Local-to-global signal transduction at the core of a Mn$^{2+}$ sensing riboswitch

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The widespread Mn$^{2+}$-sensing $yybP$-$ykoY$ riboswitch controls the expression of bacterial Mn$^{2+}$ homeostasis genes. Here, we first determine the crystal structure of the ligand-bound $yybP$-$ykoY$ riboswitch aptamer from Xanthomonas oryzae at 2.96 Å resolution, revealing two conformations with docked four-way junction (4WJ) and incompletely coordinated metal ions. In $>100$ µs of MD simulations, we observe that loss of divalents from the core triggers local structural perturbations in the adjacent docking interface, laying the foundation for signal transduction to the regulatory switch helix. Using single-molecule FRET, we unveil a previously unobserved extended 4WJ conformation that samples transient docked states in the presence of Mg$^{2+}$. Only upon adding sub-millimolar Mn$^{2+}$, however, can the 4WJ dock stably, a feature lost upon mutation of an adenosine contacting Mn$^{2+}$ in the core. These observations illuminate how subtly differing ligand preferences of competing metal ions become amplified by the coupling of local with global RNA dynamics.
Riboswitches are structured RNA domains commonly found in the 5′-untranslated regions of bacterial mRNAs, where they regulate many essential and virulence genes in response to binding of a specific ligand. Currently, there are over 40 different riboswitch classes known to respond to ligands ranging from metabolites, enzyme cofactors, signaling molecules, tRNAs, and to metal ions. Ligand binding generally stabilizes a conformation of the riboswitch that modulates either a ligand-independent transcriptional termination or translocation initiation through accessibility of the Shine-Dalgarno (SD) sequence. The static ligand-bound structures, and the ligand-recognition modes, of a number of riboswitch aptamers have been determined at atomic resolution. Often, the ligand occupies a linchpin position in the global fold where distal residues of the RNA are brought together; however, the dynamic machinery largely remain enigmatic.

The ybbP-ykoY RNA motif is one of the most widespread riboswitches across bacteria, including many human and plant pathogens. It has evolved to sensitively detect Mn²⁺ metal ions and broadly regulate a variety of genes, particularly those involved in Mn²⁺ homeostasis, at the levels of either transcription or translation. We and others have previously solved crystal structures of the aptamer domain of this riboswitch, henceforth simply referred to as “riboswitch”, revealing that it senses the charge, geometry, and Lewis-acid hardness of Mn²⁺. The global structure showed that formation of the Mn²⁺ binding site requires “locking” of two distal helical legs of a four-way junction to form a paperclip-shaped global architecture, facilitated by an A-minor interaction and a second, nonspecific divalent metal ion binding site. However, the transduction of ligand binding in this metal-sensing core into global structural changes that affect the distal helix P1.1 involved in riboswitching is not understood, rendering it an archetypical representative of our level of understanding of many crystallized riboswitches.

Here, we first solve the structures of two ligand-binding states of the ybbP-ykoY transcriptional riboswitch from the rice pathogen X. oryzae, captured in distinct conformations that offer snapshots of structural changes en route to full ligand binding. We then use these conformers for atomistic molecular dynamics (MD) simulations that reveal how ligand-dependent local structural perturbations in the metal-sensing core are linked to the stability of distal P1.1 helix to affect riboswitching. Finally, using single-molecule FRET (smFRET), we investigate the global structural dynamics of the riboswitch in the presence of varying concentrations of Mg²⁺ and Mn²⁺, as well as other transition metals, revealing a previously unobserved extended conformation of the riboswitch. We show that addition of Mg²⁺ induces two kinetically distinct docked and undocked conformations that are in dynamic equilibrium with one another. In contrast, upon addition of submillimolar Mn²⁺, the riboswitch adopts a stably docked (SD) conformation that becomes abolished upon mutation of the conserved core adenosine. Taken together, our work reveals the ligand-dependent (un)folding pathway of the Mn²⁺ sensing riboswitch as a guide for how subtle binding preferences distinguishing two similar metal ion ligands cascade through the coupling of local with global RNA conformational dynamics into powerful effects on gene regulation.

Results

Mn²⁺. Sensing by stepwise removal of metal ion hydration. Previously, we determined the structures of the Mn²⁺-bound and Mn²⁺-free states of the L. lactis (Llac-Mn) and E. coli (PDB: 4Y1I) and E. coli (PDB: 4Y1M) ybbP-ykoY riboswitches, respectively. The Mn²⁺-sensing core was shown to be first bridged via the binding of a Mg²⁺ ion to a nearby site, then undergoes further conformational changes into the final Mn²⁺-bound state. Importantly, Mn²⁺ sensing involves the complete stripping of the Mn²⁺ ion’s hydration shell, substituted with a set of selective metal–RNA contacts. Since completely dehydrating a metal ion is energetically costly, this process was speculated to take place in stepwise fashion, possibly achieved by a stepwise set of local conformational changes in the metal ion sensing loop. However, no structural intermediates have been captured so far; hence our understanding of the selective metal ion sensing process has remained vague.

