lish Consitivity DNIA Kit		C-+#F0(7.4(2)
Aglient High Sensitivity DNA Kit	Agrient	Cat#5067-4626
Powdered milk	Regilait	
Streptavidin-Alkaline Phosphatase	Promega	Cat#V5591
NBT/BCIP detection kit	Promega	Cat#S3//1
Xylene Cyanol	Biosolve	Cat#242223
DNA Ladder: Low Molecular Weight DNA Ladder	New England	Cat#N3233S
	Biolabs	
RNA Ladder: Century-Plus RNA Marker	Thermo Fisher	Cat#AM7145
	Scientific	
MinElute PCR Purification Kit TM	Qiagen	Cat#28006
RNA Clean & Concentrator [™] -5	Zymo research	Cat#R1015
Isopropanol	Sigma-Aldrich	Cat#19516-4L
GlycoBlue coprecipitant	Thermo Fisher	Cat#AM9515
	Scientific	
MiliporeSigma TM Ultrapure water for Molecular	Thermo Fisher	Cat#09-739-006
Biology	Scientific	
Avian Myeloblastosis Virus Reverse Transcriptase	Promega	Cat#M5101
Sodium Hydroxide	Sigma-Aldrich	Cat#221465
OlAguick Nucleotide removal Kit TM	Ojagen	Cat#28304
Dideoxy-nucleotides	GE Healthcare	Cat#27-2045-01
	Life Sciences	
Thermo Sequenase Dye Primer Manual Cycle	Thermo Fisher	Cat#792601KT
Sequencing Kit	Scientific	
ExoSAP-IT reagent	Thermo Fisher	Cat#PN/78200
	Scientific	
OuikChange Lightning Site-Directed Mutagenesis	Agilent	Cat#210518
Kit	Technologies	Cat#210510
Critical Commercial Account	Technologies	
		C + #F(0F077
PURExpress & RF123 & Ribosomes Kit	New England	Cat#E6850ZZ
	Biolabs	
Oligonucleotides	1	
iTP/TP_frag1_T7_RBS_ATG_f	Seip et al., 2018	Eurogentec custom
CGA-TCG-AAT-TCT-AAT-ACG-ACT-CAC-TAT-		synthesis
AGG-GCT-TAA-GTA-TAA-GGA-GGA-AAA-AAT-		
ATG		
Stop_EcoRV_r	Seip et al., 2018	Eurogentec custom
TAT-ATG-GAT-CCT-TTT-TGA-TAT-TGA-TAT-CTC-		synthesis
ATC-ACA-CCG-AGA-TCG		
iTP/TP_frag1_T7_f	Seip et al., 2018	Eurogentec custom
CGA-TCG-AAT-TCT-AAT-ACG-ACT-CAC-TAT-AG		synthesis
iTP_EcoRV_r	Seip et al., 2018	Eurogentec custom
TAT-ATG-GAT-CCT-TTT-TGA-TAT-TGA-TA		synthesis

iTP NNN15 random-library	This work	Eurogentec custom
GGA-GGA-AAA-AAT-ATG-NNN-NNN-NNN-		synthesis
NNN-NNN-NNN-NNN-NNN-NNN-NNN-		- /
NNN-NNN-NNN- GCG-ATC-TCG-GTG-TGA		
iTP 3' linker Apol	Seip et al., 2018	Eurogentec custom
/5rAPP/GGT-ATC-TCG-GTG-TGA-CTG-ACT-GAA-		synthesis
AAT-TTC-TGT-AGG-CAC-CAT-CAA-T/ddC		
iTP Linker r	Sein et al. 2018	IDT custom synthesis
ATT-GAT-GGT-GCC-TAC-AG	5cip et ui., 2010	
iTP cDNA f	Sein et al 2018	Furgentec custom
	501p et ul., 2010	synthesis
iTP Biotin standard	Sein et al 2018	Furgentec custom
$/5Biosg/AAA_AAAA_AAA_AAAAAAAAT_TAA_CTC_CAT_$	501p et ul., 2010	synthesis
CTA_A		synthesis
ITP NCS f	Soin et al 2018	Eurogentec custom
	Selp et al., 2010	synthesis
	Soin at al 2018	Eurogentec custom
	Selp et al., 2010	synthesis
iTR NCS adapter f	Soin at al 2019	Eurogontos custom
	Selp et al., 2010	curthosis
		synthesis
ATC-TOT-ATA-AGG-AGG-AAA-AAA-TAT-G	This	Fine contra constant
	I NIS WORK	Eurogentec custom
		synthesis
ITD NCC adaptor index44	This work	Functional custom
	This work	Eurogeniec custom
		synthesis
ACA-G		F
$IIP_INUS_addapter_Index/+2$	I NIS WORK	Eurogentec custom
		synthesis
IIP_NUS_adapter_index43	This work	Eurogentec custom
		synthesis
		
