Unprecedented tunability of riboswitch structure and regulatory function by sub-millimolar variations in physiological Mg²⁺

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ABSTRACT

Riboswitches are cis-acting regulatory RNA biosensors that rival the efficiency of those found in proteins. At the heart of their regulatory function is the formation of a highly specific aptamer-ligand complex. Understanding how these RNAs recognize the ligand to regulate gene expression at physiological concentrations of Mg²⁺ ions and ligand is critical given their broad impact on bacterial gene expression and their potential as antibiotic targets. In this work, we used single-molecule FRET and biochemical techniques to demonstrate that Mg2+ ions act as fine-tuning elements of the amino acid-sensing lysC aptamer's ligand-free structure in the mesophile Bacillus subtilis. Mg²⁺ interactions with the aptamer produce encounter complexes with strikingly different sensitivities to the ligand in different, yet equally accessible, physiological ionic conditions. Our results demonstrate that the aptamer adapts its structure and folding landscape on a Mg²⁺-tunable scale to efficiently respond to changes in intracellular lysine of more than two orders of magnitude. The remarkable tunability of the IvsC aptamer by submillimolar variations in the physiological concentration of Mg²⁺ ions suggests that some single-aptamer riboswitches have exploited the coupling of cellular levels of ligand and divalent metal ions to tightly control gene expression.

INTRODUCTION

Riboswitches are noncoding mRNA sequences usually found in the 5' untranslated regions of many genes involved in metabolite biosynthesis or transport, which they regulate by binding specific, related metabolites (1,2). Riboswitch architecture includes an aptamer domain that acts as the metabolite-sensing element and a downstream expression platform that interacts with the transcription or translation machinery (3–5). A cascade of local and long-range conformational changes is initiated by ligand binding to the aptamer and transmitted to the expression platform (6), biasing its structure towards one of two competing conformers that control the activation or repression of the downstream gene (Figure 1A). Riboswitches have been shown to regulate gene expression through different mechanisms including transcription termination, translation inhibition and, in some eukaryotes, by exposing an alternative mRNA splicing site (1,5,7).

To date, nearly 40 classes of riboswitches have been discovered (8), among them, three amino acid-sensing riboswitch classes that recognize lysine (8,9), glycine (10) or L-glutamine (11). Lysine riboswitches typically regulate the expression of lysine-specific enzymes in the diaminopimelate (DAP) pathway for lysine, methionine and threonine biosynthesis (12). However, despite the lysine riboswitch's

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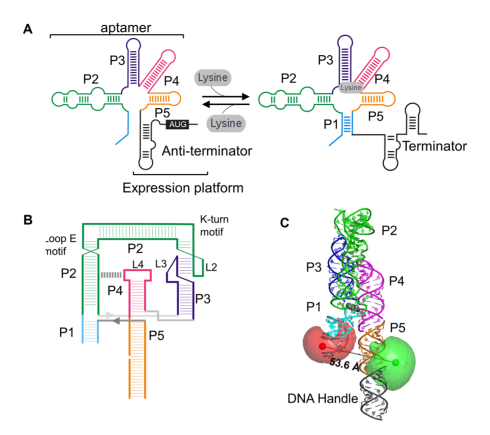


Figure 1. Structural components and regulatory mechanism of the lysine riboswitch aptamer. (**A**) Secondary structure of the *lysC* riboswitch showing the aptamer domain, comprising stems P1 to P5, and the expression platform. Predicted changes in the expression platform induced by lysine binding are also shown. In the absence of ligand an anti-terminator stem forms, allowing transcription readthrough. Upon ligand binding, the expression platform forms a terminator stem that stops transcription. (**B**) Tertiary contacts in the lysine-bound aptamer as observed in the crystal structure. In addition to the P1–P2 and P4–P5 helical stacks, the L2–L3 and P2–L4 interactions assisted by k-turn and loop E motifs, respectively, are also shown. The P5 stem of the experimental construct, shown here, has been extended to include a biotinylated DNA portion for surface immobilization. (**C**) X-ray crystal structure of the ligand-bound lysine aptamer (PDB accession code 3DIL) (20). The positions of the FRET donor Cy3 (green) and the FRET acceptor Cy5 (red) used in this study are shown. Mean dye positions were modelled using the accessible volume (AV) approach (Supplementary Methods). The ligand is shown in grey, and the line between the donor and acceptor spheres represents the predicted mean distance.

abundance (13), biological relevance (14), and proven potential as antibiotic target (15) and in bioengineering (16), its regulatory mechanism remains poorly understood.

The lysC aptamer from Bacillus subtilis comprises five helical stem-loops, labelled P1-P5 in Figure 1A, organised around a core domain (9,17–20) (Figure 1B and c). In the presence of lysine, P1 is stabilized by close association with P5, precluding the formation of an anti-terminator stem and allowing a terminator to form and eject the transcription complex (19,20). Recent studies have revealed the structural motifs and residues responsible for its ligand specificity (21,22). However, many questions about its structural dynamics remain unexplored, including how its folding landscape changes as a function of ionic environment and the ligand; and whether the ligand itself participates in folding. Small-angle X-ray scattering (SAXS) (23) and crystallographic data suggest very little difference between ligand-bound and ligand-free structures (9,20). The crystallized aptamers were from the thermophile Thermotoga maritima, which is extremely stable and has a 30-fold greater affinity for lysine than the aptamer from mesophilic B. subtilis (20). Biochemical data on the B. subtilis aptamer, on the other hand, provide evidence for a global rearrangement of the aptamer upon ligand binding (9,17).