In this study, we report the partial and complete Mn²⁺ dehydronation states, captured in the same crystal lattice of the X. oryzae ybbP-ykoY riboswitch and resolved to 2.96 Å (Xory-Mn, Fig. 1a, b and Table 1). This riboswitch controls the yebN Mn²⁺ efflux pump gene in X. oryzae, a phytopathogenic bacterium that causes rice blight. To facilitate crystallization, the WT Xory-Mn sequence was modified distally from the metal ion sensing core not to affect metal ion binding, tightening the 4WJ and stabilizing P1.1 to aid crystallization (see Methods, Fig. 1a). The crystal lattice contains two Xory-Mn riboswitch molecules (Supplementary Figs. 1–4 and 6). While their overall architecture is similar to that of the previously determined L. lactis 4Y1I (Llac-Mn) structure, with an r.m.s.d. of ~2.5 Å, they represent two distinct functional states (Fig. 1c). Conformer 1 is clearly in the canonical Mn²⁺-sensing state despite minor sequence differences in the metal ion binding core, hence providing a comparable reference point for mechanistic interpretation. Its structure superimposes well with the previously determined, canonical, Mn²⁺-bound L. lactis structure (Fig. 1c). Both feature two coaxial stacked “legs” connected at the top by the 4WJ, a superimposable metal ion sensing core, with bound Mn²⁺ and Mg²⁺ ions in superimposable positions (Fig. 1c). As in Llac-Mn, the L1 loop in Xory-Mn adopts a sarcin-ricin-like motif (SRL-like) and extrudes a conserved A10 to form a cross-helix A-minor interaction with helix P3.1 above L3 (Figs. 1c and 2a, b). The L3 loop is extruded from the opposite side of the helical stacks, adopting a highly compressed conformation and stacks below A10 in L1. The close juxtaposition of phosphates in the L1–L3 interface creates a metal ion binding hotspot. In Llac-Mn, a largely inner-sphere coordinated (five phosphoryl oxygens and one water molecule) metal ion is located above the A10 residue (Supplementary Fig. 5). Both Mg²⁺ and Mn²⁺ can access this site; we will henceforth refer to it as Mₐ₆ₛ because the physiological concentration of Mg²⁺ is usually 100-fold higher than Mn²⁺. This metal ion binding site is preserved in Xory-conformer 1, although the inner-sphere contacts to A10 and G8 are either lost or replaced by outer-sphere contacts (Fig. 2a). Of note, although 80 mM SrCl₂ were present in the crystallization buffer, discernible strontium anomalous signals were found in many sites of Xory-conformer 1 but almost none in the Mₐ₆ₛ site (Supplementary Fig. 3), suggesting that, like Llac-Mn, Xory-conformer 1 prefers smaller divalent ions such as Mg²⁺ or Mn²⁺ in Mₐ₆ₛ. Mn²⁺ coordination is specifically found in a nearby site (Mₐ₆₅ₛ), through complete dehydration and a Mn²⁺-specific “soft” contact from the N7 of an invariable adenosine residue (A41 in Llac and A48 in Xory) (Fig. 1d). In Llac-Mn, the four residues following the Mn²⁺-sensing adenosine in the L3 loop form a three-residue stack with U43 mediating a U-turn motif. Due to sequence differences (ACA₄₉U₄₉ACU₅₂ in Xory vs UCA₄₉U₄₉UCU₅₉ in Llac), while the three-residue-stacking theme is preserved, the U-turn is found to be mediated by the first residue (U₄₉) in Xory-conformer 1, rather than the second


residue (U43) in Llac-Mn (Fig. 1d and Supplementary Fig. 5). Despite the idiosyncratic folding in the L3 loop, the inner-sphere contacts to Mn$^{2+}$ are mediated by the same set of phosphates and the invariable adenosine, suggesting that they utilize the same mechanism to sense Mn$^{2+}$ (Fig. 2a).

Whereas Xory-conformer 1 serves as a reference point to reveal common themes in Mn$^{2+}$ sensing among yybP–ykoY riboswitches, Xory-conformer 2 provides a rare, intermediate snapshot of the riboswitch sampling for Mn$^{2+}$ through a partial dehydration process. The Mg$^{2+}$ ion is more loosely coordinated in the MA,Mg site; an inner-sphere coordination from U52 of the L3 loop is lost (Fig. 2b). This is because the L3 loop has yet to adopt its final Mn$^{2+}$-sensing conformation, as the first four residues in L3 have not formed the ordered stacks atop the invariable A48. Importantly, a partially dehydrated metal ion is already present at the MB,Mn site despite lacking half of the contacts seen in Conformer 1, where only the inner-sphere contacts by the phosphates of A45 and A46 are preserved (Fig. 2b). Interestingly, the electron density is sufficiently clear to suggest that the invariable adenosine residue (A48) rotates slightly and orients its N1 group closer than N7 towards MB,Mn (a 2.6 Å outer-sphere contact). Also of note, the temperature B-factors for L3 are higher in Conformer 2 than Conformer 1 (Fig. 2c, d), indicating elevated conformational flexibility. Importantly, this includes larger flexibility around P1.1, suggesting a possible link between the flexibilities of these two key regions. We interpret this conformation as representing the functional state of the yybP–ykoY riboswitch sampling metal ions.
in solution. The partial dehydration process selects a metal ion with an ionic radius similar to Mn\(^{2+}\). Mg\(^{2+}\) could in theory access the site, however, the N1 and N7 functional groups in the invariable adenosine residue would favor a softer metal ion such as Mn\(^{2+}\) (Fig. 2b). We envision that a further conformational transition to that in Conformer 1 would complete the metal ion selection process, as this process would exert additional stringency by not only selecting ionic radius, but also specifying an octahedral coordination and a clear preference for a "soft" transition metal ion (Fig. 2a, b). Taken together, our crystal structures provide important insights into the stepwise metal-sensing process and form the basis for the following MD simulations.

### Effect of M\(_{A,Mg}\) and M\(_{B,Mn}\) inner-sphere contacts on L3 stacking

To provide deeper insights into the RNA dynamics associated with metal ion identity, we performed 43 atomistic MD simulations, equivalent to a total of 113 \(\mu\)s of real-time (see Supplementary Table 1, Methods), with different metal ions in the M\(_{A,Mg}\) and M\(_{B,Mn}\) sites and starting either from Xory-Mn Conformers 1 or 2 (as reported here) or from L. lactis structures (PDB IDs 4Y1I and 6CB3)\(^{12,19}\). We found that the arrangement of the ion binding sites in chain A of the 6CB3 L. lactis structure involved three Cd\(^{2+}\) ions and required the deprotonated form of uracil U44\(^{\text{\textdagger}}\) to prevent loss of its first-shell coordination in the simulations. In addition, the replacement of the Cd\(^{2+}\) ions by one Mn\(^{2+}\) and two Mg\(^{2+}\) ions in the M\(_{B,Mn}\) M\(_{A,Mg}\) and M\(_{C}\) ion binding sites resulted in modest reorganization of their first-shell coordination spheres even in the presence of a deprotonated U44\(^{\text{\textdagger}}\). Therefore, this particular arrangement of ion binding sites appears to be specific for large Cd\(^{2+}\) ions, likely due to their enhanced affinity to nitrogen ligands and preference for hepta-coordination over the native Mn\(^{2+}\) ion (Supplementary Discussion, Supplementary Note 3 and Supplementary Figs. 11–14).

Therefore, we view this specific arrangement around the toxic and xenobiotic Cd\(^{2+}\) ions as not fully relevant for physiological Mn\(^{2+}\)/Mg\(^{2+}\) inner-sphere conditions.