IIP_NGS_adapter_index6+1	This work	Eurogentec custom
CAA-GCA-GAA-GAC-GGC-ATA-CGA-GAT-ATT-		synthesis
CTT-CCG-ATC-GAT-TGA-TGG-TGC-CTA-CAG		
ITP_NGS_adapter_index22+2	This work	Eurogentec custom
CAA-GCA-GAA-GAC-GGC-ATA-CGA-GAT-CGT-		synthesis
ACG-GTG-ACT-GGA-GTT-CAG-ACG-TGT-GCT-		
CTT-CCG-ATC-GAT-TGA-TGG-TGC-CTA-CAG		
TP_trag2_r	This work	Eurogentec custom
CTT-GCC-TGC-GCA-CGA-AGA-GTA-CGG-ATG-		synthesis
TTG-TTC-AGA-GTC-AGT-TAT-TAT-TCG-CT		

TD frag? $N/1$ r	This work	Eurogoptoc custom
IF_IIdg2_INVI_I	THIS WORK	supplied custom
		synthesis
	TI · I	
IP_short_r	This work	Eurogentec custom
GGT-TAT-AAT-GAA-TTT-TGC-TT		synthesis
TP_RT_yakima-yellow_r	This work	Eurogentec custom
/5YakYel/GGT-TAT-AAT-GAA-TTT-TGC-TTA-TTA-		synthesis
AC		
TP_MAAAPQKCAAA*_f	This work	Eurogentec custom
GTA-TAA-GGA-GGA-AAA-AAT-ATG-GCA-GCA-		synthesis
GCA-CCG-CAG-AAG-TGT- GCA-GCA-GCA-TGA-		7
AGC-GAA-TAA-TAA-CTG-ACT-CTG		
nZa MAAAPOKC fud	This work	Furgentec custom
		synthosis
		synthesis
AGI-ICA-ITA-ICA-ACC-AAA-CAA-AAA-ATA-AG		
pZα_MAAAPQKC _rvs	This work	Eurogentec custom
CTT-ATT-TTT-TGT-TTG-GTT-GAT-AAT-GAA-CTG-		synthesis
TAC-ACT-TCT-GCG-GTG-CTG-CTG-CCA-TAT-		
GTT-TTC-CTC-CTT-ATA-AGA-CTT-AAT-TA		
Software and Algorithms		
Image Lab	Bio-RAD	https://www.bio-
		rad com/webroot/web/pdf
		/lsr/literature/100007695
		3 pdf
puthon 2.9		bttps://www.pythop.org/
		https://www.python.org/
numpy 1.19.2		https://numpy.org/
pandas 1.1		https://pandas.pydata.org/
matplotlib 3.3.2		https://matplotlib.org/
seaborn 0.11		https://seaborn.pydata.org/
scipy 1.5		https://www.scipy.org/
PEAR v0.9.10		https://cme.h-
		its.org/exelixis/web/softwa
		re/pear/doc.html
DESeq2		https://bioconductor.org/p
		ackages/release/bioc/html/
		DESeq2 html
SerialEM		https://bio3d.colorado.edu
Schulem		/SerialEM/
Polion v4 0		https://rolion.roadthodocs
Kenon v4.0		is (as /valesses 4.0/
		10/en/release-4.0/
CIFFIND v4.1		https://grigoriefflab.umass
		med.edu/cttfind4
Chimera v1.16		https://www.cgl.ucsf.edu/
		chimera/
Phenix v1.20.1		http://www.phenix-
		online.org/
Coot v0.9.7		https://www2.mrc-
		Imb.cam.ac.uk/personal/p
		emsley/coot/
		51115107/0000

Isolde v1.3		https://isolde.cimr.cam.ac. uk/
ChimeraX v.1.3		https://www.rbvi.ucsf.edu/ chimerax/
PyMOL v.2.5.0	Schrödinger	https://pymol.org/2/

Supplementary Table 2 | Cryo-EM statistics

70S-MAAAPQKC-TcmX Data collection and processing Magnification 45,000 Voltage (kV) 200 Electron exposure (e^{-7} Å²) 35.7 Defocus range (µm) -0.4 to -2.5 Pixel size (Å) 0.93 Symmetry imposed C1 Initial particle images (no.) 838,561 Final particle images (no.) 91,383 Map resolution (Å) 2.7 FSC threshold 0.143 Map resolution range (Å) 2.67-8.30 Refinement Initial model used (PDB code) 6TBV Model resolution (Å) 2.7 FSC threshold 0.143 Model resolution range (Å) 2.5-8.7 Map sharpening B factor ($Å^2$) -10 **Model composition** Non-hydrogen atoms 148,592 Protein residues 5,702 **RNA** bases 4,786 Ligands 410 **B-factor** (A²) RNA 101.3 Protein 101.8 Ligand 85.8 **R.m.s.** deviations Bond lengths (Å) 0.0299 Bond angles (°) 4.62 Validation Model-to-map CC 0.83 MolProbity score 1.09 Clashscore 2.39 Poor rotamers (%) 0.67% **Ramachandran plot** Favored (%) 97.66% Allowed (%) 2.32% Disallowed (%) 0.02%



