Interrogating RNA—Small Molecule Interactions with Structure Probing and Artificial Intelligence-Augmented Molecular Simulations

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ABSTRACT: While there is increasing interest in the study of RNA as a therapeutic target, efforts to understand RNA-ligand recognition at the molecular level lag far behind our understanding of protein—ligand recognition. This problem is complicated due to the more than 10 orders of magnitude in time scales involved in RNA dynamics and ligand binding events, making it not straightforward to design experiments or simulations. Here, we make use of artificial intelligence (AI)-augmented molecular dynamics simulations to directly observe ligand dissociation for cognate and synthetic ligands from a riboswitch system. The site-specific flexibility profiles from our simulations are compared with in vitro measurements of flexibility using selective 2′-hydroxyl acylation analyzed by primer extension and mutational profiling (SHAPE-MaP). Our simulations reproduce known relative binding affinity profiles for the cognate and synthetic ligands, and pinpoint how both ligands make use of different aspects of riboswitch flexibility. On the basis of our dissociation trajectories, we also make and validate predictions of pairs of mutations for both the ligand systems that would show differing binding affinities. These mutations are distal to the binding site and could not have been predicted solely on the basis of structure. The methodology demonstrated here shows how molecular dynamics simulations with all-atom force-fields have now come of age in making predictions that complement existing experimental techniques and illuminate aspects of systems otherwise not trivial to understand.

1. INTRODUCTION

RNA regulates diverse cellular processes, far beyond coding for protein sequence or facilitating protein biogenesis. 1 In this role, RNA can interact with proteins 2 or small molecule factors 3,4 to directly or indirectly impact gene expression and cellular homeostasis. 5 For example, riboswitches are sequences found primarily in bacterial mRNAs that contain aptamers for small molecules and regulate transcription or translation. 6 In humans, mutations in noncoding RNAs can directly impact neurodegenerative diseases and cancer. 7,8 The regulatory role of bacterial riboswitches, coupled with the broader link between RNA and human disease, has also led to interest in RNA as a therapeutic target for small molecules. 9−11 However, in comparison to proteins, methods to computationally and experimentally interrogate RNA structure are relatively less mature. 12,13 Thus, more robust tools are needed to better understand RNA structure, dynamics, and recognition of small molecules. In spite of its clear relevance to fundamental science and drug discovery efforts, the highly dynamic nature of RNA makes it difficult to study due to the diverse conformational ensembles it adopts. 14−16 Luckily, riboswitches represent a valuable model for RNA-small molecule recognition. When ligands bind to riboswitch aptamers, the RNA undergoes a conformational change that modulates gene expression. 17,18 The cognate ligand for a riboswitch is typically associated with its upstream gene, often part of the ligand’s biosynthetic pathway. 18,19 This feedback mechanism, paired with well-established structural analyses, makes riboswitches excellent models for RNA-ligand binding since the target ligand is known and often highly specific. 18

Although we can use riboswitches to model RNA-small molecule interactions, visualizing riboswitch-ligand dissociation pathways with high spatiotemporal resolution remains a challenge for in vitro and in silico experiments. Such a femtosecond and all-atom resolution is hard to directly achieve in in vitro experiments, while the associated time scales are several orders of magnitude too slow for molecular dynamics (MD) simulations performed even on the most powerful supercomputers. Previously, ref 20 used Gō-model simulations...
of PreQ₁ riboswitches, specifically transcriptional *Bacillus subtilis* (Bs) and translational *Thermoanaerobacter tengcongensis* (Tte) aptamers. Their work suggests that in respective cases the ligand binds at late and early stages of riboswitch folding, indicating that perhaps these riboswitches respectively fold via mechanisms of conformational selection and induced fit. Since ref 20, RNA force-fields have become more detailed and accurate, but the time scales involved in RNA-small molecular dissociation continue to stay beyond reach.21