As expected, given the limited timescales compared to experiments\(^{22,23}\), we found that our simulations generally maintained quite stable inner-sphere contacts for divalent metal ions, including their coordination with water molecules (Fig. 3a). To accelerate these dynamics, we performed additional MD simulations that replaced one or both divalent ions at the M\(_{A,Mg}\) and M\(_{B,Mn}\) ion binding sites with monovalent K\(^{+}\) ions (Supplementary Table 1). Replacement of divalent ions by monovalents increases flexibility of the starting ion-binding arrangement, which allows us to observe more local dynamics on the simulation timescale while retaining most relevant aspects of ion binding, since monovalent and divalent ions do compete for the same binding sites.\(^{22,23}\) Indeed, the global architecture of the riboswitch was maintained except for local perturbations of L3 and a variable P2–P4 interhelical angle (Supplementary Figs. 7, 8, and Supplementary Note 1).

The simulations revealed that the inner coordination spheres of monovalent ions in both the M\(_{A,Mg}\) and M\(_{B,Mn}\) sites were significantly more stable when the second binding site was occupied by a divalent rather than monovalent ion (Fig. 3a). This observation suggests that binding of the divalent ion into one binding site helps pre-organize and stabilize the second binding site, consistent with the notion of cooperativity between M\(_{A,Mg}\) and M\(_{B,Mn}\). Furthermore, while some experimentally observed inner-sphere contacts remained very stable upon replacement with a monovalent, other contacts were lost (Fig. 3a–c). Notably, the bridging inner-sphere contacts of the US2 phosphate with the M\(_{A,Mg}\) ion and those of the C51 and G9 phosphates with the M\(_{B,Mn}\) metal ion were found to be most labile across all simulations (Fig. 3a, Supplementary Table 2, and Supplementary Figs. 15–19). These are the same inner-sphere contacts found to be lost in the crystallized Conformers 1 and 2 compared to the 4Y1I and 6CB3 (chain B) L. lactis structures, supporting the notion that the crystallized X. oryzae Conformers are relevant as en route to (un) folding of the metal ion binding sites. In contrast, the inner-shell contacts of the A46 phosphate with the ions in both M\(_{A,Mg}\) and M\(_{B,Mn}\) as well as that of the G9 phosphate with the ion in M\(_{A,Mg}\) were stable across all simulations (Fig. 3a, Supplementary Table 2, and Supplementary Figs. 15–19). These interactions are likely stabilized by coupling of the A-minor interaction of A10 to the G66–C44 base pair and associated stacking of A10 on the neighboring A46. All these interactions are parts of the L1–L3 tertiary docking contact and remained stable across all simulations (Fig. 3d).

We also found the dynamics of the entire L3 loop to be sensitive to the type of ion in the M\(_{B,Mn}\) site. When Mn\(^{2+}\) in the M\(_{B,Mn}\) site was replaced with K\(^{+}\), the flexibility of the L3 loop was increased so that the loop populated various conformations with broken stacking patterns (Fig. 3e and Supplementary Note 4). These perturbations originated from the weakened inner-sphere contact of the A48(N7) nitrogen to the M\(_{B,Mn}\) metal, suggesting that the L3 loop is involved in direct sensing of the Mn\(^{2+}\) ion by forming a linchpin to support a tight stacking patterns only in the presence of the native Mn\(^{2+}\) (Fig. 3e).

### Stability of the L1 loop requires an A-minor interaction

To complement our probing of structural dynamics in the docked state with those in the undocked state, we performed additional

| Table 1 X-ray crystallographic data collection and refinement statistics for the X. oryzae yybP-ykoY Mn\(^{2+}\) riboswitch |
|-----------------------------------------------|
| **Xory yybP-ykoY riboswitch** | **Crystal 1** |
| Data collection | |
| Space group | P 21 21 21 |
| Cell dimensions | a, b, c (Å) | 81.32, 85.19, 92.49 |
| α, β, γ (°) | 90, 90, 90 |
| Resolution (Å) | 62.66–2.96 (3.07–2.96)* |
| R\(_{\text{sym}}\) or R\(_{\text{merge}}\) | 0.110 (1.51) |
| Completeness (%) | 99.83 (99.63) |
| Redundancy | 81.90 |
| CC1/2 | 1 (0.599) |
| CC* | 1 (0.866) |
| Refinement | |
| Resolution (Å) | 62.66–2.96 (3.07–2.96) |
| No. of reflections | 13901 (1361) |
| R\(_{\text{work}}\)/R\(_{\text{free}}\) | 0.205 (0.220) |
| No. of atoms | RNA 4214 |
| Ligand/ion | 120 |
| Water | 7 |
| B-factors | |
| RNA | 69.17 |
| Ligand/ion | 93.43 |
| Water | 66.69 |
| R.M.S. deviations | |
| Bond lengths (Å) | 0.005 |
| Bond angles (°) | 0.59 |
| PDB accession code | 6N2V |

Note that the data were mildly anisotropic, with reflections along the reciprocal axis stronger.

Thus, U/σ for these reflections was well over 1.2 (indeed, over 1.5 up to 2.85 Å resolution)

*One crystal was used. Values in parentheses are for highest-resolution shell

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MD simulations of the segment consisting only of P1.1, P1.2, and L1. We started these undocked simulations from either of the two Xory-Mn Conformers and Llac-Mn structure (Supplementary Table 1), with the aim to reveal their structural dynamics in the absence of the L1–L3 docking contact. While the L1 loop in the context of the entire riboswitch with its docked L1–L3 interaction always populated the SRL-like conformation, two of our three undocked simulations lost this motif (Fig. 3f, g, Supplementary Fig. 22, and Supplementary Note 5). In particular, in one of the simulations we observed loss of the G9–G93 trans Watson–Crick/Hoogsteen and G8–A94 trans Sugar-Edge/Hoogsteen base pairing24 as well as the S-turn25 that forms part of the MA,Mg site (Fig. 3f, g). The former two base pairs are coaxially stacked on the P1.1 stem, suggesting that their loss may highlight the beginning of a transduction path by which P1.1 could become destabilized.

Fig. 2 Conformational differences in the metal ion binding sites and B-factors in the structures. Close-up view of the M\textsubscript{A,Mg} (black) and M\textsubscript{B,Mn} (purple) metal ion binding sites of a Conformer 1 and b Conformer 2 showing different contacts with surrounding L1 and L3 residues. Waters are shown as red spheres. The lost metal ion contacts are shown as red dashed lines with their distances. The anomalous difference map at 4\(\sigma\) (magenta mesh) shows that Sr\textsuperscript{2+} is not observed in Conformer 1 and weakly occupies the M\textsubscript{A,Mg} site but not the M\textsubscript{B,Mn} site in Conformer 2. The Llac-Mn structure is shown in Supplemental Fig. 5 in the same orientation for comparison. Overall X. oryzae Conformer 1 (c) and Conformer 2 (d) structures, colored by individual atomic temperature B-factors (range: 38–135 Å\(^2\) for Conformer 1, 44–170 Å\(^2\) for Conformer 2). Notice both are well-structured in L1 and in P3 around the binding site. However, Conformer 1 is much more tightly structured in L3 than is Conformer 2. Also, the metal ions in Conformer 1 are somewhat less variable. In both molecules, there is flexibility around the 4WJ
**Fig. 3** Structural dynamics of inner-shell ligands of the MA,Mg and MB,Mn sites, L3 and L1 loops as revealed by MD simulations. 