In this work, we have quantified the *in vivo* concentration of Mg²⁺ ions in B. subtilis and used single-molecule FRET (24–26) to explore the conformational dynamics and ligandbinding response of the B. subtilis lysC aptamer within the in vivo range of Mg²⁺ concentrations. We reveal a remarkable tunability of the lysC folding pathway in response to subtle changes in the physiological concentration of Mg²⁺ ions. We demonstrate that sub-millimolar variations in Mg²⁺ impact aptamer function, alter the structure of the aptamer ligand encounter complex, and change the ligand-binding affinity by two orders of magnitude. The extreme sensitivity of the lysC aptamer suggests it acts as a dual-input sensor that relies on Mg²⁺ and lysine levels to ensure tight control over gene expression. Our results imply that cellular levels of Mg²⁺ ions not only participate in RNA compaction and stabilization but could also act as modulators of gene expression. Importantly, the lysC aptamer does not fold into its global holo state in the absence of lysine, regardless of the Mg²⁺ concentration. This is unusual for the riboswitch aptamers that have been studied with single-molecule techniques to date, which often adopt a ligand-free folded conformation that may be only kinetically distinguishable from the ligand-bound state.

MATERIALS AND METHODS

Labelling and purification of RNA oligonucleotides

The experimental constructs for the wild type and variant lysine aptamers incorporate two RNA strands and a biotincarrying DNA tether (see Supplementary Methods for details). The longest strand was generated by in vitro transcription, whilst the other two were synthetic (IDT Inc., USA). The sequences and a schematic of the construct are shown in Supplementary Table S1 and Supplementary Figure S3. Fluorophores were incorporated at the appropriate location either during solid-phase chemistry or post-synthesis using succinimide ester derivatives of Cy3 and Cy5 and following the protocol provided by the dye manufacturer (GE Lifesciences, USA) as described in the Supplementary Methods. Purification of single-strand RNA sequences and hybridized constructs was carried out using polyacrylamide gel electrophoresis, also as described in the Supplementary Methods.

Single-round in vitro transcription

DNA templates were prepared by PCR using the B. subtilis gly Os promoter followed by the B. subtilis lys C riboswitch aptamer domain fused to a 95 nt sequence downstream of the expression platform. To allow transcription to be initiated by the ApC dinucleotide, the promoter region ends with an adenine and was fused to the C17 position of the riboswitch sequence. The truncated product terminates at position 269, whereas for the full-length product the RNAP continues 95 nt further. Reactions were analysed by gel electrophoresis as previously described (17) and performed at least three times.

Intracellular Mg²⁺ measurement

Bacillus subtilis strain 168 was grown in a Spizizen medium at 37°C (150 rpm) for 4 h (27). Every hour, an aliquot of the culture was collected, and cells were pelleted by centrifugation (Beckman Coulter_{TM} J-25I with JLA 16,250 rotor; 6500g, 20°C, 7 min). The cells were then suspended in 4% paraformaldehyde in PBS and incubated 7 min at room temperature for fixation. Cells were pelleted by centrifugation and washed with an oxalate/EDTA solution (0.1 M/0.05 M) at room temperature for 7 min to remove external metals. The pellet was then suspended in 10 ml sterile 0.5 M NaCl. From this 10 ml, 2 ml were coloured with SYTO9 and used for cell volume calculation. Cellular volume was obtained using fluorescent microscopy pictures of B. subtilis taken with the fluorescent dye and it was calculated as a cylinder and two half-spheres using maximum Feret diameter (length) and minimal Feret diameter (width) evaluated by image analysis (CellProfiler 3). Images were taken using a Zeiss Axio Observer Z1 microscope. The other 8 ml were washed again with oxalate/EDTA and pelleted by centrifugation (6500g, 20°C, 7 min). These cells were digested on an SCP science Digiprep Jr with 600 µl of nitric acid (trace metal grade, Fisher Chemical) at 65°C for 45 min. The SCP

science Digiprep Jr is a 24-position Teflon-coated block digestion system that can reach 180°C with 1.0°C uniformity across the surface. After digestion, each tube was filled with 10 ml Milli-O water. Samples were analyzed for phosphorus and metal content on an inductively-coupled-plasma mass spectrometer (ICP-MS; Thermo Scientific, XSeries2) as previously described (28). ICP-MS allows detection and quantification of metals at low concentration (ppt). Analvsis is achieved by ionizing an acid-digested sample with a plasma torch of argon. Ions are then extracted through a series of cones into a quadrupole and separated by their mass/charge ratio before reaching the detector. Phosphorus content was used as a proxy to evaluate cell number.

Single-molecule FRET

Single-molecule FRET experiments were carried out using a Total Internal Reflection microscope that has been described elsewhere (29). The experimental buffer contained 50 mM Tris at pH 7.8, 100 mM K⁺ unless otherwise noted, and Mg²⁺ and L-lysine as indicated. Trolox was used as triplet state quencher (30) and PCA-PCD as the oxygen scavenging system (31). The temporal resolution of the single-molecule movies was 50 ms/frame except to capture the slow dynamics in Figure 5B, where 200 ms/frame was used. FRET was calculated from the raw donor and acceptor intensity traces using $E_{\rm app} = I_{\rm A}/(I_{\rm D} + \alpha I_{\rm A})$ where $\alpha = 0.88$ accounts for 12% leakage into the acceptor detection channel (32). When constructing FRET population histograms, the first ten frames of each movie were averaged to produce a 'average' value for that movie. Between 1100 and 1500 representative FRET values, and 35-450 structural state dwell times in Figure 5B, were used to construct each distribution in this work.

RESULTS

Measurement of the in vivo Mg²⁺ concentration in B. subtilis

Previous work on the lysine riboswitch has addressed aptamers from multiple organisms, both thermophilic (20,33) and mesophilic (12,23,34), in a wide variety of physiological and non-physiological cationic environments. Recent studies on the Escherichia coli metalome suggest free concentrations of Mg²⁺ ions between 1.5 and 3 mM, whereas the total Mg²⁺ concentration, mostly interacting with nucleic acids, proteins and other metabolites, can reach values of ~ 50 mM (35). Given the lack of data regarding the intracellular concentration of Mg²⁺ ions in B. subtilis, we determine the Mg²⁺ concentration range in growth-phase B. subtilis, relevant to lysC riboswitch function in vivo, using inductivelycoupled plasma mass spectrometry (ICP-MS).

