

Chapter 9 1

Multiscale Modeling of RNA Structure 2

and Dynamics 3

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Abstract We have developed a multiscale approach for RNA folding using discrete molecular dynamics (DMD), a rapid conformational sampling algorithm. We use a coarse-grained representation to effectively model RNA structures. Benchmark studies suggest that the DMD-based RNA model is able to accurately fold small RNA molecules (<50 nucleotides). However, the large conformational space and force field inaccuracies make it difficult to computationally identify the native states of large RNA molecules. We devised an automated modeling approach for prediction of large and complex RNA structures using experimentally derived structural constraints and tested it on several RNA molecules with known experimental structures. In all cases, we were able to bias the DMD simulations to the native states of these RNA molecules. Therefore, a combination of experimental and computational approaches has the potential to yield native-like models for the diverse universe of functionally important RNAs, whose structures cannot be characterized by conventional structural methods. 5
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9.1 Introduction 19

RNA molecules play a wide range of functional roles in gene expression, from regulating transcription and translation [e.g., riboswitch regulator motifs (Edwards et al. 2007)] to decoding genetic messages (tRNA), catalyzing mRNA splicing [spliceosome RNA or self-splicing introns (Vicens and Cech 2006)] and protein synthesis (rRNA). Knowledge of the underlying RNA structure in these and many other molecules is a fundamental prerequisite to a complete understanding of RNA function. Methods such as X-ray crystallography and NMR spectroscopy offer critical 20
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27 insight into the details of RNA structure–function relationships. However, many
28 RNAs contain both structured and functionally important but flexible elements.
29 These RNAs are not amenable to structure determination in their intact forms by
30 crystallography or NMR. Hence, molecular modeling of RNA to predict three-
31 dimensional structure and dynamics is crucial for our understanding of RNA functions.

32 Currently, RNA folding tools focus mainly on predicting RNA secondary
33 structure (Hofacker 2003; Mathews 2006; Zuker 2003). Using a dynamic program-
34 ming approach (Eddy 2004), secondary structures are inferred by scoring nearest-
35 neighbor stacking interactions with adjacent base pairs (Mathews 2006). These
36 RNA secondary structure prediction methods play an important role in the current
37 study of RNA. However, in order to model the tertiary structure of RNA molecules,
38 it is necessary to explicitly model RNA in 3D. Cao and Chen designed a simplified
39 diamond-lattice model for predicting folded structure and thermodynamics of RNA
40 pseudoknots (Cao and Chen 2005, 2006). This approach quantitatively predicts the
41 free energy landscape for sequence-dependent folding of RNA pseudoknots, in
42 agreement with experimental observations (Cao and Chen 2005, 2006). However,
43 due to lattice constraints and the dynamic issues associated with predefined Monte
44 Carlo moves (Baumgartner 1987), off-lattice models are necessary to accurately
45 model RNA 3D structure.

46 Computational tools for manually constructing RNA models have been devel-
47 oped for RNA 3D structure prediction (Shapiro et al. 2007). These methods use
48 comparative sequence analysis to manually construct 3D models, with or without
49 reference to a known, homologous 3D structure. Their accuracy is enhanced by use
50 of experimental probes of secondary or tertiary structure and libraries of modular
51 3D motifs (Jossinet and Westhof 2005; Major et al. 1991, 1993; Massire et al. 1998;
52 Massire and Westhof 1998; Shapiro et al. 2007; Tsai et al. 2003). Recently,
53 significant progress has been made toward ab initio modeling of RNA 3D structures
54 (Das and Baker 2007; Ding et al. 2008; Parisien and Major 2008). These studies
55 show that starting only with sequence, it is possible to predict the structures of some
56 small RNA motifs with atomic-level accuracy. However, as RNA length increases,
57 the conformational space increases exponentially and the inherent inaccuracies of
58 the force field accumulate, limiting the ability of current methods to predict the
59 structures of large RNAs automatically. De novo prediction of large RNA
60 structures with nontrivial tertiary folds from sequence alone remains beyond the
61 realm of current ab initio algorithms.