- **a** The population of native inner-shell contacts in the MA,Mg and MB,Mn sites in percent as a function of the type of ion in these sites. A representative structure of the MA,Mg and MB,Mn site as typically observed in MD simulations with a K⁺ ion occupying the site. The most labile inner-shell contacts are depicted in cyan.
- **b** Partial unfolding of the stacking pattern in the L3 loop observed upon replacement of Mn²⁺ with K⁺ in the MB,Mn site.
- **c** The loss of native G9–G93 trans Watson-Crick/Hoogsteen and G8–A94 trans Sugar-Edge/Hoogsteen base pairing, and 
- **d** loss of the S-turn backbone conformation involving a phosphate notch forming part of the MA,Mg ion binding site in response to loss of the L1-L3 tertiary interaction with its A10-G66 = C44 A-minor interaction.
smFRET reveals an unlocked state that docks upon Mg$^{2+}$ addition. To probe the global structural dynamics in the presence of Mg$^{2+}$ and Mn$^{2+}$, we used smFRET to monitor fluorophores positioned on the distal legs of the Xory riboswitch (Fig. 4a, Supplementary Fig. 23, and Methods). smFRET traces at 100 mM KCl in the absence of any divalent metal ions showed a stable low-FRET value of ~0.1 without global dynamics (Fig. 4b), with a population FRET histogram displaying a single peak centered on ~0.13 ± 0.10 (mean ± standard deviation) (Fig. 4c). The non-dynamic nature of the traces is also evident as an on-diagonal contour centered at ~0.13 in the transition occupancy density plot (TODP), which represents as a heat map the fraction of single contour centered at ~0.13 in the transition occupancy density plot (TODP), which represents as a heat map the fraction of single molecule traces that exhibit any given speci
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Submillimolar Mn$^{2+}$ uniquely yields a SD riboswitch. We next asked what specific effect Mn$^{2+}$ has on the folding of the riboswitch. In the presence of 1 mM Mg$^{2+}$, addition of 0.1 mM Mn$^{2+}$ results in the appearance of a unique population of SD (~43%) traces residing in the high-FRET state for >30 s (k_{undock} < 0.03 s$^{-1}$) before photobleaching (Fig. 5a). Accordingly, the FRET histogram shows two peaks with mean FRET values of 0.17 ± 0.14 and 0.69 ± 0.12 and an increased ~68% population of the docked conformation (Fig. 5a). The SD population is evident in the TODP as a new on-diagonal contour centered on the ~0.7-FRET value (Fig. 5a). Double-exponential kinetics similar to the 1 mM Mg$^{2+}$ alone condition are observed, with k_{fast, dock} = 5.55 s$^{-1}$, k_{slow, dock} = 0.46 s$^{-1}$, k_{fast, undock} = 3.84 s$^{-1}$ and a ~5-fold slower k_{slow, undock} of 0.23 s$^{-1}$. Notably, in the presence of Mn$^{2+}$, most of the dynamic traces show DD character. These data demonstrate that Mn$^{2+}$ binding stabilizes the docked conformation while uniquely enabling an SD state. FRET histograms further show that out of a variety of divalent metal ions tested, only Cd$^{2+}$ is effective in promoting docked conformations over 1 mM Mg$^{2+}$ alone; Ni$^{2+}$, Co$^{2+}$, Sr$^{2+}$, or Zn$^{2+}$ had little effect (Fig. 5b and Supplementary Fig. 26). Interestingly, in the absence of Mg$^{2+}$, while 0.1 mM Mn$^{2+}$ alone leads to the appearance of DD and SD traces with ~62% docked population (mean FRET 0.67 ± 0.12) (Fig. 5c), 0.1 mM of Ni$^{2+}$, Co$^{2+}$, Sr$^{2+}$, or Zn$^{2+}$ does not affect SU traces and Cd$^{2+}$ has only a small effect in promoting DD traces (Supplementary Note 6 and Supplementary Fig. 26). These results suggest that while the Xory riboswitch has some degree of plasticity in recognizing ligands in a background of Mg$^{2+}$, it preferentially recognizes Mn$^{2+}$ and—to a lesser extent—Cd$^{2+}$.

To probe how the kinetically distinct SU, DU, and DD traces respond to Mn$^{2+}$, we observed the same set of molecules at 1 mM Mg$^{2+}$, before and after the addition of Mn$^{2+}$. We find that all three populations respond to Mn$^{2+}$ and are capable of forming the stably docked (SD) population (Fig. 5d). In particular, the SU traces convert into DU, DD, and SD conformations with similar probabilities, suggesting that they are correctly folded with metal-sensing sites poised to bind ligand. By comparison, a majority of DD traces convert into SD traces, whereas DU traces adopt DD and SD behavior upon Mn$^{2+}$ addition. Only a small fraction of traces show no response at low Mn$^{2+}$ suggesting that they may be misfolded.