Bacillus subtilis strain 168 was grown in a Spizizen (27) medium at 37°C and 150 rpm, including 1.6 mM final Mg²⁺ concentration provided as MgSO₄.7H₂O. Maximum and minimum values of intracellular Mg²⁺ were calculated each hour for 4 h and nine biological replicates. The maximum and minimum Mg²⁺ values were calculated using the maximal and minimal cell volumes observed, respectively (Supplementary Figure S1). The intracellular Mg²⁺ concentrations exhibited very little variation during the growth phase. The minimum average values ranged from 0.8 ± 0.2 to 1.1

 \pm 0.2 mM, and the maximum average values from 2.6 \pm 0.6 mM to 3.7 \pm 0.2 mM. These values are similar to those reported for the free Mg²⁺ concentration in *Salmonella enterica* cells either immersed in a Mg²⁺-free medium ([Mg²⁺]_{free} \sim 0.9 mM) or in 10 mM extracellular Mg²⁺ concentration ([Mg²⁺]_{free} \sim 1.6 mM) (36,37).

Transcriptional regulation by *lysC* riboswitches at physiological Mg²⁺ concentrations

Next, we used a single-round in vitro transcription assay (12,17) to characterize riboswitch regulatory function within the determined physiological range of Mg²⁺ ions (Supplementary Figure S2A and B). In the absence of lysine, a similar transcription termination baseline of $\sim 25\%$ was observed in 1, 2 and 10 mM Mg²⁺. However, when a saturating concentration of lysine ligand (5 mM) was added to the transcription buffer, termination efficiency reached a value of 75% in 2 mM Mg²⁺, which decreased to 51% in 1 mM Mg²⁺ (Supplementary Figure S2C). A further increase in the concentration of Mg²⁺ to 10 mM only slightly increased the termination efficiency to a value of 82%. Transcription termination levels of $\sim 30\%$ in the absence of ligand and maximal termination efficiencies of ~80% at saturating ligand concentrations are typical for many riboswitches, including the lysine riboswitch (21).

The lysine concentration required to induce a 50% transcription termination (T_{50}) decreased by half when the Mg²⁺ concentration increased from 1 to 2 mM, from 129 \pm 15 μ M to 61 \pm 7 μ M. However, T_{50} only decreased by an additional 15%, to 52 \pm 7 μ M, in 10 mM Mg²⁺ (Supplementary Figure S2c). A 50% variation in T_{50} in the 1–2 mM range suggests that *lysC* regulatory function can be strongly modulated by sub-millimolar variations in the physiological concentration of Mg²⁺ ions.

Influence of Mg²⁺ ions on *lysC* conformation

We wanted to determine the conformational response of the lysC aptamer to small changes in Mg²⁺ levels. Therefore, we began by characterizing the structure and dynamics of the ligand-free state as a function of the Mg²⁺ concentration using smFRET in a total-internal reflection (TIR) microscope (38–40). We assembled our experimental lysC aptamer from three oligonucleotide strands (Supplementary Figure S3 and Supplementary Table S1). The Cy5 acceptor was incorporated internally in the P1 stem, and the Cy3 donor on a P5 stem that was extended by 15 bp to hybridize a biotinylated DNA strand for surface attachment (Figure 1B and C and Supplementary Figure S3). We have chosen to monitor the P1-P5 distance based on chemical probing and previous FRET data that suggested a significant structural re-organization of these stems on ligand binding (34). We obtained a molecular model of the doubly labelled aptamer using the X-ray structure of the ligand-bound state (PDB: 3DIL) (20) and the mean dye positions calculated using the accessible volume (AV) approach (41) (see Supplementary Methods) (Figure 1C). The modelled Cy3–Cy5 distance was \sim 53.6 Å. Assuming a Förster distance of 60 Å we expected an apparent FRET efficiency (E_{app}) of ~ 0.66 for the ligand-bound state.

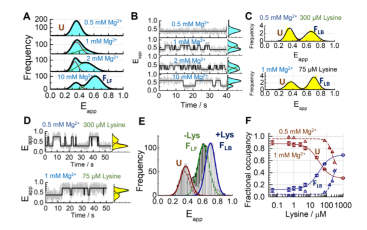


Figure 2. Ligand-binding to unstructured aptamers promotes direct folding into the native state. Single-molecule FRET histograms (A) and corresponding FRET trajectories (B) obtained at the indicated concentrations of Mg²⁺ ions. (C) Comparison of single-molecule FRET histograms obtained in a background of 100 mM K⁺, 0.5 mM Mg²⁺ and 300 μ M lysine ligand (top), and in a background of 100 mM K⁺, 1 mM Mg²⁺ and 75 μM lysine (bottom). (D) Representative single-molecule FRET trajectories obtained in the same conditions as (C). The solid line represents the idealized FRET trajectory obtained using Hidden Markov modelling. (E) Unfolded (U, dark red), ligand-free folded (F_{LF}, green) and ligand-bound folded (FLB, blue) states have distinct FRET efficiencies. Comparison of single-molecule FRET histograms obtained in the absence (dark grey) and presence of lysine (light grey) are shown. The single-molecule histogram in the absence of lysine with a predominant F_{LF} state corresponds to that shown at 10 mM Mg²⁺ in panel a. The single-molecule histogram with a predominant F_{LB} state corresponds to that shown in panel C. (F) Relative contributions of the U (dark red) and F_{LB} (dark blue) states as a function of lysine concentration obtained in a background of 100 mM K⁺ and 0.5 mM (Δ) or 1 mM (o) Mg²⁺ ions. The solid lines represent the result of a global fit of the contributions of the U and F_{LB} states to a Hill model as a function of lysine concentration (Supplementary Methods).

