Mechanism of RNA recognition by a Musashi RNA-binding protein

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1. Introduction

Protein-RNA interactions play crucial roles in various cellular activities and their dysfunction leads to a wide range of human diseases (Wu, 2020; Hentze et al., 2018; Conlon and Manley, 2017). Identification of small molecules that modulate interactions between RNA-binding proteins (RBPs) and RNA is progressing rapidly. It represents a novel strategy for discovery of drugs with new mechanisms (Wu, 2020). The Musashi (MSI) RBPs have been shown to regulate translation of target mRNAs and participate in the maintenance of cell stemness and tumorigenesis. They have been suggested as potential drug targets for treating many types of human cancer, including acute myeloid leukemia, ovarian cancer, colorectal cancer and bladder cancer (Kudinov et al., 2017). The MSI protein family has two members: MSI1 and MSI2. Each MSI protein contains two N-terminal RNA recognition motifs (RRM1 and RRM2) that mediate the binding to their target mRNAs (Sakakibara et al., 2001). MSI1 binds to the 3'-untranslated region of Numb mRNA and represses its translation, which confers to the upregulation of Notch signaling. This leads to increased cell proliferation and survival, and decreased apoptosis of cancer cells (Kudinov et al., 2017). Understanding the molecular mechanism of MSI1-Numb RNA interaction is important in both basic biology and applied medical research.

Rational design of small molecules targeting protein-RNA interactions requires structural characterizations of the RBP-RNA complexes. Due to high flexibility of MSI proteins and lack of potent ligands, only a few MSI structures have been resolved so far, including the apo structure of MSI1/2-RRM1 (Lan et al., 2026; Lan et al., 2017; Nagata et al., 1999; Miyanoiri et al., 2003; Minuesa et al., 2019) and RNA-bound structure of MSI1 (Ohyama et al., 2011; Iwaoka et al., 2017). These structures have greatly facilitated structure-based modeling and drug design targeting the MSI-RNA interactions (Lan et al., 2015; Clingman et al., 2014; Lan et al., 2018; Lan et al., 2020). For example, we have recently identified one potent compound Aza-9 by combining fluorescence polarization (FP) assay, surface plasmon resonance (SPR), nuclear magnetic resonance (NMR) spectroscopy and molecular docking (Lan et al., 2020). However, experimental structures are rather static images and the dynamic mechanism of MSI-RNA interactions remains unknown, which has largely hindered the development of potent inhibitors of MSI proteins.

Molecular dynamics (MD) is a powerful technique that enables all-atom simulations of biomolecules. MD simulations are able to fully...
account for the flexibility of the RBP and RNA during their interactions (Cheatham III and Case, 2013; Sponer et al., 2017; Borisik et al., 2019). In 2015, Krepl et al. (Krepl et al., 2015) provided systematic benchmarking data by simulating six structurally diverse protein/RNA complexes over multiple microsecond timescale MD runs and evaluating the simulations’ stability. Their results suggested that current force fields are able to handle microsecond MD simulations of protein/RNA complexes in many cases. For most systems, MD was possible to achieve a good but imperfect agreement with the experimental structure. However, MD could not maintain the initial experimental structure in one among six cases (3K5Y). The same group further presented a joint MD and NMR study to interpret and expand the available structural data of two RBPs bound with their single-stranded target RNAs (Krepl et al., 2016). They collected more than 50 μs simulations and showed that the MD simulation was robust enough to reliably describe structural dynamics of RBP-RNA complexes (Krepl et al., 2016). However, due to the slow dynamics and limited simulation timescale, it is rather challenging for conventional MD (cMD) simulations to sufficiently sample RBP-RNA interactions and obtain proper free energy profiles to quantitatively characterize RBP-RNA interactions.

To overcome limitations of cMD, enhanced sampling methods have been developed to improve biomolecular simulations (Mlýnský and Bussi, 2018; Abrams and Bussi, 2014; Spiwok et al., 2015; Gao et al., 2019; Bussi, 2018; Abrams and Bussi, 2014; Spiwok et al., 2015; Gao et al., 2019). First, for any two arbitrary potential values \( E_1 \) and \( E_2 \) are automatically determined on three enhanced sampling principles. First, for any two arbitrary potential values \( E_1 \) and \( E_2 \) found on the original energy surface, if \( V_1(\tilde{r}) < V_2(\tilde{r}), \Delta V \) should be a monotonic function that does not change the relative order of the biased potential values; i.e., \( V_1'(\tilde{r}) < V_2'(\tilde{r}) \). Second, if \( V_1(\tilde{r}) < V_2(\tilde{r}) \), the potential difference observed on the smoothed energy surface should be smaller than that of the original; i.e., \( V_2'(\tilde{r}) - V_1'(\tilde{r}) < V_2(\tilde{r}) - V_1(\tilde{r}) \). By combining the first two criteria and plugging in the formula of \( V' \) and \( \Delta V \), we obtain:

\[
V_{\text{max}} \leq E \leq V_{\text{min}} + \frac{1}{k}
\]