In order to better understand the ligand-riboswitch dissociation process, in this work we employ a combination of (i) enhanced sampling methods combining statistical physics with artificial intelligence (AI)22−25 and (ii) experimental techniques to measure RNA flexibility and ligand binding.26−28 The computational sampling techniques allow us to study the dissociation process in an accelerated but controlled manner, while the selective 2′ hydroxyl acylation analyzed by primer extension and mutational profiling (SHAPE-MaP), microscale thermophoresis (MST), and fluorescence intensity assay (FIA) provide a rigorous validation of the computational findings. Specifically, we study the Tt-PreQ₁ riboswitch aptamer interacting with its cognate ligand PreQ₁ (7-aminomethyl-7-deazaguanine) and a synthetic ligand (2′-[(dibenzo[b,d]furan-2- yl)oxy]-N,N-dimethylethan-1-amine).29 See Figure 1 for an illustration of the riboswitch and chemical structures of the two ligands. First, we perform 2 μs long unbiased MD simulations of the ligand-bound and ligand-free systems to quantify the flexibility of individual nucleotides. Comparing the flexibility profiles from unbiased MD with SHAPE-MaP reveals some broad overall agreement; however, a more detailed quantitative analysis reveals the limitations of both approaches. To simulate the dissociation process, which is too slow for MD simulations,21 a key challenge in most enhanced sampling methods is the need for an *a priori* estimate of the dominant dissociation mechanisms as captured through the reaction coordinate (RC). However, it is very challenging to estimate such an RC without already having simulated the dissociation process, especially for flexible biomolecules like RNA.30 Here, we use our AI-based sampling method RAVE to directly observe the dissociation of both of these ligands from the wild-type (WT) Tt-PreQ₁ riboswitch. RAVE automates the learning of RCs through a scheme that iterates between MD to generate data and deep learning to construct an approximate RC from this data. The MD-deep learning iteration continues until we see no further improvement in accelerating the dissociation events as measured through typical MD time to observe dissociation. The RC itself is learned as the most informative low-dimensional representation expressed as a past-future information bottleneck.31

By efficiently sampling multiple independent riboswitch-ligand dissociation events for both cognate and synthetic ligands, we are able to reproduce relative binding affinity measurements from MST and FIA and give a qualitatively reasonable fit to flexibility measurements from SHAPE-MaP experiments, though the limitations are also strikingly obvious. Furthermore, we are able to pinpoint which nucleotides play the most critical roles in dissociation. These are found to be the nucleotides U22 and A32 for the cognate and synthetic ligands, respectively. On the basis of these, we design and perform a set of *in vitro* mutagenesis experiments, which validate our mutation predictions. We note that these predicted mutations are unlikely to have been found simply from a structural perspective, as they are distal from the binding site and do not directly interact with either of the ligands. Our findings thus reveal the complex nature of the interaction between RNA and small molecules, and might help guide the design of RNA-binding small molecules in other systems in the future. Thus, this methodology will be broadly useful in understanding increasingly complex, biologically important RNA-small molecule interactions.

### 2. RESULTS

#### A. Ligand Binding Induces Flexibility Change in Riboswitch Aptamer

Our first test is to compare the nucleotide-specific flexibility from all-atom simulations to those from SHAPE-MaP measurements. We perform 2 μs unbiased MD simulations (4 independent sets of 500 ns each for both ligands) each for PreQ₁ riboswitch bound with the cognate and synthetic ligand, respectively. See Supporting Information for details of the simulation setup. While these simulations are not long enough to observe ligand dissociation, they can still give a robust measure of how the flexibility varies between the two different ligand-bound systems and provides nucleotide-level insight into how specific ligand binding induces different conformation changes in the riboswitch. Specifically, we compare the fluctuations of the distances between consecutive C2 atoms obtained from these simulations with SHAPE-MaP measurements (see Supporting Information for details of SHAPE-MaP). We performed SHAPE-MaP analyses using a PreQ₁ riboswitch construct consisting of the entire riboswitch (both the aptamer domain and expression platform). Probing was performed using a recently reported SHAPE reagent, 2A3, that provides improved mutagenesis rates during reverse transcription.32 We analyzed data in the absence of any ligand, and in the presence of either cognate ligand (PreQ₁) or synthetic ligand. Our results were in good agreement with previous studies reporting SHAPE-MaP on PreQ₁ riboswitches using different SHAPE reagents and a slightly shorter construct.33 As discussed

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Figure 1. Conformational change of the PreQ₁ riboswitch and chemical structures of PreQ₁ and synthetic dibenzofuran ligands.
previously, classical force-fields accurately model the behavior of complex RNAs and highlights the need for continued method development from both the computational and experimental sides that can facilitate more direct comparisons of nucleotide-dependent flexibility. The lack of unequivocal quantitative agreement could be due to many reasons. First, while SHAPE is a measurement of flexibility, it is strictly speaking a surrogate measurement relating to the kinetics of acylation of each nucleotide. Although flexibility of each nucleotide is a main factor in determining the probability of a nucleotide being acylated, there are other factors that can determine how the SHAPE profile looks like. Second, our unbiased simulations are not long enough to reduce the statistical error. Finally, as pointed out by the reviewer, we only simulated the aptamer domain of the PreQ₁ riboswitch, while in the experiment the SHAPE reactivity of the whole PreQ₁ riboswitch was measured. The presence of other domains may also contribute to flexibility changes in the system.