At low concentrations of monovalent ions (Supplementary Figure S4A) or in the presence of EDTA (Supplementary Figure S5A), the single-molecule FRET histograms showed a single Gaussian peak centered at $E_{\rm app} \sim 0.32 \pm$ 0.01(mean \pm s.d.), and the smFRET trajectories of individual aptamers remained constant until photobleaching occurred (Supplementary Figures S4B and S5B). We designated this state the unfolded conformation, U. As we increased the concentration of Mg²⁺ in a background of 100 mM K⁺, a decrease in the relative contribution of the low-FRET peak and a concomitant increase in the contribution of a higher-FRET peak were observed (Figure 2A). In 10 mM Mg²⁺ (Figure 2A, bottom panel), 70% of the population has shifted into a state centered at $E_{app} = 0.6 \pm 0.01$, which we designated the ligand-free folded state, F_{LF}. A representative single-molecule trace for each Mg²⁺ concentration investigated is shown in Figure 2B. As the Mg²⁺ concentration increases, the dwell time in U decreases while the dwell time in the F_{LF} state increases.

Using the average FRET efficiency obtained from the single-molecule histograms, we fitted the Mg^{2+} binding isotherm to a Hill equation (Supplementary Figure S6A). We obtained a dissociation constant of 1.8 ± 0.3 mM and a Hill coefficient of 2.6 ± 0.4 . When these experiments were repeated in the absence of K^+ ions, the smFRET histograms and trajectories followed a similar trend (Supple-

mentary Figure S6B and C). In these conditions we obtained a slightly lower K_D value (1.1 \pm 0.1 mM) and a similar Hill coefficient (2.2 \pm 0.4) (Supplementary Figure S6A). These data confirm that K⁺ only plays a role during the metabolite-recognition phase by mediating lysine-RNA interactions, as observed in the crystal structure of the ligandbound aptamer.

smFRET analysis of two aptamer variants, L2X and L4X, which disrupt the formation of the L2-L3 and P2-L4 interactions (18), respectively (Supplementary Figure S7A and Supplementary Table S1), confirmed that the stabilization of the ligand-free folded state (F_{LF}) requires the formation of both tertiary contacts (Supplementary Figure S7b). Interestingly, an aptamer variant carrying a G39C substitution that was expected to only abolish ligand binding (BPX) (21) also severely compromised the formation of the F_{LF} state (Supplementary Figure S7B), suggesting that the binding pocket becomes at least partially structured in that state.

Metabolite sensing by unstructured lysC aptamers

Next, we investigated whether the ligand could directly interact with aptamers in the U state. In a background of 0.5 mM Mg²⁺, where the aptamer is exclusively in the U state (Figure 2A, upper panel), we observed a progressive increase in the contribution of a new high-FRET state as we increased the ligand concentration. In 300 µM lysine, the relative contribution of this high-FRET state is \sim 42% (Figure 2C, upper panel and Supplementary Figure S8A). When we repeated these experiments in 1 mM Mg²⁺, we observed an increase in ligand binding affinity. We found that the addition of only 75 µM lysine ligand was enough to induce a shift of \sim 65% of the *lysC* aptamer population into the high-FRET state (Figure 2D, bottom panel and Supplementary Figure S9a). This high-FRET population was centred at $E_{\rm app} = 0.66 \pm 0.01$, in good agreement with the modelled distance (Figure 1C) and higher than the FRET efficiency of the F_{LF} state ($E_{app} \sim 0.6$) (Figure 2E), so we designated it the ligand-bound folded state, F_{LB} .

smFRET trajectories obtained in 0.5 mM Mg²⁺ and increasing lysine concentrations exhibited only fluctuations between U and F_{LF} (Figure 2D, upper panel and Supplementary Figure S8B). In 1 mM Mg²⁺, the smFRET traces showed shorter dwell times in U, reflecting the shift in the equilibrium populations towards the F_{LB} state (Figure 2D, bottom panel and Supplementary Figure S9). We concluded that ligand binding to tertiary-unstructured aptamers induces the direct transition to the ligand-bound folded state, F_{LB} , distinguished from F_{LF} by at least a closer juxtaposition of stems P1 and P5. The fractional occupancies of the U and F_{LB} states as a function of the lysine concentration were extracted by calculating the normalized areas of the U and F_{LB} Gaussian populations at each ligand concentration (Figure 2F). Fitting the fractional occupancies to a two-state binding model yielded K_D values of 180 \pm 20 μ M and 33 \pm 3 μ M at 0.5 mM and 1 mM Mg²⁺, respectively.

Ligand-induced dynamics in unstructured lysC aptamers

We quantified the dynamics of $U \leftrightarrow F_{LB}$ transitions by calculating the distribution of dwell times in the U and F_{LB} states