62 We have developed a multiscale approach (Ding and Dokholyan 2005) for RNA
63 modeling based on a coarse-grained RNA model for discrete molecular dynamics
64 (DMD) simulations (Ding et al. 2008). DMD is a special type of molecular dynamics
65 simulation in which pairwise interactions are approximated by stepwise functions.
66 This approximation enables DMD to sample conformational space more efficiently
67 than traditional molecular dynamics simulations (Dokholyan et al. 1998). Using the
68 coarse-grained RNA model with DMD simulations, we were able to accurately fold a
69 set of 150 small RNA molecules (<50 nt) within 6 Å (a majority within 4 Å) to their
70 native states (Ding et al. 2008). To solve the folding problem of large RNA molecules
71 with complex tertiary 3D structures, we proposed to incorporate experimentally

derived structural information into our structure determination protocol. Long-range 72
constraints for RNA modeling can be inferred from a variety of biochemical and 73
bioinformatic techniques, ranging from chemical footprinting and cross linking to 74
sequence covariation (Gutell et al. 1992; Juzumiene et al. 2001; Michel and Westhof 75
1990; Ziehler and Engelke 2001). Experimental constraints derived from these bio- 76
chemical and bioinformatics techniques are generally of lower than atomic resolution, 77
but can be readily incorporated into the coarse-grained RNA model for structure 78
determination. The all-atom RNA model can then be reconstructed from the coarse- 79
grained structural model. 80

First, we will describe our coarse-grained representation of RNA models for 81
DMD simulations. Then, we will describe and evaluate the applications of the 82
DMD–RNA procedure to ab initio folding of a set of small RNA models and 83
structure determination using experimental constraints. 84

9.2 Coarse-Grained RNA Modeling Using Discrete Molecule 85 Dynamics 86

We use DMD as the conformational sampling engine. A detailed description of the 87
DMD algorithm can be found elsewhere (Dokholyan et al. 1998; Rapaport 2004; 88
Zhou and Karplus 1997). The difference between discrete molecular dynamics and 89
traditional molecular dynamics is in the interaction potential functions. Interatomic 90
interactions in DMD are governed by stepwise potential functions (Fig. 9.1a). 91
Neighboring interactions (such as bonds, bond angles, and dihedrals) are modeled 92
by infinitely high square well potentials (Fig. 9.1b). By approximating the continu- 93
ous potential functions with step functions of pairwise distances, DMD simulations 94
are reduced to event-driven (collision) molecular dynamics simulation. In a DMD 95
simulation, atoms move with constant velocity until they collide with another atom. 96
As soon as the potential of interaction between the two atoms changes (i.e., the 97
pairwise distance is at the step of the stepwise potential function), the velocities of 98
the two interacting atoms change instantaneously (Fig. 9.1a). These velocity 99
changes are required to conform to the conservation laws of energy, momentum, 100
and angular momentum. Each such collision is termed an “event.” The sampling 101
efficiency of DMD over traditional MD is mainly due to rapid processing of 102
collision events and localized updates of collisions (only colliding atoms are 103
updated at each collision). In the limit of infinitesimally small steps, the discrete 104
step function approaches the continuous potential function, and DMD simulations 105
become equivalent to traditional molecular dynamics. 106

We approximate the single-stranded RNA molecule as a coarse-grained “beads- 107
on-a-string” polymer with three beads representing each nucleotide, one for sugar (S), 108
one for phosphate (P), and one for nucleotide base (B) (Fig. 9.2a). The P and S beads 109
are positioned at the centers of mass of the corresponding phosphate group and the 110
5-atom ring sugar, respectively. For both purines (adenine and guanine) and 111

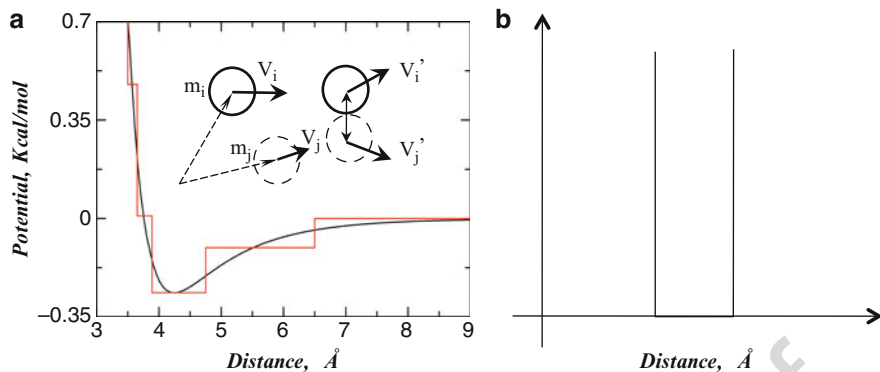


Fig. 9.1 Discrete molecular dynamics simulations. (a) Schematic of the DMD potential. The stepwise function used in DMD is the approximation of the continuous function in traditional molecular dynamics. The insert depicts the collision of two atoms with masses of m_i and m_j at the initial position of r_i and r_j , respectively. The two atoms move with constant velocities (v) until they meet at distance of R_{ij} . (b) Schematic of the potential energy of bonds in DMD. The atom pairs remain within the distance range during the simulation