(2)

Where \( V_{\text{max}} \) and \( V_{\text{min}} \) are the system minimum and maximum potential energies. To ensure that Eq. (2) is valid, \( k \) has to satisfy: \( k \leq 1/(V_{\text{max}} - V_{\text{min}}) \). Let us define: \( k = k_0 \times 1/(V_{\text{max}} - V_{\text{min}}) \), then \( 0 < k_0 \leq 1 \). Third, the standard deviation (SD) of \( \Delta V \) needs to be small enough (i.e. narrow distribution) to ensure accurate reweighting using cumulant expansion to the second order: \( \sigma_{\Delta V} = k(E - \bar{E}_{\text{avg}}) \sigma_{E} \leq \sigma_{0} \), where \( \bar{E}_{\text{avg}} \) and \( \sigma_{E} \) are the average and SD of \( \Delta V \) with \( \sigma_{0} \) as a user-specified upper limit (e.g., 10kBT) for accurate reweighting. When \( E \) is set to the lower bound \( E = V_{\text{max}} \) according to Eq. (2), \( k_0 \) can be calculated as:

\[
k_{0} = \min(1.0, \Delta k) = \min \left( 1.0, \frac{\sigma_{0}}{\sigma_{E}} \frac{V_{\text{max}} - V_{\text{min}}}{V_{\text{max}} - V_{\text{avg}}} \right)
\]

(3)

Alternatively, when the threshold energy \( E \) is set to its upper bound \( E = V_{\text{min}} + 1/k \), \( k_0 \) is set to:

\[
k_{0} = k_{0} \equiv \left( 1 - \frac{\sigma_{0}}{\sigma_{E}} \right) \times \frac{V_{\text{max}} - V_{\text{min}}}{V_{\text{avg}} - V_{\text{min}}} \cdot
\]

(4)

If \( k'_{0} \) is calculated between 0 and 1. Otherwise, \( k_{0} \) is calculated using Eq. (3).

2.2. Energetic reweighting of GaMD simulations

For energetic reweighting of GaMD simulations to calculate potential of mean force (PMF), the probability distribution along a reaction coordinate is written as \( p'(A) \). Given the boost potential \( \Delta V(\tilde{r}) \) of each frame, \( p'(A) \) can be reweighted to recover the canonical ensemble distribution \( p(A) \), as:

\[
p(A) = p'(A) \frac{e^{\beta \Delta V(\tilde{r})}}{\sum_{i=1}^{M} e^{\beta \Delta V(\tilde{r})}} , \quad j = 1, \ldots, M
\]

(5)

where \( M \) is the number of bins, \( \beta = k_{B}T \) and \( e^{\beta \Delta V(\tilde{r})} \) is the ensemble-averaged Boltzmann factor of \( \Delta V(\tilde{r}) \) for simulation frames found in the \( j \)th bin. The ensemble-averaged reweighting factor can be approximated.

### Table 1

Summary of GaMD simulations performed on the Numb RNA (apo), MSI1 protein without RNA Numb (apo) and with RNA Numb started from the Bound and Unbound states.

| System                  | \( N_{\text{max}} \) | Length(ns) | Boost potential (kcal/mol) |
|-------------------------|-----------------------|------------|---------------------------|
| Numb RNA (apo)          | 14,616                | 500 x 3    | 9.84 ± 3.02              |
| MSI1 protein (apo)      | 28,742                | 500 x 3    | 12.28 ± 3.54             |
| MSI1-Numb (Bound)       | 29,368                | 500 x 3    | 14.08 ± 3.79             |
| MSI1-Numb (Unbound)     | 29,569                | 1,200 x 19 | 11.49 ± 3.36             |
using cumulant expansion:

$$e^{\Delta \alpha_{(r)}} = \exp \left\{ -\sum_{k=1}^{\infty} \frac{\beta^k}{k!} C_k \right\}$$  \hspace{1cm} (6)

where the first two cumulants are given by:

$$C_1 = \Delta V,$$

$$C_2 = \Delta V^2 - \Delta V^2 = \sigma^2. \hspace{1cm} (7)$$

The boost potential obtained from GaMD simulations usually follows near-Gaussian distribution (Miao and McCammon, 2017). Cumulant expansion to the second order thus provides a good approximation for computing the reweighting factor (Miao et al., 2015; Miao et al., 2014). The reweighted free energy $$F(A) = -k_B T \ln p(A)$$ is calculated as:

$$F(A) = F^*(A) - \sum_{k=1}^{\infty} \frac{\beta^k}{k!} C_k + F_c,$$  \hspace{1cm} (8)

where $$F^*(A) = -k_B T \ln p'(A)$$ is the modified free energy obtained from GaMD simulation and $$F_c$$ is a constant.