Mutation of A48 results in complete loss of SD conformation. The highly conserved discriminator base A48 in L3 is positioned to confer Mn$^{2+}$ specificity via its N7 and also helps maintain L3 in a stacked conformation. We, therefore, tested the effect of a single A48U mutation on folding and Mn$^{2+}$ sensing of the riboswitch. Similar to the WT, at 100 mM KCl without divalents, the mutant riboswitch shows SU population with a mean FRET value of 0.11 ± 0.12 (Fig. 6a). At 1 mM Mg$^{2+}$, we observe dynamic traces with excursions into the docked higher FRET states, and the FRET histogram shows a major (64%) 0.14 ± 0.11 low-FRET peak and a minor (36%) broad 0.52 ± 0.21 mid-FRET peak that corresponds to more extended docked conformations (Fig. 6b). Of note, 100% of the dynamic mutant traces show DU character under this condition (Fig. 6b). As a result, we observe fast single-exponential kinetics with k_{undock} = 6.67 s$^{-1}$, while k_{dock} was double-exponential with a major k_{slow, dock} = 3.70 s$^{-1}$ (95%) and a minor k_{fast, dock} = 0.46 s$^{-1}$ (5%) (Supplementary Fig. 27). The TODPs show SU behavior in the absence of divalents, whereas both SU and DU behaviors are observed at 1 mM Mg$^{2+}$ (Fig. 6a, b).
Next, we asked whether the mutant riboswitch can still respond to Mn$^{2+}$. In the presence of 1 mM Mg$^{2+}$ and 0.1 mM Mn$^{2+}$, most smFRET traces show DD behavior, while only a small fraction remain in the SU state, similar to WT (Fig. 6c). FRET histograms show a similar ~0.14 ± 0.13 (57%) low-FRET undocked state but display a docked state now with a higher mean-FRET value of 0.63 ± 0.15 and a larger ~43% population, compared to the 1 mM Mg$^{2+}$ only condition. The kinetics under these conditions are double-exponential with a $k_{\text{fast dock}}$ = 6.67 s$^{-1}$ (89%), $k_{\text{slow dock}}$ = 0.64 s$^{-1}$ (11%), $k_{\text{fast undock}}$ = 3.10 s$^{-1}$ (94%), $k_{\text{slow undock}}$ =
Fig. 4 smFRET analysis of the WT Xory Mn^{2+} riboswitch. a Schematic of the smFRET experiment using TIRFM indicating the fluorophore labeling positions on the riboswitch. b Representative smFRET traces under different buffer conditions (top–bottom): no divalents (~0.1 mM EDTA), 0.5 mM Mg^{2+} and 1 mM Mg^{2+}, respectively. Green, Cy3; Red, Cy5; Black, FRET. c Population FRET histograms showing the equilibrium distribution of two FRET states under the conditions in panel (b). Gaussian peaks for the low- and high-FRET states are shown in red and green, respectively with the cumulative fit shown in black. Reported are the percentages of FRET states at equilibrium, as well as the number of molecules N analyzed. d TODPs showing the static and dynamic traces as “on-diagonal” and “off-diagonal” heat map contours, respectively. The color code indicates the fraction of each population. e Fraction of the high-FRET state as a function of Mg^{2+} concentration, fit with a standard Hill equation (red). f Kinetics of structural dynamics as a function of Mg^{2+} and Mn^{2+} concentration. The diamond symbols represent rates in 1 mM Mg^{2+} and 0.1 mM Mn^{2+} while the triangle symbols represent rates in 0.1 mM Mn^{2+} alone.

g Exemplary dynamic docked (DD, top) and dynamic undocked (DU, bottom) traces in the presence of 1 mM Mg^{2+}. The stable docked (SD) conformation appears only in the presence of Mn^{2+}. Interestingly loses the WT’s ability to form the SD conformation, with the histogram showing a major single peak around 0.19 ± 0.13 FRET, further corroborated by a single on-diagonal ~0.2 FRET contour in the TODP (Fig. 6d). These data demonstrate that the conserved discriminator A48 is essential for Mn^{2+} inducing a SD riboswitch conformation. The presence of Mg^{2+} partially recovers the loss of Mn^{2+} sensing by the mutant, yet does not restore its ability to form the SD conformation.

Fig. 5 Effect of Mn^{2+} on the folding of the riboswitch. a Representative smFRET trace, FRET histogram and TODP for the WT riboswitch in the presence of 0.1 mM Mn^{2+} and 1 mM Mg^{2+}. The stable docked (SD) conformation appears only in the presence of Mn^{2+}, as highlighted by the dashed yellow box in the TODP. b FRET histograms for the WT riboswitch at 0.1 mM concentration of different transition metals in the presence of 1 mM Mg^{2+}. Over 150 molecules were analyzed for each histogram. The corresponding TODPs are shown in Supplementary Fig. 26. c FRET histogram and TODP with SD population highlighted by dashed yellow box for the WT riboswitch in the presence of 0.1 mM Mn^{2+} alone. d Example smFRET traces for the WT riboswitch showing transitions between SU, DU, DD, and SD traces upon addition of 0.4 mM Mn^{2+}. The time of Mn^{2+} addition is indicated by the red arrow. e Exemplary trace showing conversion of a DU trace to DD trace upon chelation of Mn^{2+} with EDTA (indicated by the red arrow) and reintroducing 1 mM Mg^{2+}. Source data for panels (e) and (f) are provided as a Source Data file.
Possibly this reflects the ability of Mg\(^{2+}\) to bind at M\(_B\) in A48U, though with weaker affinity, via the uridine O4, as suggested in the A41U L. lactis structure\(^{12}\).

**Mn\(^{2+}\)-binding enhances the stability of switch helix P1.1.** The proposed mechanism of transcription antitermination of the yybP-\(ykoY\) riboswitch, supported by our structural and dynamic data, involves Mn\(^{2+}\)-mediated stabilization of the inherently weak P1.1 switch helix. To more directly test this prevailing hypothesis, we probed the accessibility of the 3’-half of the P1.1 hairpin (last 12-nt) in trans by a complementary Cy5-labeled DNA oligonucleotide probe using our established single molecule kinetic analysis of RNA transient structure (SiM-KARTS) assay (Fig. 7a)\(^{36}\). In the current configuration, SiM-KARTS mimics the strand invasion by the downstream terminator sequence and reports on the stability of P1.1 helix at the single-molecule level. In the presence of 10 mM Cy5-labeled oligonucleotide and 1 mM Mg\(^{2+}\), repeated short binding events are observed with a probe binding rate constant \(k_\text{on}\) of 9.67 ± 1.24 \(\times\) 10\(^6\) M\(^{-1}\) s\(^{-1}\) and a dissociation rate constant \(k_\text{off}\) of 5.88 ± 0.35 s\(^{-1}\) for the WT riboswitch (Fig. 7b, c). Addition of increasing concentrations of Mn\(^{2+}\) results in a decrease in the frequency of binding events, with a \(k_\text{on}\) value significantly reduced by ~2.5-fold in the presence of 200 μM Mn\(^{2+}\) (Supplementary Figs. 28 and 29). These data report on a decreasing accessibility of the 3’-segment of the P1.1 helix and thus a more stable P1.1 duplex in the presence of Mn\(^{2+}\). Interestingly, the \(k_\text{off}\) value also decreases with the addition of Mn\(^{2+}\), consistent with the notion that the RNA can still sense Mn\(^{2+}\) via interaction of the free 5’-half of P1.1—extending into the L1 loop—with loop L3, resulting in a stable L1 and P1.2 that can stack on top of the RNA–DNA hybrid. Remarkably, the effect of Mn\(^{2+}\) is completely lost in the A48U mutant, where the \(k_\text{on}\) and \(k_\text{off}\) values remain essentially unchanged at 8.24 ± 0.30 \(\times\) 10\(^6\) M\(^{-1}\) s\(^{-1}\) and 4.54 ± 0.41 s\(^{-1}\), respectively, even at high (500 μM) Mn\(^{2+}\) concentration, showing no change in the frequency of individual binding events (Fig. 7d, e and Supplementary Figs. 28 and 30). These data demonstrate that the stability of P1.1 is enhanced once the riboswitch encounters Mn\(^{2+}\), an effect lost upon mutation of the Mn\(^{2+}\) sensing adenosine. These results thus provide a direct evidence for the prevailing mechanistic hypothesis of riboswitching by the yybP-\(ykoY\) RNA, wherein Mn\(^{2+}\) binding stabilizes the antiterminating P1.1 switch helix through long-range signal transduction from the metal ion sensing core, thereby preventing formation of the competing terminator hairpin.