112 pyrimidines (uracil and cytosine), we represent the base bead (B) as the center of the
 113 6-atom ring. The neighboring beads along the sequence, which may represent moieties
 114 that belong to the same or a neighboring nucleotide, are constrained to mimic the chain
 115 connectivity and local chain geometry (Fig. 9.2a). Types of constraints include
 116 covalent bonds (solid lines), bond angles (dashed lines), and dihedral angles
 117 (dotted–dashed lines). The parameters for bonded interactions mimic the folded
 118 RNA structure and are derived from a high-resolution RNA structure database
 119 (Murray et al. 2003) (Table 9.1). Nonbonded interactions are crucial to model the
 120 folding dynamics of RNA molecules. In our model, we include base-pairing
 121 (Watson–Crick pairs of A–U and G–C and Wobble pair of U–G), base-stacking,
 122 short-range phosphate–phosphate repulsion, and hydrophobic interactions, which
 123 are described in the following section with the parameterization procedure.

124 *Base Pairing.* Two base-paired nucleotides have bases facing each other with the
 125 corresponding sugar and base beads aligned linearly. We use the “reaction” algorithm
 126 to model the orientation dependence of base-pairing interactions. The details of the
 127 algorithm can be found in (Ding et al. 2003). Briefly, to model the orientation
 128 dependence, we introduce auxiliary interactions in addition to the distance-dependent
 129 interactions between hydrogen bond donor and acceptor atoms (Fig. 9.2b). For
 130 example, when the two nucleotides (e.g., A–U, G–C, or U–G, represented as B_i and
 131 B_j in Fig. 9.2b) approach the interaction range, we evaluate the distances between $S_i B_j$
 132 and $S_j B_i$, which define the relative orientations of these two nucleotides. A hydrogen
 133 bond is allowed to form only when the distances fall within predetermined ranges. A
 134 schematic of the auxiliary interaction potential is shown in Fig. 9.2c, and the
 135 corresponding interaction parameters are listed in Table 9.2.

136 *Hydrophobic Interactions and Overpacking.* Buried inside the double-helix, the
 137 planar surface of bases are hydrophobic in nature. We include a weak attraction



Fig. 9.2 Coarse-grained structural model of RNA employed in DMD simulations. (a) Three consecutive nucleotides, indexed $i-1$, i , $i+1$ are shown. The S , P , and B symbols correspond to loci of sugar, phosphate, and base beads in the RNA, respectively. Covalent interactions are shown as *thick lines*, angular constraints as *dashed lines*, and dihedral constraints as *dashed-dotted lines*. Additional steric constraints are used to model base stacking. (b) Hydrogen bonding in RNA base pairing. The base-pairing contacts between bases $B_{i-1}:B_{j+1}$ and $B_i:B_j$ are shown in *dashed lines*. A reaction algorithm is used (see Methods) for modeling the hydrogen-bonding interaction between specific nucleotide base pairs. (c) Schematic of the potential function for the auxiliary base-pairing interactions. (d) Histogram of the number of neighboring bases within a cutoff of 6.5 \AA

between all the base beads. Due to the coarse-graining feature of our model, the assignment of attraction between bases results in overpacking (e.g., the symmetrically attractive interactions tend to form close packing). In order to avoid the artifact of overpacking, we first evaluate the packing observed in experimental 3D structures (<http://ndbserver.rutgers.edu>). We compute for each base the number of neighboring bases within a cutoff distance of 6.5 \AA . The histogram of the number of neighbors is shown in Fig. 9.2d. Indeed, we find that the average number of neighbors is much smaller than that of close packing, 12. In order to avoid unrealistic close-packing due to the coarse-graining process, we introduce an effective energy term to penalize overpacking of bases:

$$E_{\text{overpack}} = dE\Theta(n_c - n_{\text{max}}), \tag{9.1}$$

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148 where $\Theta(x)$ is a step function,

$$\Theta(x) = \begin{cases} x & x > 0 \\ 0 & x \leq 0 \end{cases}, \quad (9.2)$$

149 n_c is number of contacts, and n_{\max} is the maximum number of contacts; dE is the
150 repulsion coefficient. Based on the histogram of the number of base neighbors
151 (Fig. 9.2d), we assign the value 4.2 for n_{\max} and 0.6 kcal/mol for dE .