2.3. System setup

Two models were prepared for simulations of MSI1-RNA interactions. One was obtained from the first model of NMR structure of the Numb RNA-bound MSI1 protein (PDB: 2RS2, denoted as the “Bound” state) (Ohyama et al., 2011). Another one was generated with the first model by moving the Numb RNA ~30 Å away from its binding site in MSI1 (denoted as the “Unbound” state). The models for only the Numb RNA (denoted as the “apo” state) were obtained from the Numb-MSI1 complex by deleting the corresponding binding partner. The four systems were solvated in explicit water using tleap in the AMBER 20 package (D.A. Case, D.S. Cerutti, T.E. Cheatham, III, T.A. Darden, R.E. Duke et al., 2020, AMBER, 2020, University of California, San Francisco). The system charges were then neutralized at 0.15M NaCl using tleap. The AMBER ff14SBionlyc force field parameters (Maier et al., 2015), RNA.LJbb (Zgarbová et al., 2011) and TIP3P model (Jorgensen et al., 1983) were applied for the protein, RNA and water molecules, respectively.

2.4. Simulation protocol

GaMD simulations were performed by using the GPU-accelerating program pmemd.cuda in AMBER 20 (D.A. Case, D.S. Cerutti, T.E. Cheatham, III, T.A. Darden, R.E. Duke et al., 2020, AMBER, 2020, University of California, San Francisco). The hydrogen-heavy atom bonds were constrained using the SHAKE algorithm and the simulation time step was set to 2.0 fs. The particle mesh Ewald (PME) method was employed to compute the long-range electrostatic interactions and a cutoff value of 9.0 Å was applied to treat the non-bonded atomic interactions. The temperature was controlled using the Langevin thermostat with a collision frequency of 1.0 ps$^{-1}$. Each system was minimized using steepest descent for 50,000 steps and conjugate gradient for another 50,000 steps. After minimization, the system was heated from 0 to 300 K in 1 ns simulation by applying 1 kcal/(mol·Å$^2$) harmonic position restraints to the protein and RNA heavy atoms with a constant number, volume and temperature (NVT) ensemble. Each system was further equilibrated using a constant number, pressure and temperature (NPT) ensemble at 1 atm and 300 K for 1ns with same restraints as in the NVT run. Another 1.2 ns cMD simulations were performed to collect potential energy statistics (including the maximum, minimum, average and standard deviation). Then 24 ns GaMD equilibration after applying the boost potential was performed. Previous studies showed that hundreds-of-nanosecond GaMD simulations with multiple replicas could provide good sampling of native biomolecular complexes (Miao and McCammon, 2016a,b; Bhattarai et al., 2020; Draper-Joyce et al., 2021; Wang and Miao, 2019) and microsecond GaMD simulations were able to capture ligand binding from solvent to target proteins (Miao and McCammon, 2016a,b; Pawnikar and Miao, 2020; Miao et al., 2018). Therefore, 3 independent 500 ns GaMD simulations with randomized initial atomic velocities were performed on the MSI1-Numb bound system (Table 1). While 19 independent 1,200 ns GaMD simulations were performed on the unbound system with the aim to capture at least 5 events of successful RNA binding to the MSI1 protein (Table 1). In addition, three independent 500 ns GaMD production runs with randomized initial atomic velocities were performed on the apo states of the MSI1 protein and RNA, respectively. Simulation frames were saved every 0.4 ps for analysis. The boost potential in GaMD simulation was calculated using potential energies of the present frame, as well as potential statistics including the minimum, maximum, average and standard deviation that are collected through short cMD and GaMD equilibration runs in the same simulation. The AMBER input files of GaMD equilibration and production simulations are provided in Supporting Information.

2.5. Simulation analysis

CPPTRAJ (Roe and Cheatham, 2013) and VMD (Humphrey et al., 1996) were used to analyze the GaMD simulations. Important reaction coordinates were identified from the simulation trajectories such that they involved dynamic regions (e.g., the [j-3] loop of MSI1) and could be used to differentiate conformational states of the MSI1-Numb system. Therefore, root-mean-square deviations (RMSDs) of the backbone of core RNA (central three nucleotides UAG in Numb) and the [j-3] loop of MSI1 relative to the first NMR conformation in the PDB with alignment of the MSI1 protein core (excluding the highly flexible C-terminus residues 96 to 103), the number of native contacts between MSI1 and Numb RNA ($N_{contacts}$), the radius of gyration ($R_g$) and end-to-end distance of the Numb RNA were selected as reaction coordinates. Root-mean-square fluctuations (RMSFs) were calculated for the protein residues and RNA nucleotides, averaged over the GaMD production simulations and color coded for schematic representation of each system. Since only the Sim1 to Sim6 GaMD trajectories successfully captured complete binding of the Numb RNA to MSI1, these trajectories were used separately for structural clustering to identify the RNA binding pathways using the hierarchical agglomerative algorithm in CPPTRAJ (Roe and Cheatham, 2013). The RMSD cutoff was set to 3.0 Å for the core RNA backbone to form a cluster. The PyReweighting (Miao et al., 2014) toolkit was applied to reweight GaMD simulations to recover the original free energy or PMF profiles of the two MSI1-Numb systems. Three 500 ns GaMD production simulations were combined for calculating the PMF profiles of the Bound MSI1-Numb system. Six of the 1,200 ns GaMD production simulations that successfully captured spontaneously binding of Numb RNA to MSI1 started from the Unbound state and were combined to calculate the corresponding PMF profiles. A bin size of 1.0 Å was used for the core RNA backbone RMSD, the MSI1 [j-2-3] loop backbone RMSD, the Numb $R_g$ and the end-to-end distance of Numb. A bin size of 100 was used for $N_{contacts}$. The cutoff was set to 500 frames for all 2D PMF calculations.