B. Reaction Coordinates and Approximate Free Energy Profile. Encouraged by the agreement between flexibility profiles from unbiased MD and SHAPE, we next aim to calculate the free energy profiles and reaction coordinates for ligand dissociation from PreQ₁. As these time scales are far beyond unbiased MD, here we use our recent AI-augmented MD method RAVE which learns the dominant slow degrees of freedom on-the-fly by iterating between rounds of sampling and AI. See Supporting Information for further details of the protocol. After four rounds of such iterations, we obtain converged RCs for both systems, expressed as a linear combination of different riboswitch-ligand heavy atom contacts.

Figure 3 shows different contacts and their weights as obtained from AMINO and RAVE simulations. It is interesting to note that the RC for the cognate ligand has many more components than the RC for synthetic ligand, highlighting differences between the molecules. Most of these contacts in the RC are present also in the crystal structure (PDB 6E1W, 6E1U), except carbon S-U2, carbon S-A23 for the cognate ligand, and carbon 1-A23 for the synthetic ligand (ligand atom numbering corresponds to numbering in the PDB files). We would like to highlight here that AMINO provides a basis set for RAVE to build the reaction coordinate as a function thereof, and the precise output of AMINO could differ depending on which precise trajectory was used with it. However, as shown previously in ref 22, we do not expect the final result from RAVE to significantly vary with the precise basis set fed into RAVE.

We then used this optimized RC and ran 10 independent biased MD simulations each using well-tempered metadynamics to sample the free energy profiles of the systems. The probability distribution was estimated by using the reweighting technique introduced in ref 36. In Figure 4, we provide the free energy profiles as functions of the ligand-riboswitch coordination number for both systems. Note that for both systems there were two predominant pathways as indicated through arrows in Figure 5. A key difference between the two paths is that in the first path the ligand unbinds through the space in between stem 2 and the backbone connecting stem 1 and loop 1 (Figure 5, blue arrows). In the second path, the ligand unbinds through the space in between loop 2 and stem 2 (Figure 5, red arrows). In order to compare the binding free energies of the two systems, we combine the information from both of the pathways by plotting the free energy as a function of the ligand-riboswitch shared coordination number C. During the dissociation process, the coordination number gradually decreases to 0 irrespective of

Figure 2. SHAPE reactivities for the Tt-PreQ₁ riboswitch aptamer obtained using the 2A3 reagent (red circles joined by solid lines) are compared with the fluctuations of C2-C2 distances (blue triangles joined by dashed lines) for the PreQ₁ riboswitch (a) in the absence of ligand, (b) bound to cognate ligand, and (c) bound to synthetic ligand. Pearson correlation coefficients $R$ are shown in upper left corners.
where $N$ and $M$ are the number of ligand and riboswitch atoms, respectively. $r_{ij}$ is the distance between ligand atom $i$ and riboswitch atom $j$ and $r_0$ is set to be 3 Å. It can be seen from Figure 4 that when bound to the riboswitch the cognate ligand has a higher $C$ relative to when the synthetic ligand is bound to it, illustrating that the synthetic ligand makes overall fewer contacts per ligand atom with the riboswitch, as also observed in crystal structures (PDB 6E1W and 6E1U). Though we were not able to simulate the rebinding of ligands to get a converged free energy profile, the probability distributions used to construct these free energies are obtained by averaging over multiple independent dissociation events. This leads to the relatively small error bars shown in Figure 4, which indicates that we can use the free energy profile to give a rough estimation of the strength of the binding affinity. By considering $C \lesssim 5.6$ as the unbound states, we can calculate that the binding affinity difference between bound and unbound state is 2–4 kcal/mol higher for the cognate ligand than that for the synthetic ligand. We however caution the reader that this is an approximate free energy profile in the absence of strict rebinding events during metadynamics. In principle, specialized methods can be used to address this limitation and will be the subject of careful investigation in future work. This is in quantitative agreement with relative affinity measurements ($3.0 \pm 1.4$ kcal/mol) through fluorescence titrations as reported in ref 29.

C. Predicting Critical Nucleotides for Mutagenesis Experiments. We now analyze our 10 independent dissociation trajectories each obtained for the synthetic ligand bound and cognate ligand bound systems to predict critical nucleotides in the riboswitch that play a role in the dissociation process. We are interested in not just nucleotides we could have predicted from looking at the static structures of ligand-bound complexes but also in nucleotides distal from the binding site. Those nucleotides might play an essential role in determining the structural ensemble of a riboswitch. Thus, changing the properties of those nucleotides can alter the interaction between ligand and other nucleotides. Such critical nucleotides are hard to predict from a purely structural perspective, but our all-atom resolution dissociation trajectories are ideally poised to make such predictions. Predictions made in this section are then validated through mutation experiments reported in section 2D.