152 *Base Stacking.* To model stacking interactions, we assume that each base bead
153 makes no more than two base–base stacking interactions and that three consecutively
154 stacked base beads align approximately linearly. To determine the stacking interaction
155 range between base beads, we compute center-to-center distances between base beads
156 from known RNA structures. We find that distribution depends on base type (purine or
157 pyrimidine) and identify stacking cutoff distances as 4.65 Å between purines, 4.60
158 between pyrimidines, and 3.80 Å between purine and pyrimidine. To approximately
159 model the linearity of stacking interactions, two bases that form a stacking interaction
160 to the same base are penalized for approaching closer than 6.5 Å. As a result, these
161 three bases effectively define an obtuse angle. Next, we discuss the energy parameter-
162 ization of base-stacking, base-pairing, and hydrophobic interactions.

163 *Parameterization of Base-Pairing, Base-Stacking, and Hydrophobic Interactions.*
164 In order to determine the pairwise interaction parameters for stacking and hydropho-
165 bic interactions for all pairs of a base, we decomposed the sequence-dependent free
166 energy parameters of the individual nearest-neighbor hydrogen bond model (INN-
167 HB) (Mathews et al. 1999). We assume that the interaction of neighboring base pairs
168 in INN-HB is the sum of all hydrogen-bond, base-stacking, and hydrophobic
169 interactions. In a nearest neighboring base-pair configuration (Fig. 9.1), B_{i+1} and B_i
170 (B_{j-1} and B_j) on one strand usually stack on top of each other. However, if both bases
171 B_{i+1} and B_j are purines, we found that they tend to stack instead. The distance
172 between bases B_i and B_{j-1} is usually greater than the cutoff distance of 6.5 Å for
173 hydrophobic interactions. Therefore, we used the following equations to estimate the
174 strength of pairwise interactions, where the first equation applies when B_{i+1} , B_j are
175 both purines and the second equation applies otherwise:

$$E\left(\begin{matrix} 5'B_iB_{i+1}3' \\ 3'B_jB_{j-1}5' \end{matrix}\right) = \left(E_{B_iB_j}^{\text{HB}} + E_{B_{i+1}B_{j-1}}^{\text{HB}}\right) + E_{B_iB_{i+1}}^{\text{Stack}} + E_{B_iB_{i+1}}^{\text{hydrophobic}} + E_{B_jB_{j-1}}^{\text{hydrophobic}}, \quad (9.3)$$

$$E\left(\begin{matrix} 5'B_iB_{i+1}3' \\ 3'B_jB_{j-1}5' \end{matrix}\right) = \left(E_{B_iB_j}^{\text{HB}} + E_{B_{i+1}B_{j-1}}^{\text{HB}}\right) + E_{B_iB_{i+1}}^{\text{Stack}} + E_{B_jB_{j-1}}^{\text{Stack}} + E_{B_{i+1}B_j}^{\text{hydrophobic}}. \quad (9.4)$$

176 Here, E^{stack} , E^{HB} , and $E^{\text{hydrophobic}}$ are the interaction strengths of base-stacking,
177 base-pairing, and hydrophobic interactions, respectively. Given the experimentally
178 tabulated energies between all possible neighboring base pairs (Mathews et al. 1999),
179 we were able to determine values of E^{stack} , E^{HB} , and $E^{\text{hydrophobic}}$ that are consistent
180 with experimental measurements using singular value decomposition (Khatun et al.
181 2004; Press et al. 2002). The interaction parameters are listed in Tables 9.2 and 9.3.