3. Results

3.1. GaMD simulations captured complete binding of the Numb RNA to the MSI1 protein

Extensive GaMD production simulations were performed on the MSI1-Numb system, including three independent 500 ns runs started from the Bound state and 19 independent 1,200 ns runs started from the Unbound state (Table 1). The GaMD simulations started from the Bound state recorded average and SD of the boost potential as 14.08 kcal/mol and 3.79 kcal/mol, respectively (Table 1). The GaMD simulations started from the Unbound state showed similar average and SD of boost potential.
with 11.49 kcal/mol and 3.36 kcal/mol, respectively (Table 1). The GaMD simulations of “apo” state of MSI protein showed similar average and SD of boost potential with 12.28 kcal/mol and 3.02 kcal/mol, respectively (Table 1). In contrast, the GaMD simulations of “apo” state of Numb RNA with fewer atoms recorded lower average and SD of boost potential with 9.84 kcal/mol and 3.02 kcal/mol, respectively (Table 1). The Bound MSI1-Numb complex was found to maintain the NMR structure with ~3 Å RMSD of the core RNA backbone during most of the GaMD simulations (Fig. 1A). In the GaMD simulations started from the Unbound state, the core RNA backbone RMSD relative to the NMR structure in 6 of 19 simulations (~31.5%) decreased to ~3.0 Å, suggesting that complete binding of the Numb RNA from free diffusion in the solvent to the MSI1 target site was successfully captured (Fig. 1B & S1). Spontaneous binding of RNA was observed in the Sim1 after ~100 ns with the RNA backbone RMSD decreased to ~3.0 Å relative to the first NMR structure (Fig. 1B). In Sim2, the Numb RNA bound to MSI1 during ~1,010–1,130 ns and then dissociated to the solvent (Fig. 1B). The Numb RNA bound to MSI1 after ~800 ns in Sim3, Sim4 and Sim5 (Fig. 1B). In Sim6, spontaneous binding of RNA was observed after ~1000 ns (Fig. 1B). Multiple RNA binding events captured in the present GaMD simulations allowed us to characterize the dynamic interactions between the MSI1 protein and Numb RNA.

3.2. Variations of structural flexibility upon MSI1-Numb RNA binding

We analyzed structural flexibility of both the MSI1 protein and Numb RNA in the GaMD simulations. During GaMD simulations started from the Bound NMR structure, the MSI1 protein underwent small fluctuations except the loop connecting β2 and β3 strands (the β2-β3 loop) and the C-terminus (Fig. S2A). The fifth nucleotide in the Numb RNA exhibited significantly higher flexibility than the other nucleotides, especially the central three ones UAG (denoted as the core RNA). Similar findings were observed in the simulations of the apo Numb RNA (Fig. 2D). This suggested that interactions between the core RNA and the MSI1 were strong. Thus, the core RNA might play an important role in the interactions between the MSI1 protein and Numb RNA. Furthermore, the MSI1 β2-β3 loop and C-terminus and the Numb RNA exhibited significantly higher fluctuations in the GaMD simulations started from the Unbound state than those in simulations started from the Bound state (Fig. S2B). Note that both Bound and Unbound conformations of MSI1-Numb were observed in the GaMD simulations (Sim1-Sim6) started from the Unbound state (Fig. 1B). In this regard, trajectories of Sim7 to Sim19 started from the Unbound state that did not capture the Numb RNA binding were used for RMSF calculation of the system in the Unbound state (Fig. 1B).

The three GaMD trajectories started from the Bound state (Fig. 1A) plus the 100–1200 ns trajectory of Sim1, 850–1200 ns trajectory of Sim3, 800–1200 trajectory of Sim5 and 1000–1200 ns trajectory of Sim6 started from the Unbound state (Fig. 1B) were used for RMSF calculation of the system in the Bound state (Fig. 2A). Results showed that the flexibility of loop β2-β3 and C-terminus of MSI1 in the Bound state was significantly lower than in the Unbound and apo state (Fig. 2), being similar to the findings observed in Fig. S2. These motifs were suggested to be important for binding of the Numb RNA (Ohyama et al., 2011) and small molecules (Clingman et al., 2014) to the MSI1 protein.