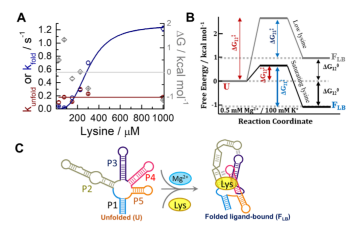


Figure 3. Ligand binding to unstructured aptamers promotes direct folding into the native state. (A) Kinetic rates obtained for the $U \rightarrow F_{LB}$ (dark blue) and $F_{LB} \rightarrow U$ (dark red) transitions as a function of lysine concentration in a background of 100 mM K⁺ and 0.5 mM Mg²⁺. The solid lines represent fits to a straight line for the $F_{LB} \to U$ transition, and to a Hill model for the $U \rightarrow F_{LB}$ transition. Free energy values in kcal/mol obtained from the kinetics rates are also plotted (\Diamond , grey), and $\Delta G^{\circ} = 0$ is indicated by a solid grey line. (B) Free energy landscape of the lysC aptamer in 0.5 mM Mg²⁺ ions at low (\sim 25 μ M, grey line) and saturating lysine concentrations (1 mM, black line). The diagram shows the free energy differences (ΔG°) and barrier heights (ΔG^{\ddagger}) between states U (denoted state 1 in subscripts for simplicity) and F_{LB} (denoted state 2). The absolute barrier heights are representative only but have been scaled such that the magnitudes of the relevant changes $\Delta \Delta G^{\ddagger}$ (Supplementary Table S6) are accurately represented. (C) Single-step mechanism proposed for the interplay between folding and induced-fit ligand binding in the lysC aptamer at sub-saturating concentrations of Mg²⁺ (<1 mM).

as a function of lysine concentration (Figure 3A, Supplementary Figure S10 and Supplementary Table S2). In 0.5 mM Mg²⁺, the rate of the $F_{LB} \rightarrow U$ transition was independent of lysine concentration, and when fitted to a straight line, it yielded a value of $0.18 \pm 0.03 \text{ s}^{-1}$ (Supplementary Table S3). In contrast, the $U \rightarrow F_{LB}$ docking rate increased by \sim 60-fold, from $0.019 \pm 0.003 \text{ s}^{-1}$ in 75 μM lysine to 1.22 $\pm 0.04 \text{ s}^{-1}$ in 1 mM lysine (Supplementary Table S2). The association rates were fitted to a two-state model (Supplementary Equation 2) and yielded a dissociation constant of $310 \pm 50 \,\mu\text{M}$ and a Hill coefficient of 3.1 ± 0.7 , implying a strong coupling between ligand binding and aptamer folding at sub-saturating Mg²⁺ concentrations.

To investigate this mechanism in more detail, we calculated the free energy differences between structural states (ΔG°) and the changes in the barrier height $(\Delta \Delta G^{\ddagger})$ from the single-molecule kinetic data (Figure 3B). In 0.5 µM lysine, U is more energetically favourable than F_{LB} (ΔG = 0.89 ± 0.02 kcal/mol) (Supplementary Table S3). This scenario reverses at high ligand concentrations. For instance, in 150 μ M lysine, F_{LB} is 1.1 \pm 0.1 kcal/mol more favourable than U, and the height of the energy barrier for docking and folding has decreased by \sim 1.9 \pm 0.1 kcal/mol, while the dissociation energy barrier has changed only by -0.34 ± 0.08 kcal/mol. A ligand-dependent docking rate coupled with a ligand-independent unfolding rate fulfills the canonical definition of an induced-fit ligand-binding mechanism (Figure 3C), in which lysine binding re-structures the junction, promotes L2-L3 and P2-L4 contacts, and reorients the P1 and

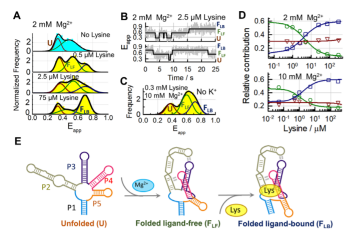


Figure 4. Ligand-induced re-organization of tertiary pre-organized lysine aptamers (A) Single-molecule FRET histograms obtained in the absence of ligand (cyan) and at the indicated concentrations of lysine ligand (yellow) in a background of 100 mM K⁺ and 2 mM Mg²⁺. The result from fitting each histogram to three Gaussians (black line) and the relative contributions of the U (red), F_{LF} (green) and F_{LB} (blue) states are also shown. (B) Representative FRET trajectories obtained in 2 mM Mg²⁺, 100 mM K^{+} and 2.5 μ M lysine showing fluctuations between U, F_{LF} and F_{LB} states within single aptamers. The solid line represents the idealized FRET trajectory obtained using Hidden Markov modelling. (C) Single-molecule FRET histogram obtained in the absence of K⁺ ions at 10 mM Mg²⁺ ions and 300 μ M lysine. (D) Variation in the relative contributions of U (Δ , red), F_{LF} (o, green), and F_{LB} (\square , blue) states as a function of lysine concentration obtained in a background of 100 mM K⁺ at 2 mM (upper panel) and 10 mM (bottom panel) Mg²⁺. The solid lines represent the result of globally fitting the contributions of the F_{LF} and F_{LB} states as a function of lysine ligand concentration to a Hill model (Supplementary Methods). Dissociation constants of $4.5 \pm 1 \,\mu\text{M}$ and $1.3 \pm 0.5 \,\mu\text{M}$ were obtained in 2 mM and 10 mM Mg²⁺, respectively. (E) Proposed two-step folding and ligand-binding mechanism for the lysC aptamer at saturating concentrations of Mg²

P5 stems, thus orchestrating the formation of the global tertiary structure.

Metabolite sensing by partially-folded lysC aptamers

Our analysis of the folding of the lysC aptamer revealed that the F_{LF} state can be efficiently populated at the high end of the physiological Mg²⁺ concentration range (~2 mM). We hypothesised that F_{LF} might function as a high-affinity binding scaffold for lysine in these conditions. To test this hypothesis, we titrated the aptamer with lysine in a background of 2 mM (Figure 4A) or 10 mM (Supplementary Figure S11) Mg²⁺. The smFRET histograms showed that as the lysine concentration increased, a third Gaussian population appeared centred at $E_{\rm app} \sim 0.69 \pm 0.01$, consistent with the $F_{\rm LB}$ state. With a difference in $E_{\rm app}$ of 0.09 \pm 0.01, the F_{LB} and F_{LF} states can be clearly distinguished in singlemolecule traces (Figure 3B). The relative contribution of the **F_{LB}** population in 75 μM lysine and 2 mM Mg²⁺was 59% (Figure 4A). Increasing the Mg²⁺ concentration to 10 mM only increased the fractional occupancy to 64% (Supplementary Figure S11), indicating that the F_{LB} state is already efficiently formed in physiological Mg²⁺ concentrations.