Table 9.1 The averages and standard deviations of the bonded atom pairs

Bonded atom pair	Distance range (Å)	
$P_i S_i$	4.55 ± 0.09	t1.1
$S_i P_{i+1}$	4.10 ± 0.07	t1.2
$S_i A_i$	4.85 ± 0.15	t1.3
$S_i U_i$	3.74 ± 0.08	t1.4
$S_i G_i$	4.81 ± 0.14	t1.5
$S_i C_i$	3.70 ± 0.13	t1.6
$P_i P_{i+1}$	6.25 ± 0.95	t1.7
$S_i S_{i+1}$	5.72 ± 0.45	t1.8
$P_i A_i$	7.45 ± 0.45	t1.9
$P_i U_i$	5.57 ± 0.37	t1.10
$P_i G_i$	7.43 ± 0.43	t1.11
$P_i C_i$	5.57 ± 0.37	t1.12
$A_i P_{i+1}$	7.25 ± 0.42	t1.13
$U_i P_{i+1}$	6.40 ± 0.20	t1.14
$G_i P_{i+1}$	7.20 ± 0.43	t1.15
$C_i P_{i+1}$	6.40 ± 0.20	t1.16
$P_{i-1} S_i$	9.25 ± 0.95	t1.17
$S_{i-1} P_{i+1}$	8.96 ± 0.44	t1.18
$A_{i-1} S_i$	5.68 ± 0.68	t1.19
$U_{i-1} S_i$	6.38 ± 0.73	t1.20
$G_{i-1} S_i$	5.68 ± 0.68	t1.21
$C_{i-1} S_i$	6.38 ± 0.73	t1.22
$S_{i-1} A_i$	7.25 ± 0.60	t1.23
$S_{i-1} U_i$	5.66 ± 0.54	t1.24
$S_{i-1} G_i$	7.25 ± 0.60	t1.25
$S_{i-1} C_i$	5.66 ± 0.54	t1.26

All the bonds, angles, and dihedrals are effectively modeled using a bonded interaction in the DMD simulations (Fig. 9.1b). A, U, G, and C corresponds to four types of bases (B) t1.27

Table 9.2 The parameters for base pairing, modeled by hydrogen bonds between A–U, G–C, and U–G t2.1

Atom pair	d_{\min} (Å)	d_0 (Å)	d_1 (Å)	d_{\max} (Å)	
Ci–Gj base pair					t2.2
Si Gj	7.70	8.08	8.63	9.00	t2.3
Ci Sj	9.74	10.10	10.53	10.82	t2.4
Ai–Uj base pair					t2.5
Si Uj	9.76	9.94	10.50	10.76	t2.6
Ai Sj	7.72	7.92	8.82	9.00	t2.7
Ui–Gj base pair					t2.8
Si Gj	7.00	7.44	8.24	8.70	t2.9
Ui Sj	9.50	10.25	10.80	11.35	t2.10

The details of the DMD algorithm for the hydrogen bond can be found in Ding et al. (2003). The schematic interaction potential is shown in Fig. 9.2c. The hydrogen bond strengths, E^{HB} , for A–U, G–C, and U–G are 0.5, 1.2, and 0.5 Kcal/mol, respectively. The interaction potential between the donor and acceptor is $-E^{\text{HB}}$ t2.11

t3.1 **Table 9.3** The stacking and hydrophobic interaction strengths, expressed in kcal/mol units

t3.2	E^{Stack}	A_U	U_A	G_C	C_G	G_U	U_G
t3.3	A_U	-0.45	-0.50	-0.75	-0.95	-0.42	-0.70
t3.4	U_A	-0.50	-0.40	-0.55	-0.60	-0.35	-0.35
t3.5	G_C	-0.75	-0.55	-0.81	-0.95	-0.48	-0.92
t3.6	C_G	-0.95	-0.60	-0.95	-1.10	-0.47	-0.51
t3.7	G_U	-0.42	-0.35	-0.48	-0.47	-0.52	0.62
t3.8	U_G	-0.70	-0.35	-0.51	-0.51	0.62	-0.44
t3.9	$E^{\text{Hydrophobic}}$	A_U	U_A	G_C	C_G	G_U	U_G
t3.10	A_U	-0.25	-0.40	-0.40	-0.50	-0.25	-0.35
t3.11	U_A	-0.40	-0.30	-0.25	-0.25	-0.25	-0.25
t3.12	G_C	-0.40	-0.25	-0.25	-0.45	-0.25	-0.41
t3.13	C_G	-0.50	-0.25	-0.45	-0.50	-0.25	-0.41
t3.14	G_U	-0.25	-0.25	-0.25	-0.25	-0.30	0.25
t3.15	U_G	-0.35	-0.25	-0.41	-0.41	0.25	-0.25

t3.16 The subscript indicates that the base bead is paired. For example, A_U is a base bead A that has been paired with a U bead. The cutoff distance for stacking interactions is 6.0 Å. The cutoff distance for hydrophobic interactions is 6.5 Å. The hardcore distance between all beads is set as 3.0 Å