3.3. Free energy profiles of RNA binding to the MSI1 protein

Free energy profiles were calculated from the GaMD simulations using the core RNA backbone RMSD relative to the NMR structure and the number of native contacts between MSI1 and Numb RNA (N_contacts) as reaction coordinates. Only one low-energy minimum of the “Bound” conformation was identified from the GaMD simulations on the NMR structure, in which the core RNA backbone RMSD and N_contacts centered around (1.2 Å, 1600) (Fig. 3A). Five low-energy minima were identified from GaMD simulations started from the Unbound state, including the “Bound”, “Intermediate I1”, “Intermediate I2”, “Intermediate I3” and “Unbound” states, in which the core RNA backbone RMSD and N_contacts centered around (2.0 Å, 1500), (5.2 Å, 480), (9.5 Å, 200), (25.0 Å, 10) and (40 Å, 0), respectively (Fig. 3B). The intermediate I1, I2 and I3 conformational states are shown in Fig. 4. The Numb RNA binding to MSI1 involved large conformational changes in both the RNA and protein (Figs. 4 and 5B, Movies S1 and S2). In the Unbound state, RNA diffused far away from the protein with ~40 Å RMSD in the core RNA backbone relative to the NMR complex structure, while the protein loop β2-β3 could adopt a conformation close to the NMR structure with ~1.0 Å RMSD (Fig. 5B). As the RNA moved towards the MSI protein in the I2 and I3 intermediate states, the Numb RNA exhibited RMSDs of 10.8 Å and 22.3 Å from the target binding conformation and interacted with the protein loop β2-β3 and C-terminus, respectively (Fig. 4). Such interactions induced significant conformational changes of the corresponding regions, which appeared to pull the Numb RNA to the protein target site (Movies S1 and S2). Then, the Numb RNA moved closer to the target site in the I1 intermediate state with reduced RMSD of ~5.2 Å. Meanwhile, the protein loop β2-β3 and C-terminus in the I1 intermediate state showed RMSDs of ~4.0 Å and ~21.2 Å, respectively, relative to the NMR structure (Figs. 4A & 5B). Finally, the protein (especially the β2-β3 loop and C-terminus) and RNA rearranged their conformations and formed complex in the “Bound” state that was similar to the NMR structure.
experimental structure (Fig. 4D). Remarkably, positively charged residues (Arg61 and Arg99) in the β2-β3 loop and C-terminal region of the MSI1 protein formed favorable salt-bridge and hydrogen bond interactions with the central nucleotide A106 of Numb RNA. In the intermediate I1 state, the Numb RNA formed interactions with both the β2-β3 loop and C terminus of MSI1, leading to large conformational changes of these two regions (Fig. 4A). Notably, the sidechain of residue Arg99 in the C terminus of the MSI1 protein formed three hydrogen bonds with the sidechain of nucleotide A106 in the Numb RNA (Fig. 4A). In the intermediate I2 state, the Numb RNA formed interactions with the β2-β3 loop of MSI1, leading to a large conformational change of this loop (Fig. 4B). The sidechain of residue Arg61 in MSI1 could flip out to the solvent, forming salt-bridge with the backbone (oxygen atom in the phosphate group) of the nucleotide A106 of Numb RNA (Fig. 4B). In the intermediate I3 state, Arg99 in the C terminus of MSI1 formed a hydrogen bond and a salt-bridge with sidechain and backbone of the nucleotide A106 in the Numb RNA (Fig. 4A).
Fig. 5. 2D PMF profiles of the MSI1 β2-β3 loop backbone RMSD and core RNA backbone RMSD relative to the first NMR conformation (PDB: 2RS2) are calculated from GaMD simulations started from the (A) Bound and (B) Unbound states of the MSI1-Numb system.

Numb RNA, respectively, for which a large conformational change of the protein C terminus was observed (Fig. 4C). No contact between MSI1 Arg99 and Numb A106 was found in the NMR experimental structure (Fig. 4C). The distance between MSI1 Arg99 and Numb A106 and the Numb Core RMSD were chosen as reaction coordinates to calculate 2D free energy profiles (Fig. S3). Indeed, a salt-bridge interaction between MSI1 Arg99 and Numb A106 was identified in low-energy intermediate states (I1 and I2) during RNA binding to MSI1 (Fig. S3B). Thus, the electrostatic interaction between MSI1 Arg99 and Numb A106 played a significant role in the recognition and binding of the Numb RNA to MSI1 protein.