Analysis of the tertiary structure variants, L2X and L4X, and the binding knockout aptamer, BPX, confirmed that none of the aptamer variants can adopt the F_{LB} conforma-

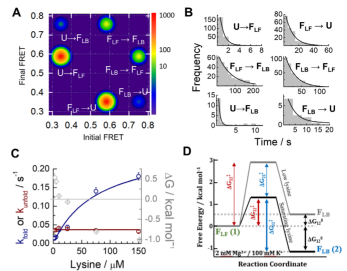


Figure 5. Single-molecule kinetic analysis confirms an induced-fit model for ligand binding at all experimental conditions. (A) Transition density plot (TDP) obtained in 2 mM Mg $^{2+}$, 100 mM K $^+$ and 2.5 μ M lysine, showing the number of times a given transition from $E_{initial}$ to E_{final} was identified. fied in FRET traces showing dynamics. The TDP indicates a prevalence of $U \rightarrow F_{LF}$ and $F_{LF} \rightarrow F_{LB}$ transitions over direct $U \leftrightarrow F_{LB}$ dynamics. (B) Single-molecule dwell-time histograms for each type of FRET transition shown in the TDP. The solid line represents the fit to a mono-exponential decay function. (C) Kinetic rates for the interconversion between the $F_{LF}\,$ and F_{LB} states obtained as a function of lysine concentration. The F_{LB} - F_{LF} rate (dark red) stays constant at 0.037 ± 0.001 s⁻¹, while the F_{LF} - F_{LB} rate (blue) increases with ligand concentration. The free energy difference between F_{LF} and $F_{LB},\,\Delta \emph{G}^{\circ}$ (grey), is also plotted. The F_{LB} state becomes energetically favourable ($\Delta G^{\circ} < 0$) around 10 μ M lysine concentration. (D) Free energy differences ΔG° and barrier heights ΔG^{\ddagger} between F_{LF} (denoted state 1 for clarity) and F_{LB} (state 2). The grey line represents the landscape at 2 mM Mg^{2^+} and 2.5 μM lysine and the black line represents the landscape in 300 µM lysine. The absolute barrier heights are representative only but have been scaled such that the magnitudes of the relevant changes $\Delta \Delta G^{\ddagger}$ (Supplementary Table S6) are accurately represented.

tion, even in high concentrations of Mg^{2^+} or lysine (Supplementary Figure S12). Furthermore, the native aptamer cannot form the F_{LB} state efficiently when K^+ is omitted or substituted with Na^+ (Supplementary Figure S13), even in lysine concentrations as high as 300 μM (Figure 4c). This confirms the specific role of K^+ ions in stabilizing key RNA–ligand interactions.

We plotted the fractional occupancies of each state (U, F_{LF} and F_{LB}) as a function of lysine in both 2 mM and 10 mM Mg²⁺ (Figure 4D). The fractional occupancy of the U state was higher in 2 mM (~35%) than in 10 mM Mg^{2+} (~25%), but it remained nearly constant when increasing the concentration of lysine, while the F_{LF} and F_{LB} states exchanged populations. A global fitting of the binding isotherms for these two states yielded K_D values of 3 \pm 1 μ M in 2 mM and 1.3 \pm 0.5 μ M in 10 mM Mg²⁺, two orders of magnitude lower than in 0.5 mM Mg²⁺ (\sim 310 \pm 50 μ M). Omitting K⁺ ions from the medium increased the K_D to $850 \pm 35 \,\mu\text{M}$ (Supplementary Figure S14), comparable to the value of $660 \pm 50 \,\mu\text{M}$ reported by Batey et al. using a 2-aminopurine fluorescence assay (22). As summarized in Figure 4E, at sufficiently high concentrations of Mg²⁺, the aptamer domain is mostly organized into the F_{LF} state, in

which the L2–L3 and P2–L4 contacts are formed. Lysine binding to F_{LF} 'zips up' the binding pocket by coordinating the two base pairs at the top of the P1 stem and promoting the stacking of P1/P2 and P4/P5 as observed in the crystal structure.

Conformational dynamics of partially-folded *lysC* aptamers

Our analysis of the single-molecule equilibrium populations in 2 mM Mg^{2+} revealed the coexistence of the U, F_{LF} and FLB states in the presence of lysine ligand and physiologically relevant Mg²⁺ concentrations (Figure 4A). To characterize the interconversion kinetics between these states, we examined the dynamics of single aptamers in 2 mM Mg²⁺ ions and 2.5 µM lysine ligand. In these conditions, all three states are populated (Figure 4A), and transitions between them are regularly observed in smFRET trajectories (Figure 4B).