182 *Loop Entropy.* Loop entropy plays a pivotal role in RNA folding kinetics and
 183 thermodynamics (Tinoco and Bustamante 1999). Hence, RNA folding prediction
 184 methods should take this entropic effect into account, either implicitly as in all-
 185 atom MD simulations (Sorin et al. 2004) or explicitly as in Monte Carlo or dynamic
 186 programming methods (Mathews 2006; Rivas and Eddy 1999). However, the
 187 reduction of degrees of freedom in our simplified RNA model causes entropy to
 188 be underestimated in DMD simulations. For example, we often observe formation of
 189 large loops that traps RNA molecules in nonnative conformations for significant
 190 simulation times. To overcome such artifacts arising from the coarse-graining pro-
 191 cess, we developed a simple modification of DMD simulation to model loop entropy
 192 explicitly. We use the free energy estimations for different types of loops, including
 193 hairpin, bulge, and internal loops (Mathews et al. 1999). Loop free energies were
 194 obtained from experimental fitting for small loops and extended to arbitrary lengths
 195 according to polymer theory. We compute the effective loop free energy in DMD
 196 simulations based on the set of base pairs formed in simulations. Upon the formation
 197 or breaking of each base pair, the total loop free energy changes according to the
 198 changes in either the number or size of loops. We estimate the changes in loop free
 199 energy, ΔG^{loop} , for each base pair formed during the simulation and determine the
 200 probability of forming such a base pair by coupling to a Monte Carlo procedure using
 201 a Metropolis algorithm with probability $p = \exp(-\beta\Delta G^{\text{loop}})$. If the base pair is
 202 allowed to form stochastically, the particular base pair will form only if the kinetic
 203 energy is sufficient to overcome the possible potential energy difference before and
 204 after the base-pair formation. Upon breaking of a base pair, the stochastic procedure
 205 is not invoked since base-pair breakage is always entropically favorable. The break-
 206 ing of a base pair is only governed by the conservation of momentum, energy, and
 207 angular momentum before and after the base-pair breakage.

The total potential energy, E , is obtained by adding all interaction terms, as given in (9.5):

$$E = E_{\text{Bonded}} + E_{\text{Hbond}} + E_{\text{Stack}} + E_{\text{Hydrophobic}} + E_{\text{Overpacking}} + G_{\text{loop}}, \quad (9.5)$$

and is used to perform DMD simulations of RNA molecules. The energy landscape of RNA molecules is very rugged with a vast number of local minima due to the high degeneracy of nucleotide types (only 4 compared to the 20 different amino acids found in proteins). In order to efficiently sample the conformational space of RNAs, we utilize the replica-exchange sampling scheme (Okamoto 2004; Zhou et al. 2001).

Replica Exchange DMD. In replica exchange computing, multiple simulations or replicas of the same system are performed in parallel at different temperatures. Individual simulations are coupled through Monte Carlo-based exchanges of simulation temperatures between replicas at periodic time intervals. For two replicas, i and j , maintained at temperatures T_i and T_j and with energies E_i and E_j , temperatures are exchanged according to the canonical Metropolis criterion with exchange probability p , where $p = 1$ if $\Delta = (1/k_B T_i - 1 - k_B T_j)(E_j - E_i) \leq 0$, and $p = \exp(-\Delta)$, if $\Delta > 0$. For simplicity, we use the same set of eight temperatures in all replica exchange simulations: 0.200, 0.208, 0.214, 0.220, 0.225, 0.230, 0.235, and 0.240. The temperature is in the abstract unit of kcal/(mol k_B). Note that we approximate the pairwise potential energy between coarse-grained beads with the experimentally determined free energy of nearest neighboring base pairs, instead of the actual enthalpy. As a result, the temperature does not directly correspond to physical temperatures. In DMD simulations, we maintain constant temperature using an Anderson thermostat (Andersen 1980).