From our collection of dissociation trajectories, we observe that dissociation mainly happens through two pathways (Figure 5) and that for the cognate ligand, 7 out of 10 simulations went through the top/blue pathway, while for the synthetic ligand, 6 out of 10 simulations went through the bottom/red pathway. These observations indicate how two ligands interact with the riboswitch differently before fully dissociating from the pocket. To more rigorously quantify the role played by different nucleotides, we monitor the change in the RMSD of every nucleotide in the enhanced MD simulation. In principle, a comprehensive study of ligand–receptor interactions should also include simulations of the rebinding events. As previously stated, that will require the implementation of specialized methods and thus is not explored in this study. Nonetheless, dissociation trajectories should be able to give a good sampling for bound state ensembles, allowing us to make predictions about the effects of mutations on binding affinity. Our motivation here is that the nucleotides showing greatest relative movement during the dissociation process are the ones most likely to be impacted once mutated. As shown in Supplementary Figure 2, every nucleotide’s RMSD changes with the coordination number $C$, which captures the extent of
dissociation. These changes indicate different conformations the riboswitch adopts during ligand dissociation. Through this procedure, we identify U22 and A32, highlighted in Figure 5, to be two such critical nucleotides for the cognate and synthetic ligands respectively (see Supporting Information for further details of this protocol). Figure 6 shows the changes in the RMSDs of these two selected nucleotides with different coordination numbers calculated from the enhanced MD trajectories. For U22, in RAVE simulations the RMSD increases much more for the cognate ligand-bound system than for the synthetic ligand-bound system. However, the reverse is found to be true for the nucleotide A32, which displays more enhanced movement during the dissociation of the synthetic ligand bound complex relative to the cognate ligand bound complex.

On the basis of the above observations, we can predict that for the cognate ligand, mutating U22 should display more tangible effects on the strength of the complex than mutating A32. For the synthetic ligand, we predict an opposite trend—mutating A32 should have a more pronounced effect on the interaction relative to mutating U22.

D. Validating Predicted Critical Nucleotides through Mutagenesis Experiments. In section 2C, we predicted on the behalf of our dissociation trajectories that mutating nucleotides U22 and A32 will have differing effects on the cognate ligand-bound and synthetic ligand-bound systems. Also, we would like to note that, as we can see from Figure 6, the RMSD changes appear in the range of coordination numbers that correspond to the bound state. These changes imply the role of these nucleotides in determining the bound state interactions and suggest that the effects of mutations can be reflected by changes in binding affinity. Here we report in vitro tests of our predictions. As shown in Figure 7, we measure the equilibrium dissociation constant \( K_D \) for six different complexes, namely, the two wild-type complexes, U22A-cognate ligand, U22A-synthetic ligand, A32U-cognate ligand, and A32U-synthetic ligand. We also performed in silico mutagenesis experiments to validate that the mutations change the bound state ensemble of both systems.

To experimentally test whether the A32U and U22A mutations impacted \( K_D \) values, we used microscale thermopho-
K\textsubscript{D} values were measured using the fluorescently labeled wild type, A32U, and U22A PreQ\textsubscript{1} riboswitch aptamer constructs for the cognate ligand. For PreQ\textsubscript{1}, we observed a K\textsubscript{D} of 11.7 ± 0.35 nM for the WT construct. In contrast, the A32U mutant had a K\textsubscript{D} of 22.5 ± 0.42 nM, while the U22A mutant had a K\textsubscript{D} of 49.4 ± 1.5 nM. For the synthetic ligand, changes were observed in fluorescence of the labeled RNA upon incubation with the ligand, and thus a fluorescence intensity assay (FIA) was used to measure K\textsubscript{D} values rather than MST. Using FIA, we observed a K\textsubscript{D} of 1.6 ± 0.7 μM for the wild type aptamer. In contrast, both the U22A and A32U mutant constructs displayed significantly weaker K\textsubscript{D} values (150 ± 1.3 μM, and ≥ 100 μM respectively). These K\textsubscript{D} measurements directly validate the findings from section 2C. There we had predicted that mutating U22 will effect the cognate ligand-bound system more than mutating A32, and indeed we find that K\textsubscript{D} for U22A is five times weaker than WT, while only two times weaker for A32U. At the same time, we had predicted that mutating A32 will effect the synthetic ligand-bound system more than mutating U22, and indeed we find that, while the K\textsubscript{D} for U22A is 75 times weaker than WT, there is effectively no tangible association at all for A32U with the synthetic ligand.