A transition-density plot (TDP) of the raw number of FRET transitions revealed six populations corresponding to pairwise transitions between the U, F_{LF} and F_{LB} states (Figure 5a and Supplementary Table S4. Note the logarithmic intensity scale in Figure 5A). The statistical predominance of the folding route involving all three states $(U \leftrightarrow F_{LF} \leftrightarrow F_{LB})$ over direct $U \leftrightarrow F_{LB}$ transitions can be clearly observed. In the \sim 450 transitions analysed in which U was the initial state, less than 10% of them involved a direct $U \leftrightarrow F_{LB}$ transition (Supplementary Table S4). We extracted the six kinetic rates involved in the $U \leftrightarrow F_{LF} \leftrightarrow F_{LB}$ mechanism in 2.5 µM lysine by fitting the dwell-time histogram of each transition with a mono-exponential decay function (Figure 5b and Supplementary Table S4). The undocking transition $F_{LB} \rightarrow F_{LF}$ was 7-fold slower (0.036 \pm 0.003 s⁻¹) than the rare $F_{LB} \rightarrow U$ transition (0.25 \pm 0.06 s^{-1}). In contrast, direct docking via the $U \rightarrow F_{LB}$ route, although infrequent, was 150-fold faster (2.3 \pm 0.3 s⁻¹) than docking through the $\mathbf{F_{LF}} \rightarrow \mathbf{F_{LB}}$ pathway $(0.014 \pm 0.003 \,\mathrm{s}^{-1})$ (Figure 5B and Supplementary Table S4). This suggests that complexes form faster via the direct two-state pathway $(U \leftrightarrow F_{LB})$, but this route leads to a much less stable F_{LB} state than when formed through the F_{LF} intermediate.

When we extended the kinetic analysis over a lysine concentration range spanning more than two orders of magnitude (Figure 5C and Supplementary Table S5), we found that the unfolding rate was independent of ligand concentration, whereas the $F_{LF} \rightarrow F_{LB}$ docking rate increased by a factor of 10 and became similar to that observed for the $U \rightarrow F_{LB}$ transition (Figure 5B). Fitting this liganddependent rate (Supplementary Eq. 2) yielded a K_D of 80 \pm 30 μ M and an intrinsic k_{fold} for the $\mathbf{F_{LF}} \rightarrow \mathbf{F_{LB}}$ transition of $0.27 \pm 0.08 \text{ s}^{-1}$. Using this k_{fold} and the $\mathbf{F_{LB}} \rightarrow \mathbf{F_{LF}}$ rate obtained previously, we determined the equilibrium constant for the ligand-induced conformational change and obtained a value of 0.14 \pm 0.04. The apparent K_D of the entire transition $(K_{D, app})$ was determined from the product $K_{D, fold}$ • $K_{\rm D, \, bind}$ to be 11 \pm 7 μ M, close to the value of 3 \pm 1 μ M obtained by fitting the fractional occupancies of \mathbf{F}_{LF} and \mathbf{F}_{LB} (Figure 4D, upper panel).

The free energy differences between the F_{LF} and F_{LB} states calculated from the kinetic rates (Figure 5B-D and Supplementary Table S6) suggest that the ligand 'tilts' the energy landscape of the aptamer-ligand encounter complex such that F_{LB} becomes more favourable by -1.84 ± 0.05 kcal/mol. The energy barrier for transitions from F_{LF} to $\mathbf{F_{LB}}$ also decreases by 1.49 \pm 0.09 kcal/mol, whilst the barrier to complex dissociation has increased relative to the low-Mg²⁺ case due to Mg²⁺ stabilization of the complex. However, it remained insensitive to lysine concentration $(\Delta \Delta G^{\ddagger} = +0.05 \pm 0.01 \text{ kcal/mol})$. Thus, although Mg²⁺ ions pre-organize much of the tertiary structure of the aptamer, the metabolite-binding $F_{LF} \rightarrow F_{LB}$ transition retains the kinetic features of an induced-fit mechanism.

DISCUSSION

In this study, we have revealed that the nature of the RNAligand encounter complex, the ligand-binding affinity, the folding pathway and the regulatory function of the lysC riboswitch (Figure 1) are all strongly modulated by small variations within the physiological range ($\sim 0.5-3$ mM) of Mg²⁺ concentrations (Figures 2–4). Mg²⁺-induced compaction of the RNA structure or organization of a native or near-native state are common themes in riboswitch folding (39,42). However, the observed sensitivity of the *lysC* aptamer's structure and ligand-binding affinity to small changes within the physiological Mg²⁺ concentration range is more nuanced than simple Mg²⁺-induced compaction. Our results indicate that Mg²⁺ actively tunes the structure of the ligand-free state to produce encounter complexes, U and \mathbf{F}_{LF} , with strikingly different sensitivities to the ligand (Figures 2F and 4D) in different, yet equally biologically accessible (Supplementary Figure S1), environments (37).

Our single-molecule analysis also demonstrates that the formation of the F_{LF} state and the associated increase in binding affinity to the low micromolar regime are not artefacts of saturating Mg²⁺ concentrations, as concluded in a previous study (34). Rather, the observed two-orderof-magnitude change in binding affinity reflects a Mg²⁺induced shift between these two distinct and biologically relevant binding-competent structures, U and F_{LF}.

Absolute metabolite concentrations have been reported in E. coli using liquid-chromatography-tandem mass spectrometry (43). The intracellular concentration of lysine found in glucose-fed exponentially growing E. coli was 400 μM, which is within the lysine concentration range employed in our studies, and orders of magnitude lower than the concentration at which the amino acid lysine has been recently shown to play a non-specific role in RNA stabilization (35,44). Thus, our single-molecule data collected in the 0.5-2 mM Mg²⁺ range with lysine concentrations up to 1 mM constitute a reasonable model of the extrema of in vivo function in B. subtilis, and the observed tunability of lysC function by Mg²⁺ and lysine ligand represents an intrinsic feature of its regulatory mechanism. On the other hand, the intracellular K⁺ concentration in B. subtilis is in the range 200–400 mM (45,46), and K⁺ ions have been shown to coordinate lysine aptamer folding at concentrations as low as 10 mM (22). Accordingly, in the following discussion, we treat the K⁺ background as effectively saturating, which it would be in any but the most extreme environmental conditions.