**Discussion**

Using a combination of X-ray crystallography, MD simulations and smFRET, our work here sheds light on how the binding of a single Mn\(^{2+}\) ion at a distant site can modulate the stability of the
Fig. 7 Accessibility of switch helix P1.1 probed using a SiM-KARTS assay. 

a Schematic of our SiM-KARTS assay to probe the accessibility of switch helix P1.1 in the presence and absence of Mn²⁺.

b Representative SiM-KARTS traces at different Mn²⁺ concentrations show multiple binding events of the Cy5-labeled DNA oligonucleotide probe. Green, Cy3 intensity; black, Cy5 intensity; red, idealized Cy5 intensity fitting to a two-state model.

c The association ($k_{on}$, red) and dissociation ($k_{off}$, black) rate constants for binding of probe to P1.1 as a function of Mn²⁺ concentration for the WT riboswitch. The errors for the rate-constants are obtained using a boot-strapping program, as described in Methods.

d, e Same as in (b) and (c), but for the A41U mutant riboswitch. Over 200 traces were analyzed for each condition to obtain reliable kinetic rate constants. The data for kinetic plots in (c) and (e) are provided in the Source Data file.
switch helix through a conformational cascade that couples local with global structural dynamics. While previous structures revealed the local Mn$^{2+}$ recognition mechanism of yybP-ykoY RNA$^{12,19}$, the structures reported here underscore the conformational plasticity around the metal ion binding core. Importantly, the structural differences observed between the two conformers in the crystal lead us to posit that Conformer 2 with L3 partially unfolded around M$_{B,Mn}$ is likely an intermediate between the fully Mn$^{2+}$-bound state and the empty M$_{B,Mn}$ state observed in the *E. coli* Mn$^{2+}$-free structures$^{12}$. We hypothesize that the loss of phosphate contacts from C51, U52, and G9 to Mn$^{2+}$ in Conformer 2 reflects their relative weakness as supported by MD simulations and likely represents a discrete step in Mn$^{2+}$ binding/unbinding. Conversely, the variation in M$_{B,Mn}$ site coordination we observe could represent inherent flexibility at this site, allowing it to recognize Mn$^{2+}$ even if the metal ion is not fully dehydrated. This flexibility is also seen in the recent Cd$^{2+}$-bound structures$^{19}$ (see Supplementary Discussion).

Our MD simulations show that the SRL-like motif in L1 is inherently unstable in the L1–L3 undocked state, but is stabilized by the tertiary A-minor interaction of A10 with P3.1. Similarly, L1–L3 docking stabilizes L3 in a stacked conformation by direct stacking of A10 on top of it. Thus, a stable conformation of L1 and therefore P1.1 is intimately linked to the stability of L3 and the tertiary A-minor interaction. We suggest that the SRL-like motif acts as a molecular switch that links the Mn$^{2+}$-dependent increase in docking observed by smFRET to stabilization of P1.1.

The observation of pairs of stably and dynamically docked and undocked states with double-exponential kinetics supports the existence of at least two undocked and docked dynamic (in addition to the static SU and SD) conformations, with similar FRET values and thus global folds, but likely different local conformations around the ligand sensing core. These conformations are possibly related to Conformers 1 and 2 in the crystal structure. We further find that SD conformations are formed only in the presence of submillimolar Mn$^{2+}$ or the related soft transition metal ion Cd$^{2+}$, and lost upon a single mutation of the invariant Mn$^{2+}$-sensing adenine A48. While our work was in preparation, another smFRET study on the folding of a different *Llac* Mn$^{2+}$ riboswitch became available. Using a similar labeling strategy, this *Llac* riboswitch showed two FRET states similar to the *Xory* riboswitch$^{37}$. Interestingly, the kinetics of conformational dynamics for the *Llac* riboswitch were largely single-exponential, suggesting a two-state folding behavior, that agrees with a single conformation of the ligand-bound structure. This is in contrast to the *Xory* riboswitch studied here, which shows double-exponential kinetics, consistent with the two distinct conformations of the ligand-bound structure seen in our crystal structures. Such species-specific differences in the Mg$^{2+}$ and Mn$^{2+}$ mediated folding of the yybP-ykoY riboswitches likely enable bacteria to elicit a fine-tuned response to changes in their intracellular metal ion concentrations.

Our structural snapshots, MD simulations and smFRET data support the long-range structural signal transduction model shown in Fig. 8. In the absence of any divalent ions, the *Xory* riboswitch adopts an extended conformation in which the two legs are distal and rarely interact, as shown by smFRET. Under these conditions, the metal-ion binding sites are likely unstructured and their interaction only transiently formed by L1–L3 contacts, as suggested by rare smFRET transitions, MD simulations, as well as previous ligand-free structures$^{12}$. MD simulations in particular show that the A-minor interaction formed by A10 of...
L1 loop with the G–C base pair of stem P3.1 can at least transiently form even in the absence of divalents. The formation of this A-minor interaction helps pre-organize the M_{\text{AMg}} binding site. Under physiological (1 mM) concentrations of Mg^{2+}, this ion binds first at the M_{\text{AMg}} site, allowing the riboswitch to sample dynamic folded conformations where the two legs are brought together via metal ion-mediated interaction between loops L1 and L3. These dynamic folded states then allow for local interactions mediated by L3, forming a pocket of high negative charge potential poised to sense Mn^{2+}. Mn^{2+} binding at M_{\text{AMg}} therefore facilitates binding of Mn^{2+} at M_{\text{AMn}} in cooperative fashion, as suggested by smFRET at the global level and further visualized by atomistic MD simulations. Capture of Mn^{2+} at site M_{\text{AMn}} then acts as the final linchpin to hold the two legs together most stably. This compact docked conformation with a SRL-like fold of L1 and continuous coaxial stacking between P1.2, L1, and P1.1 stabilizes the remote P1.1 switch helix, as shown by our SIM-KARTS assay and MD simulations, that prevents the strand invasion required to form the terminator hairpin, instead promoting transcriptional read-through.