As described above, binding of the Numb RNA induced higher flexibility of the MSI1 β2-β3 loop (Fig. 2B) and large conformational change of the same region was observed in the intermediate I1 and I2 states (Fig. 4A and B). Therefore, the MSI1 β2-β3 loop backbone RMSD and core Numb RNA backbone RMSD relative to the experimental structure were used as reaction coordinates to further compute 2D free energy profiles (Fig. 5). The MSI1 β2-β3 loop was highly flexible, sampling a large conformational space with the backbone RMSD ranging from ~0 Å to ~8.0 Å (Fig. 5B&S4). This loop sampled two distinct low-energy conformations, including the “Closed” (bound) (RMSD < 1 Å) and “Open” (free) states (RMSD ~3–5 Å) (Fig. 5B). Compared to the “Open” state, the MSI1 β2-β3 loop moved closer to the core domain in the “Closed” state (Fig. 4A). Five low-energy states were identified from GaMD simulations starting with the Unbound state, including the “Unbound/Close”, “Intermediate 13/Close”, “Intermediate 11/Close” and “Bound/Close” states, in which the MSI1 β2-β3 loop backbone RMSD and core RNA backbone RMSD were located around (1.0 Å, 40 Å), (1.2 Å, 25.0 Å), (5.0 Å, 11.5 Å), (4.2 Å, 5.5 Å) and (1.5 Å, 2.0 Å), respectively (Fig. 5B). The Numb RNA and MSI1 β2-β3 loop accommodated to each other to form the final bound conformation (Fig. 5B).

3.4. Pathways of RNA binding to the MSI1 protein

Here, we focused on exploring the pathway and mechanism of RNA binding, for which only six out of 19 GaMD simulations of the unbound system successfully captured the RNA binding process. Therefore, all these six trajectories were analyzed in detail to determine the RNA binding pathways. Thus, structural clustering was performed separately on the GaMD trajectories of Sim1-Sim6 to identify the representative binding pathways of the Numb RNA to the MSI1 protein. Movies S1 and S2 show GaMD trajectories of Sim1 and Sim2 started from the Unbound state in Supporting Information, respectively. The structural clusters were reweighted to obtain their original free energy values, which ranged from 0.0 kcal/mol to ~4.5 kcal/mol. The top reweighted clusters with PMF £2.0 kcal/mol were selected to represent the pathways of the Numb RNA binding to MSI1 (Fig. 6). In Sim1, Sim3, Sim5 and Sim6, the Numb RNA bound to MSI1 via interactions with the protein C terminus (Fig. 6A, C, 6E and 6F). In Sim4, the Numb RNA bound to MSI1 via interactions with the β2-β3 loop of MSI1 (Fig. 6D). In Sim2, both the β2-β3 loop and C terminus of MSI1 contributed important interactions with the Numb RNA during its binding to the protein target site (Fig. 6B). These findings revealed the important roles of the β2-β3 loop and C terminus of MSI1 in binding of the Numb RNA, especially for their large conformational changes in the three intermediate states of RNA binding (Fig. 4). It is worth to noting that no strong interactions were formed between the Numb RNA and the MSI1 β2-β3 loop and C terminus in the final “Bound” state. Rather these two regions formed important interactions with RNA in the intermediate states during the RNA binding process. This was consistent with the above RMSF analysis that higher flexibilities were observed in these two dynamic regions of MSI1 (Fig. 2).

3.5. The Numb RNA bound to the MSI1 protein via an induced fit mechanism

In order to further explore the mechanism of RNA binding to the MSI1 protein, we computed free energy profiles to characterize conformational changes of the Numb RNA upon binding to MSI1. In this regard, the radius of gyration ($R_g$) and the end-to-end distance of Numb were calculated to monitor its possible conformational changes. We used the $R_g$ and end-to-end distance of the Numb RNA and the core RNA backbone RMSDs as reaction coordinates to calculate 2D PMF profiles (Fig. 7). Notably, the Numb RNA sampled a large conformational space during binding to the MSI1 protein in the GaMD simulations started from the Unbound state (Fig. 7B). From the reweighted 2D PMF profiles, we identified a similar “Bound” low-energy well in simulations started from both the Bound and Unbound states, for which the Numb RNA $R_g$ and core RNA backbone RMSD centered around (8.5 Å, 2.0 Å) and (8.5 Å, 2.5 Å), respectively (Fig. 7A-B). Another four low-energy states were identified in GaMD simulations started from Unbound conformation, including the “Unbound”, “Intermediate 11”, “Intermediate 12” and “Intermediate 13”, for which the core RNA backbone RMSD and $R_g$ of Numb centered around (40.0 Å, 6.2 Å), (5.0 Å, 7.2 Å), (6.9 Å, 6.2 Å) and (25.0 Å, 7.5 Å), respectively (Fig. 7B).