The in silico experiments also support that the mutations alter the bound state ensemble and give more insights into the interaction change in the bound state after the mutations. As shown in Supplementary Figure 3, the free energy profiles of the bound state ensemble of the mutated system differ from that of the wild type. Further analyses of the hydrogen bond interactions between ligand and riboswitch show that even though nucleotides 22 and 32 do not directly form hydrogen bonds with both ligands in the bound pose as seen in the crystal structure, interactions between ligands and other nucleotides are still altered when they are mutated and can be seen in simulating the bound pose ensemble of states. See the Supporting Information for detailed discussions of these simulations.

3. DISCUSSION

Over recent years, a consensus view is emerging that a superior understanding of biomolecular structural dynamics, and not just static structures, can be fruitful in predicting key emergent biochemical and biophysical properties. However, in spite of the importance of the problem, tools to study biomolecular structural dynamics are still limited in their scope. This is true for proteins but even more so for RNAs, where it is especially challenging to determine the ensemble of possible structures that the RNA can adopt and pinpoint the specific importance of different structures. In this work, we demonstrate that this is indeed feasible—and how one can obtain detailed and robust insights into RNA structural dynamics through AI-augmented
molecular simulations that maintain all-atom resolution for RNA, ions, and all water molecules. All our predictions are validated through a gamut of complementary and experimental techniques that measure RNA flexibility and ligand binding. We specifically focused on the Tte-PreQ1 riboswitch aptamer system in its apo state and bound to two different ligands, where we first showed that 2 µs long unbiased MD simulations with classical all-atom force-fields for RNA and water can provide the same flexibility profile as measured in SHAPE experiments. Since the time scales for ligand dissociation are far slower than MD can typically access, we used the RAVE simulation method that enhances sampling along a low-dimensional reaction coordinate learned on-the-fly through the past-future information bottleneck framework. Through RAVE, we obtain multiple independent dissociation trajectories for both cognate and synthetic liganded systems, demonstrating how the cognate and synthetic ligands invoke different aspects of the riboswitch’s flexibility in order to dissociate. On this basis, we were able to predict pairs of mutations which would be expected to show contrasting behaviors for the cognate and synthetic ligands. The ability to make such predictions accurately and efficiently is of vital importance, as mutations that can disrupt RNA structural dynamics and ligand recognition processes have been linked to numerous human diseases.

Our predictions for the Tte-PreQ1 aptamer were validated by designing mutant aptamers and performing affinity measurements.

Our work thus demonstrates a pathway to gain predictive insights, as opposed to retrospective validation, from enhanced molecular dynamics RAVE simulations that maintain all-atom resolution while directly observing processes as slow as ligand dissociation. While the focus in this work was on predicting the thermodynamically averaged aspects of riboswitch flexibility, in future work it should also be possible to directly calculate rate constants using complementary approaches and for more complicated RNA systems. More broadly, this work indicates that long time scale MD simulations can accurately model ligand dissociation from RNA and provide validatable hypotheses about structural features that impact small molecule recognition of RNA. Thus, this strategy will be widely applicable not only to study natural systems such as riboswitches but to more completely understand the structural features and dynamics that govern how synthetic small molecules interact with RNA.

**ASSOCIATED CONTENT**

Supporting Information

These include details of: The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscentsci.2c00149.

In *vitro* transcription of Tte-PreQ1 RNA; acylation of RNA *in vitro*; primer extension and mutational profiling (MaP) reverse transcription of RNA; SHAPE-MaP library preparation, sequencing, and data analysis; microscale thermophoresis; fluorescence intensity assay; MD simulation setup; flexibility measurement from MD simulations and comparison with SHAPE data; AI-augmented enhanced sampling method RAVE; network architecture and hyperparameter setting for enhanced sampling simulation; predicting critical nucleotides for mutagenesis experiments; *in silico* mutagenesis experiments (PDF)

Transparent Peer Review report available (PDF)

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**Author Contributions**

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**Notes**

The authors declare no competing financial interest. The SHAPE-MaP sequencing reads were deposited into the NLM/NCBI Sequence Read Archive (SRA) under the BioProject ID PRJNA767082. All PLUMED input files required to perform the enhanced MD reported in this paper are available on PLUMED-NEST (www.plumed-nest.org), the public repository of the PLUMED consortium, as plumID:22.013. The GROMACS input files and raw data to reproduce the simulation results in this paper can be found at go.umd.edu/wang_ACS_central_science_2022.

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