Contrasting with the Mn^{2+} riboswitch, the NiCo riboswitches, which cooperatively sense the transition metals Ni^{2+} or Co^{2+}, resemble the overall “H” shaped architecture of the Mn^{2+} riboswitch but with a tight 4WJ that lacks an analogous tertiary docking interface. Accordingly, the NiCo riboswitch appears to act through a distinct mechanism utilizing four bound metal ions to weave together a network of interactions between the interhelical residues that stabilize the 4WJ and so prevent formation of a terminator via strand invasion. Similarly, the Mg^{2+} sensing M-box riboswitch adopts an architecture with three parallel coaxially stacked helices that are brought together by the binding of six Mg^{2+} ions, leading to formation of a terminator. Therefore, both the NiCo and Mg^{2+} sensing riboswitches achieve gene regulation by employing multiple metal ions that directly interact with and stabilize the switch helix. In contrast, the Mn^{2+} riboswitch is unique in requiring only a single-metal ion that does not make any direct contacts with switch helix P1.1. By combining high-resolution structural information with insights into both atomistic and global dynamics, our study outlines the mechanism by which a ligand as small as a single divalent metal ion couples local with global structure to give Mn^{2+} the ability to influence RNA folding and fine-tune gene expression by cooperating with a high background of Mg^{2+}. The general lessons revealed here of how a ligand-binding signal can be transduced across an RNA are likely to become a recurring theme among riboswitches where the ligand represents a distal structural linchpin for the switch helix.

**Molecular dynamics simulations.** The crystal structures of yfbY-ykoY riboswitch from *X. oryzae* from this study (both Conformers 1 and 2) and from *L. lactis* (PDB ID 6CB3 and PDB ID 6C6B) were used as starting structures for MD simulations. We studied the effect of divalents in the metal ion binding sites on the structural dynamics of the riboswitch, so we prepared a set of starting structures containing different ions (Mn^{2+}, Mg^{2+}, and K^{+}) in the M_{\text{AMg}} and M_{\text{AMn}} sites (in the case of 6CB3 chain A even the MC) in the crystal structures of this study, see Supplementary Table 1). In addition, we prepared 94 both for syn- and anti-conformations of A48 in Conformer 2 from *Xory* to verify refinement of the corresponding electron density as a syn-oriented nucleotide. Starting structures of the SRL-like motif lacking tertiary interactions with the rest of the aptamer were prepared from the crystal structures but entrapping only the P1.1, P1.2, and L1 segments without divalents. All MD simulations were carried out using pmemd.cuda (GPU code of AMBER 14 program package) with the ff99bsc0 force field. The simulation protocol was as follows. The structure with the RNA molecule and entrapped ions in the M_{\text{AMg}} and M_{\text{AMn}} (and in case of 6CB3 chain A even the MC) sites was immersed into a solvation box of SPC/E explicit water molecules. We added KCl salt excess corresponding to 150 mM KCl. We used following ionic parameters: K^{+} (r = 1.593 Å, ε = 0.4297 kcal/mol, q = 0.00295 kcal/mol, ε = 1.0460 Å, ε = 0.167 kcal/mol), and Cl^{-} (r = 1.0 Å, ε = 1.2 kcal/mol, q = 0.00295 kcal/mol, ε = 0.00295 kcal/mol). We used the previously determined ion binding site reported in this structure, see Supplementary Table 1). In addition, we prepared 94 for each specific starting structure four different ionic configurations in these ion binding sites (Supplementary Table 1). In the case of 6CB3 *L. lactis* structures we replaced the Ca^{2+} ions in the M_{\text{AMg}} or M_{\text{AMn}} sites by K^{+} monovalents so that we prepared for each specific starting structure four different ionic configurations in these ion binding sites (Supplementary Table 1). The solvated systems were equilibrated as follows. Geometrical optimization of hydrogen atoms was followed by optimization of waters and ions, while the position of the RNA molecule remained restrained. Subsequently, all RNA atoms were restrained and the solvent molecules with counter-ions were allowed to move after running a 500-ps long heating under NVT conditions. We relaxed the resulting second-ns equilibration by running an additional 100 ps. Finally, we relaxed the final density equilibration under NPT conditions for an additional 100 ps. The particle-Mesh Ewald method for treating electrostatic interactions was used, and simulation were performed under periodic boundary conditions in the NVT ensemble at 298.16 K using a weak coupling Berendsen thermostat with a coupling time of 0.2 ps. The SHAKE algorithm, with a 10–Å cut-off, was used to fix the positions of all hydrogen atoms, and a 0.0 A cut-off was applied to nonbonding interactions. Data analysis. All trajectories were analyzed with the P traj module of the AMBER package and the simulations were visualized using VMD. B-factors were calculated as average mass-weighted fluctuation of each residue over the entire trajectory aligned to the stem closest to the residue. This guarantees that the B-factors are not influenced by global motion and only local structural changes. The stacking interactions were detected using the G-vector of an eRMSD metric and were annotated using barNAb}.
RNA preparation for single-molecule FRET. The RNA for smFRET studies was annealed from two synthetic RNA oligonucleotides (Supplementary Fig. 23). Oligonucleotides are the modifications from the native 5′-Cy5 and 3′ biotin-TEG. In this two-oligonucleotide design of the smFRET construct, the donor fluorophore Cy3 and biotin-TEG were placed on the 5′ and 3′ ends of stem P1.1 while the acceptor fluorophore Cy5 docked to the 5′ end of P3.2 in the presence of MgCl₂ on either end of the distal arms that dock. For specifically probing the docking interaction, we extended the stem P1.1 to prevent end fraying. The biotin on the 3′-end of P1.1 was used for immobilization of the RNA onto a microscope quartz slide and subsequent internal reflection fluorescence (TIRF) microscopy (Fig. 4a). Optimal refolding of the RNA was achieved with a mixture of 1 µM oligonucleotide 1 and 1.5 µM oligonucleotide 2 in 20 mM HEPES pH 7.0, 50 mM MgCl₂ and under varying concentrations of the metal ions in the 1× buffer with OSS also used for smFRET. The 5′ Cy5-labeled DNA oligonucleotide is complementary to the 12-nt sequence (5′-GAGACACAGGGAAU-3′) of the 3′-half of L1 and P1.1 in the smFRET RNA construct (Supplementary Fig. 23).