Extended Data Fig. 1 | **Size distribution of inverse toeprints and diversity of the (NNN)**₁₅ **library. a**, Number of deep sequencing reads obtained for inverse toeprints of different lengths, plotted as a function of the distance between the start codon and the RNase R cleavage site. Average read numbers for three independent replicates are shown for TcmX-treated (orange) and untreated (blue) samples. Inverse toeprints within a size range featuring a well-defined three-nucleotide periodicity were retained for further analysis (gray area). The iTP-seq experiment consisted of 3 independent replicates for the TcmX-treated condition (2.8×10^6 , 2.5×10^6 and 2.3×10^6 reads), and 3 replicates for the untreated condition (2.7×10^6 , 2.3×10^6 and 2.6×10^6 reads). On average ~2.5 \times 10^6 reads were obtained per replicate. The error bands show the 95% confidence interval of the mean of the three replicates. **b.** Nucleotide frequency of the raw reads for the (NNN)₁₅ library, showing a relatively equal proportion of each dNTP for codons 2-14.



Extended Data Fig. 2 | **Enrichment of two-amino acid motifs in samples treated with TcmX.** Heatmaps showing the enrichment of individual residues at positions -2 to +1, and all possible combinations of motifs consisting of two adjacent amino acids, following treatment with TcmX. Enrichment is defined as $log_2(F_{TcmX}/F_{untreated})$, where F_{TcmX} is the frequency of occurrence of a given motif in the sample treated with TcmX and $F_{untreated}$ is its frequency in the untreated sample.



Extended Data Fig. 3 | Four-amino acid motifs at position [-2,+1] ending with LK are not significantly enriched following TcmX treatment. Volcano plot of statistical significance ($\log_{10}(p-value)$) against enrichment ($\log_2(fold change)$) for four-amino acid motifs at position [-2,+1]. Motifs ending in LK are shown in yellow and the two LK-containing motifs described in Ref. 6 (FVLK/TILK) are labeled in bold.



Extended Data Fig. 4 | **Variants of the PQKC motif that induce TcmX-dependent stalling.** Volcano plot of statistical significance $(-\log_{10}(p-value))$ against enrichment $(\log_2(fold change))$ for four-amino acid motifs at position [-2,+1]. Single amino acid variants of the PQKC motif are shown in red.



Extended Data Fig. 5 | **Codon composition does not have a major influence on TcmX-dependent ribosome stalling at QK motifs.** Violin plot showing the enrichment of various three-amino acid motifs at positions [-2,0] following treatment with TcmX. Each bar represents a motif variant with a distinct codon composition.



Extended Data Fig. 6 | **Overview of cryo-EM data processing.** Flowchart showing the workflow used to process cryo-EM data for the 70S-MAAAPQKC-TcmX complex in *Relion v4.0 (Ref. 42)*. The final reconstruction could be refined to an overall resolution of 2.7 Å, assessed using a Fourier shell correlation (FSC) cutoff of 0.143.



Extended Data Fig. 7 | **Quality of the 70S-MAAAPQKC-TcmX reconstruction. a,** Cryo-EM density map obtained for a 3D reconstruction of the 70S-MAAAPQKC-TcmX complex in *Relion v4.0 (Ref. 42) and auto-sharpened with Phenix v1.20.1 (Ref. 46),* filtered and colored by local resolution estimation values in *Chimera X v.1.3 (Ref. 50).* **b,** Cross-section of the same map showing the E, P and A sites of the ribosome. **c,** Isolated ligand densities in the E, P and A sites, filtered and colored by local resolution. **d,** The same isolated densities shown in c, displayed as a transparent surface and fitted with molecular models of the E-, P- and A-site tRNAs, mRNA, TcmX and the nascent peptide. **e,** Model to map correlation curve calculated for the 70S-MAAAPQKC-TcmX structure. **f,** Representative cryo-EM densities of three different tRNA modifications within the A- and P-site tRNAs.



Extended Data Fig. 8 | Views of the peptidyl transferase center in the uninduced, induced and TcmX-inhibited ribosome. Structural comparison of 23S rRNA nucleotides and tRNAs within the A site (a,b) and P site (c,d) regions of the peptidyl transferase center in the uninduced²³ and induced (pre-attack)²⁴ states (PDB 1VQ6 and 1VY4, respectively) (a,c), and the uninduced and TcmX-inhibited states (b,d). In the 70S-MAAAPQKC-TcmX structure, 23S rRNA residues U2506, G2583 and U2584 are in their induced conformation, but 23S rRNA nucleotide U2585 and nucleotide A76 of the P-site tRNA remain in the uninduced conformation, indicating partial aminoacyl-tRNA accommodation into the peptidyl transferase center.