The observed kinetics of ligand binding suggest that the *lysC* riboswitch regulates gene expression on a Mg²⁺-

tunable scale between kinetic and thermodynamic control. For instance, in 2 mM Mg²⁺ and 300 µM lysine, its dwell times in the F_{LB} and F_{LF} states are 27 \pm 2 s and 2.8 \pm 0.3 s, respectively (Supplementary Table S6). The time required to transcribe the expression platform without pausing is 1– 3 s at bacterial transcription speeds of 20–60 nt/s (47). An unfolded-state dwell time on the order of the transcription speed is the hallmark of kinetic control, but a folded-state dwell time longer than the transcription time is characteristic of thermodynamic control. If the aptamer does bind the ligand within the transcription window, it is unlikely to unfold while the transcription machinery is still bound. However, as the Mg²⁺ concentration decreases, the unfolding rate increases (Supplementary Table S3), becoming comparable to the transcription time and shifting the gene regulatory dynamics of the aptamer closer to kinetic control. Based on our results, we expect that the riboswitch will be able to bind physiological concentrations of lysine in any environment, but that increasing the Mg²⁺ concentration increases the fidelity of the gene-regulatory signal by accentuating the thermodynamic-like features of the regulatory mechanism.

Ligand-binding mechanisms are broadly classified as following either a 'conformational selection' or an 'induced fit' folding pathway (48,49). These pathways can be defined either structurally, in terms of whether the ligand binds after or before folding of the holo state; or kinetically, in terms of whether ligand binding reduces the unfolding rate of the holo state (selection) or increases the folding rate (induction). Many of the riboswitches studied using single-molecule microscopy to date have been shown to preorganize into a native or native-like state in the presence of Mg^{2+} but the absence of ligand (47,49–58).

This intrinsic Mg²⁺-induced compaction has led to the suggestion that ligand binding to riboswitches in general stabilizes ('selects') the native state from a pre-existing manifold. However, it is logically possible that a riboswitch could exhibit traits of either mechanism, according to either definition, and that these 'mixed' mechanisms could be environmentally dependent. For example, the Pre-O₁ aptamer transitions from an induced fit mechanism to conformational selection as a result of cation-induced folding (56), while the env8 hydroxocobalamin (HyCbl) riboswitch shows binding of the ligand to an unfolded state of the aptamer as expected of induced fit, but also the characteristic slowing of the unfolding rate with increasing HyCbl concentration that indicates conformational selection (59). The manganese-sensing riboswitch yybP-ykoY from B. subtilis has also been shown to shift between induced-fit and conformational selection pathways in response to cellular levels of Mg^{2+} (60).

Growing evidence suggests that 'mixed' ligandrecognition mechanisms with features of both conformational selection and induced fit are common, perhaps even prevalent, in ligand binding to RNA (49). However, our single-molecule population distribution analysis (Figure 3a-b) and kinetic data (Figure 5c-d) support a model where the *lysC* aptamer recognizes the ligand exclusively using an induced-fit mechanism (Figure 6) in both the structural and kinetic senses. To our knowledge, it is the first example of a natural aptamer that does so across the

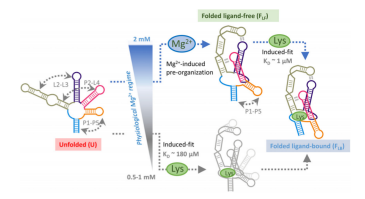


Figure 6. The folding and ligand-binding mechanism of the lysine aptamer is controlled by a narrow change in the physiological concentration of Mg²⁺ ions. Schematic of the interplay between folding and induced-fit ligand binding in the lysC aptamer, emphasising the structural changes taking place within a very narrow Mg^{2+} concentration window and the alternative folding routes predominant at each condition. At the low end of the physiological Mg²⁺ concentration range (<1 mM), the aptamer remains unstructured, and the encounter complex needs to form most tertiary contacts with the assistance of the ligand (lower route). An increase of just 1 mM in the concentration of Mg²⁺ ions is enough to shift the aptamer to a tertiary pre-organised structure (upper route). The encounter complex evolves into the native state by shortening the P1-P5 distance and closing the ligand-binding pocket. The lysine aptamer uses a dual-input signalling model to switch between a single-step, poor-affinity pathway at low concentrations of Mg²⁺ ions to a high-affinity, two-step mechanism at high physiological concentrations.

whole physiological range of Mg²⁺ concentrations. This fact is all the more remarkable given that the Mg²⁺-induced compaction widespread among riboswitches commonly organises the holo state directly, as in the case of Pre-Q₁ (56), the add and pbuE adenine riboswitches (52,61-62), and many others (63,64). In the case of lysC, Mg^{2+} tunes the affinity of the aptamer for lysine but switching still occurs exclusively in the presence of the ligand.

Other examples of environmental dependence exist among riboswitches (51,61–62). For example, the add riboswitch from V. vulnificus uses temperature-dependent switching between two ligand-free secondary structures to modulate its response to adenine inside and outside its host (61,62). The SAM-I riboswitch from T. tengcongensis achieves maximal aptamer preorganisation in the physiological Mg²⁺ concentration range, while higher concentrations favour the organisation of the expression platform (64). The lysC aptamer's dynamic dependence on Mg²⁺ concentration variations within the physiological range can be considered a new type of environmental modulation, in which Mg²⁺ tunes the ligand affinity of the aptamer and the fidelity of its regulatory outcome by varying the structure of the encounter complex without inducing holo state formation (51). In vivo experiments linking the expression of lysC to the variation in the concentrations of \hat{Mg}^{2+} and ligand in both the normal cell cycle and stressed environments is needed to illuminate the biological role of this novel duallayer control mechanism.

In summary, we have demonstrated the first example of an RNA aptamer that has optimized its response to Mg²⁺ ions to act as a fine-tuning element of its tertiary structure. By doing so, it modulates its affinity for its cognate ligand

by two orders of magnitude without the need to alter its nucleotide sequence, a strategy that might also be exploited by other regulatory elements to tightly control gene expression.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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