smFRET data acquisition and analysis. A small microfluidic channel was made by sandwiching a cleaned quartz slide and a glass coverslip using double-side tape. Beforehand, two holes were drilled into the quartz slide and afterwards tubing was attached to them to act as inlet and outlet ports that were used for flowing buffer and introducing the RNA. The two RNA oligonucleotides containing the Cy3, Cy5 and biotin-TEG arms were mixed at a concentration of 2 µM each in 1× HEPES, pH 7.2, 100 mM KCl (1× buffer). The RNA was heated at 90 °C for 2 min, left at RT for 30 s to cool down, followed by addition of 2 mM MgCl₂ and slow cooling down to RT over 15 min. This annealed RNA stock was used for carrying out smFRET experiments, for which a 15–25 pmol RNA solution was made by diluting the 2 µM annealed stock using 1× buffer.

100 µL of diluted RNA solution was flowed onto the quartz slide coated with biotinylated-bovine serum albumin and streptavidin for immobilization and incubated for 2 min. Any unbound RNA was washed off using 1× buffer. This step also washes off small (low nM) amounts of residual Mg⁺⁺ present in the stock. For experiments in the absence of divalents, 0.1× EDTA was included in the 1× buffer to chelate any contaminating divalent ions. Mg⁺⁺ titration experiments were performed on the same slide after washing off the EDTA using 1× buffer. We included an enzymatic oxygen scavenging system (OSS) containing 50 mM protocatechuate-3,4-dioxygenase and 5 mM protocatechuic acid in 1× buffer to prolong the longevity of the fluorophores. In addition, 2 mM Trolox (6-Hydroxy-2,5,7,8-tetramethylchromane-2-carboxylic acid) was included in the imaging buffer to suppress photobleaching of the dyes.

All smFRET movies were acquired at ~16 Hz, unless otherwise specified, using a prism-based TIRF microscope with an intensified CCD (P-10, Princeton Instruments) or sCMOS camera (Hamamatsu ORCA-Flash4.0 V3), essentially as previously described. 3,4 Cy3 on the RNA was excited using a 532 nm laser and the emission from both Cy3 (donor) and Cy5 (acceptor) were simultaneously detected side-by-side on the camera. Toward the end of all movies, the Cy3 was directly excited using a 640 nm laser to check for the presence of acceptor (Cy5) fluorophore. This helps in distinguishing those with a minimum combined intensity (Cy3 + Cy5 intensity) of 300, showing single-step photobleaching of the dyes, a signal-to-noise ratio of >3, and longer than 6 s were selected for further analysis. Selected traces were then background-subtracted to correct for cross-talk and (minimal) bleed-through. We calculated the FRET ratio as I3/(I3 + I5), where I3 and I5 are the background corrected intensities of the acceptor (Cy3) and donor (Cy5), respectively. FRET analyses were made using the first 50 frames (3 s) of all traces at least 200 in a given condition and fit with a sum of Gaussians using OriginPro 8.5. For kinetic analysis, traces were idealized with a two-state model corresponding to undocked (low-FRET) and docked (high-FRET) states using the segmental k-means clustering algorithm.

Beforehand, two holes were drilled into the quartz slide and afterwards tubing was added, samples were allowed to cool to RT for 10 min, then put on ice until further use. To check assembly, ~1 pmol of each oligonucleotide (1 µM for crystallization was present in the smFRET RNA. Oligonucleotide 1 has a 5′-Cy5 and oligonucleotide 2 has 5′-Cy3 and 3′ biotin-TEG. In this two-oligonucleotide design, we used the nanosecond k-means clustering algorithm to identify subpopulations of traces. In addition, the TODPs shown here also include static traces that appear as on-diagonal contours and are often critical in understanding the molecular mechanisms of riboswitches. 6,7 The on-diagonal contours in the TODPs (at −0.2 and −0.7 FRET values) thus do not represent any transitions, but rather highlight static traces that lack any FRET transitions over their entire observation length and are counted exactly once in the TODP heatmap.

SiM-KARTS assay. The SiM-KARTS assay to probe P1.1 accessibility was performed on a prism-based TIRF microscope. 8 Totally ~20 pm of the riboswitch was bound to the slide and the Cy5 on the riboswitch was first photo-bleached by excitation with a red laser. Kinetic measurements of P1.1 accessibility were performed in the presence of 10 nM of a Cy5-labeled DNA oligonucleotide and under varying concentrations of the metal ions in the 1× buffer with OSS also used for smFRET. The 5′ Cy5-labeled DNA oligonucleotide is complementary to the 12-nt sequence (5′-GAGACACAGGGAAU-3′) of the 3′-half of L1 and P1.1 in the smFRET RNA construct (Supplementary Fig. 23). Movies were acquired at 100 ms time resolution under dual excitation with both a green and red laser to localize the Cy3-labeled surface-bound riboswitch molecules and to image the short binding events of the Cy5-labeled oligonucleotide probe, respectively. The Cy5 versus intensity time traces were idealized to a two-state model using the segmental k-means algorithm implemented in QuB to obtain the bound and unbound dwell times. The unbound and bound lifetimes were obtained by single-exponential fits to the cumulative dwell time distributions, the inverse of which gives the rate constants kon and koff, respectively. The errors of the rate constants were estimated using a bootstrapping algorithm by taking three random subsets from all the traces in a condition and calculating the rate constants. The means and standard deviations from the three calculated rates are reported in Fig. 7.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. A reporting summary for this Article is available as a Supplementary Information file. Atomic coordinates and structure factors for the reported crystal structure have been deposited with the Protein Data Bank under accession number 6N2V. The source data underlying Figs. 4e, h and 7c, e are provided as a Source Data file. The data supporting the findings of this study are available from the corresponding authors upon reasonable request.

Code availability. Custom scripts used to analyze the smFRET and MD simulations data can be made available upon request.

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K.C.S., I.R.P., S.S.D., A.K. and N.G.W. designed the experiments; P.B. and J.Š. designed the MD simulations; K.C.S. and S.S.D. performed the single-molecule experiments and analyzed the data; I.R.P. and A.K. performed the X-ray crystallography experiments and analyzed the data; M.J., P.K. and P.B. performed and analyzed the MD simulations; K.C.S., I.R.P., A.K., P.B. and N.G.W. wrote the bulk of the paper. All authors read and commented on the paper.

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