Since the DMD code is highly optimized, we have found that the computational timescales linearly with respect to the system size. The folding simulation of a 50-nucleotide-long RNA sequence (median size of RNA chains in the sample) for 2×10^6 DMD simulation time units takes approximately 7 h of total wall-clock time, utilizing eight 2.33-GHz Intel Xeon compute nodes.

9.3 Ab Initio Folding of Small RNA Molecules

For each RNA molecule, we initially generated a linear conformation using the nucleotide sequence alone. Starting from this extended conformation, we performed replica exchange simulations at different temperatures as described above. From the simulation trajectories, we extracted sampled RNA conformational states, including the lowest energy state, the folding intermediate state, and the corresponding thermodynamic data. In Fig. 9.3, we illustrate the folding trajectory of one of the replicas for a turnip yellow mosaic virus (TYMV) pseudoknot (PDB ID: 1A60). An RNA pseudoknot structure has nonnested base pairing and minimally comprises base pairing between a loop region and a downstream RNA segment. Pseudoknots serve diverse biological functions, including

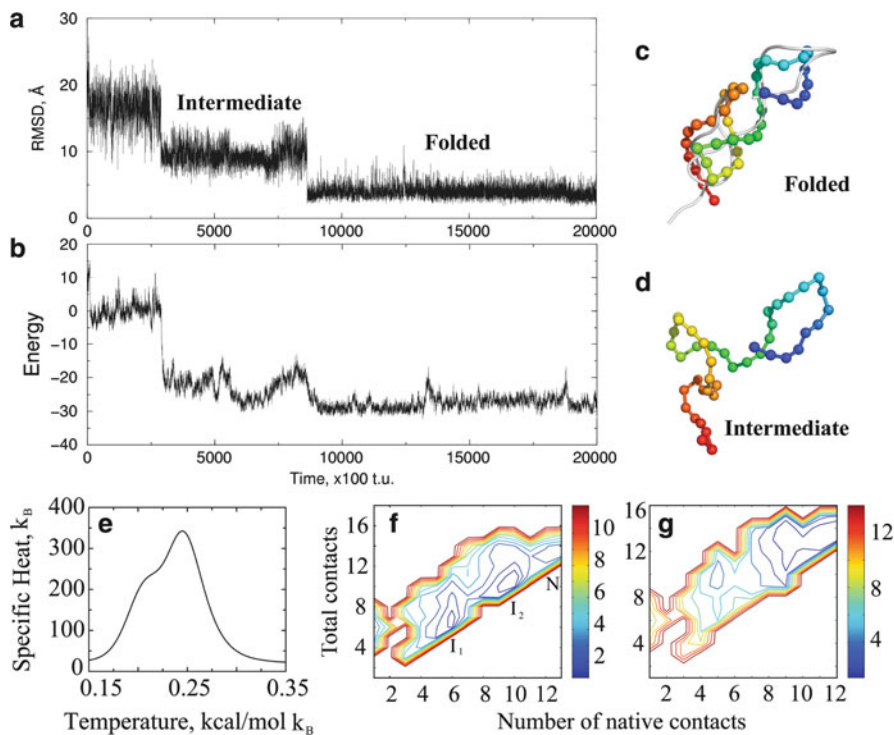


Fig. 9.3 Folding of a pseudoknot. For one replica, we present the RMSD (a) and energy (b) as the function of simulation time. Before folding into its native state (c), the molecule samples a folding intermediate state (d). (e) Specific heat is computed from the replica exchange trajectories using WHAM. (f) Two-dimensional potential of mean force 2D-PMF (potential mean force) for pseudoknot folding at $T^* = 0.245$ (corresponds to the major peak in the specific heat). The two intermediate states and the native state are indicated by I_1 , I_2 , and N , respectively. (g) The 2D-PMF plot at $T^* = 0.21$