Furthermore, we calculated 2D PMF profiles regarding the core RNA backbone RMSD and the end-to-end distance of Numb RNA (Fig. 7C-D). Only one low-energy state (“Bound”) was identified in the 2D PMF profile calculated from the GaMD simulations of the bound NMR structure, in which the Numb RNA adopted primarily the “Extended” conformation (Fig. 7C). The Numb end-to-end distance and core RNA backbone RMSD centered around (22.5 Å, 1.8 Å) (Fig. 7C). In contrast, six distinct low-energy states were identified from the 2D PMF profile in
Fig. 6. Binding pathways of the Numb RNA to the MSI1 protein revealed from GaMD simulations: Starting from free diffusion in the solvent, the Numb RNA spontaneously bound to the target site of the MSI1 via intermediate conformations in the (A) “Sim1”, (B) “Sim2”, (C) “Sim3”, (D) “Sim4”, (E) “Sim5” and (F) “Sim6” GaMD trajectories. In the intermediate conformations, the Numb RNA interacted with the C-terminus (A, C, E and F), the $\beta_2-\beta_3$ loop (D) or both the C-terminus and $\beta_2-\beta_3$ loop (B) of MSI1. The MSI1 protein is shown in blue ribbons. The Numb RNA structural clusters (sticks) are colored by the reweighted PMF free energy values in a green (0.0 kcal/mol)-white-red (2.0 kcal/mol) color scale.
the GaMD simulations started from the Unbound conformation, including the “Unbound”, “Intermediate I1”, “Intermediate I2”, “Intermediate I3” and “Bound”, in which the Numb RNA adopted primarily the “Curled”, “Curled”, “Curled”, and “Extended” conformations, respectively (Figs. 7D and 4). The end-to-end distance of Numb RNA and core RNA backbone RMSD centered around (15.0 Å, 40.0 Å) in the “Unbound/Curled” state, (15.2 Å, 25.0 Å) in the “Intermediate I3/Curled” state, (~15–10 Å, 10–17 Å) in the “Intermediate I2/Curled” state, (12.5 Å, 5.2 Å) in the “Intermediate I1/Curled” state and finally (22.0 Å, 2.5 Å) in the “Bound/Extended” state (Fig. 7D).

In comparison, the Numb RNA sampled a larger conformational space with a wider range of Rg or end-to-end distance in the “Bound” state than in the “Unbound” and even the three Intermediate conformations (Fig. 7B and D), suggesting binding of the Numb RNA to the MSI1 protein involved largely induced fit. To further support the “induced-fit” mechanism of RNA binding to MSI1, we have added 500 ns x 3 GaMD simulations on the apo form of both MSI1 protein and Numb RNA (Table 1). In the RNA-bound MSI1, the Numb RNA sampled the “Bound” low-energy state with Rg and end-to-end distance centered around (8.5 Å, 22.0 Å) (Fig. S5A). In contrast, the apo Numb RNA could not sampled the “Bound” state (Fig. S5F). In this context, Numb RNA in GaMD simulations of the RNA unbound system sampled a large conformational space covering that of both the bound and apo systems (Fig. S5E). Similar findings were observed for the MSI1 protein. The MSI1 β2-β3 loop backbone RMSD and C-terminus backbone RMSD centered around (3.0 Å, 20 Å) (Fig. S5A). It is worth noting that the large RMSD of MSI1 C-terminus observed in the “Bound” state (~20 Å) was reasonable because similar values were obtained among the 20 conformations in the NMR experimental structure (PDB: 2RS2). MSI1 in GaMD simulations of the RNA unbound system sampled a large conformational space covering that of both the bound and apo systems (Fig. S5B). In comparison, the apo MSI1 could rarely sample these conformations (Fig. S5C). Together, these results suggested that the RNA binding to MSI1 protein adopted an “induced-fit” mechanism.

4. Discussion

In this study, we have applied extensive all-atom GaMD simulations with a total length of 24,300 ns to investigate dynamic interactions between the Numb RNA and MSI1 protein. The GaMD simulations unprecedentedly captured multiple times of spontaneous and highly accurate binding of the Numb RNA from bulk solvent to the MSI1 protein with <2 Å RMSD in the core RNA backbone compared with the experimental structure. Proper energetic reweighting of the GaMD simulations allowed us to calculate free energy profiles to characterize the MSI1-Numb binding process.

Relatively low-energy conformational states of RNA binding to MSI1 protein were identified from the GaMD simulations, including the Unbound, Intermediate I1, Intermediate I2, Intermediate I3, and Bound states (Fig. 4). In the intermediate states, the β2-β3 loop and C terminus of the MSI1 protein were found to be essential for recognition and binding.
of the Numb RNA (Fig. 4A–C & 6). The charged residues Arg61 and Arg99 of MS1I formed critical hydrogen bond and salt-bridge interactions with the Numb RNA during the GaMD simulations, particularly in the Intermediate and Bound states. The important role of Arg99 was consistent with the previous finding that mutation of the corresponding residue in MS12 (Arg100Ala) decreased the binding affinity of Numb RNA (Minuesa et al., 2019). The salt bridge between Arg61 (MS1I protein) and A106 (Numb RNA) was also observed in the bound NMR structure (Fig. 4B). Furthermore, Arg61 was characterized as a key residue for inhibitor binding as the MS1I Arg61Glu mutant exhibited ~5 fold decrease in the inhibitor binding affinity (Clingman et al., 2014). Additionally, strong binding between the RNA core and MS1I protein was observed in the GaMD simulations started from the Bound state. This agreed well with the previous finding obtained by Zearfoss et al. (Zearfoss et al., 2014) that the central UAG RNA nucleotides form the MS1 recognition element and make major contributions to the binding affinity. Therefore, our GaMD simulations revealed that long-range electrostatic interactions played an important role in the Numb RNA binding to the MS1I protein and identified two critical protein residues (Arg61 and Arg99) for RNA recognition, being highly consistent with previous experimental findings.

Conformation selection (Tsai et al., 1999; Tsai et al., 1999; Kumar et al., 2000) and induce fit (Okazaki and Takada, 2008; Williamson, 2000) are two common models for describing biomolecular recognition. In this context, our GaMD simulations have revealed that binding of the Numb RNA to the MS1I protein involved predominantly an induced fit mechanism, in which both the RNA and protein underwent significant conformational changes during binding (Figs. S4 and S5). This is consistent with previous studies of other protein-RNA interactions (Williamson, 2000; Leulliot and Varani, 2001; Suryadi et al., 2005, Piticì et al., 2002), including ribosomal protein S15-rRNA and U1A-RNA complexes. A major conformational change of the rRNA was found upon binding to the S15 protein through comparison of the free and bound structures of S15 and rRNA, suggesting induced fit of the protein and RNA (Williamson, 2000). MD simulations combined with available structure analysis also indicated that binding of the U1A protein and RNA followed an induced fit mechanism (Piticì et al., 2002). For the U1A protein, MD simulations indicated that induced fit upon binding involved a non-native thermodynamic substate, in which the structure is preorganized for binding. In contrast, induced fit of the RNA involved a distortion of the native structure to an unstable form in solution.

It is important to note that the presented GaMD simulations captured six binding events of Numb RNA to MS1I protein. In addition to free energy profiles calculated by combining all six successfully RNA binding simulations (Fig. 3B), we calculated free energy profiles from each of the six successful GaMD simulations of RNA binding to as shown in Fig. S6. While the Unbound, Bound and Intermediate low-energy states could be sampled in the individual simulations, these free energy profiles showed differences in terms of the positions and values of the free energy minima and barrier heights. Results suggested that our current GaMD simulations were still not converged. Nevertheless, Numb RNA could bind to the MS1I protein in two main pathways via the protein β2-β3 loop and C-terminus interaction sites, which were observed for 4 and 2 times during the six successful GaMD simulations, respectively. For future studies, more events of RNA binding and unbinding would still need to be simulated in order to calculate the RNA binding free energies and kinetic rates quantitatively. Nevertheless, the focus of this study was to uncover the dynamic pathways and mechanism of RNA binding to the MS1I protein. The binding of RNA to MS1I protein via interacting with the two main binding sites (β2-β3 loop and C-terminus) were observed multiple times during GaMD simulations. Therefore, the six GaMD trajectories should reveal accurate Numb RNA binding pathways. More binding and unbinding events would still need to be simulated in order to calculate the RNA binding free energies and kinetic rates quantitatively. In this regard, our recently developed selective GaMD algorithms (Miao et al., 2020; Wang and Miao, 2020) could be useful to address the challenge. In particular, the peptide GaMD (Pep-GaMD) method (Wang and Miao, 2020), which works by selectively boosting the essential peptide potential energy, has been demonstrated to capture repetitive binding and unbinding of highly flexible peptides to the target protein within microsecond simulations (Wang and Miao, 2020). Apart from enhanced conformational sampling, accurate force fields are also needed especially for the RNA (Sponer et al., 2018; Tan et al., 2018; Kührova et al., 2019; Cesari et al., 2019) in order to simulate repetitive RNA dissociation and binding to RBPs. Even the force field works well individually for the protein and RNA, combination of protein and RNA force fields in the MD simulations could introduce additional challenges (Sponer et al., 2017; Krepl et al., 2015; Sponer et al., 2018). Nevertheless, we have observed multiple complete binding events of the Numb RNA to the MS1I protein with our current force field settings and GaMD simulations, which shall guide future studies of RNA-protein interactions.

5. Conclusions

In summary, all-atom GaMD simulations with unconstrained enhanced sampling and free energy calculations have provided important insights into the mechanism of the Numb RNA binding to the MS1I protein. The results and the methods used in this study would help in simulating binding process of RNA to the RBPs in general, to accurately predict the binding mechanism of protein–RNA interactions. For future studies, the effects of small molecule binding in the MS1I-Numb interactions still need to be determined and our simulation findings await validation in the wet-lab experiments. Further studies are planned to simulate both dissociation and binding of RNA to the RBPs and accurately predict the thermodynamics and kinetics of protein–RNA interactions. These efforts are expected to greatly facilitate rational drug design targeting the MS1I and other RBPs.

CRediT authorship contribution statement

Jinlan Wang: performed the research, analyzed the data, wrote the paper. Lan Lan: analyzed the data. Xiaoqing Wu: analyzed the data. Yinglong Miao: designed research, wrote the paper.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix B. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.crstbi.2021.12.002.
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