247 formation of protein recognition sites that mediate replication and translational
 248 initiation, participation in self-cleaving ribozyme catalysis, and induction of
 249 frameshifts in translation of mRNA by ribosomes (Staple and Butcher 2005). For
 250 example, 1A60 is composed of a 5'-stem and a 3'-pseudoknot (Fig. 9.3c). From the
 251 simulation trajectory (Fig. 9.3), we observe folding of the RNA model within 5 Å
 252 root-mean-square deviation (RMSD) to the native state, and the lowest RMSD from
 253 the simulations is 2.03 Å. The lowest potential energy conformation, computed
 254 across all replicas using the effective free energy function in (9.5), has all native
 255 base pairs formed and an RMSD of 4.58 Å to the native state. Interestingly, we find
 256 that during the folding process the RNA molecule samples a stable folding inter-
 257 mediate state (Fig. 9.3a, b). The intermediate state forms a 5'-stem and a partially
 258 folded 3'-pseudoknot with one of the stems. Our identified folding intermediate
 259 state is consistent with the NMR studies of the solution structures of the TYMV
 260 pseudoknot and its 3'-stem (Kolk et al. 1998). Therefore, our DMD simulation not

only allows the prediction of the native state but also enables us to identify folding intermediate states that might be important for the function of the RNA. The availability of multiple folding trajectories at different temperatures allows quantitative characterization of the folding thermodynamics.

We used the weighted histogram analysis method (WHAM) to calculate folding thermodynamics. The WHAM method utilizes multiple simulation trajectories with overlapping sampling along the reaction coordinates. The density of states $\rho(E)$ is self-consistently computed by combining histograms from different simulation trajectories (Kumar et al. 1992). Given the density of states, the folding specific heat (C_v) can be computed at different temperatures according to the partition function, $Z = \int \rho(E) \exp(-E/K_B T) dE$. To compute the potential of mean force (PMF) as a function of reaction coordinate A , we compute the conditional probability $P(A|E)$ of observing A at given energy E , which is evaluated from all simulation trajectories. Here, the reaction coordinate A can be any physical parameter describing the folding transitions, such as the number of native base pairs, the radius of gyration, or RMSD. The conditional probability $P(A|E)$ can be estimated from the histogram of parameter A for conformation states whose potential energies are within the range of $[E, E + dE]$. The PMF is computed as

$$\text{PMF}(A) = -\ln\left(\int P(A|E)\rho(E) \exp(-E/K_B T) dE\right) + C. \quad (9.6)$$

Here, C is the reference constant, and we assign the lowest PMF a value of zero. Since our simulations start from fully extended conformations, we exclude the trajectories from the first 5×10^5 time units and use those of the last 1.5×10^6 time units for WHAM analysis. We used the trajectories from all replicas to compute histograms. In Fig. 9.3e–g, we illustrate the folding thermodynamics of 1A60 using WHAM analysis, including the specific heat and potential mean field. The specific heat (Fig. 9.3e) has one peak centered at temperature $T^* = 0.245$ and a shoulder near $T^* = 0.21$, suggesting the presence of intermediate states in the folding pathway. The thermodynamic folding intermediate species is characterized by computing the two-dimensional potential of mean force (2D-PMF) as a function of the total number of base pairs (N) and the number of native base pairs (NN). The 2D-PMF plots at temperatures corresponding to the two peaks in the specific heat (Fig. 9.3f, g) show two intermediate states with distinct free energy basins: the first intermediate state corresponds to the folded 5'-hairpin, while the second intermediate corresponds to the formation of one of the helix stems for the 3'-pseudoknot. For example, the 2D-PMF plot at $T^* = 0.21$ (Fig. 9.3g) shows that the shoulder in the specific heat plot corresponds to the formation of the second intermediate state. The basins corresponding to the two intermediate states have a weak barrier, resulting in a lower peak height in the specific heat plot. Therefore, the coarse-grained RNA model combined with the DMD sampling algorithm allows the modeling of RNA structure as well as folding thermodynamics.

We benchmarked the DMD–RNA model on a set of 153 RNAs with length up to 100 nucleotides (Ding et al. 2008). For a majority of the simulated RNA sequences,

302 the lowest energy structures from simulations have a percentage of native base
303 pairs, or Q -value, close to unity, suggesting the correct formation of native base
304 pairs in simulations. Here, we only considered the base pairs of A–U, G–C, and
305 U–G. The other commonly observed Wobble pairing, A–G, was not included in the
306 benchmark study but will be included in future studies. The average Q -value for all
307 153 RNA molecules studied is 94%. For comparison with available secondary
308 structure prediction methods, we also computed the Q -values using Mfold, which
309 yielded an average Q -value of 91%. Given the high percentage of correctly
310 predicted base pairs (94%) and the relatively simple topology of the studied RNA
311 molecules, the average number of incorrectly predicted base pairs is less than one.

312 The RMSD between predicted and experimental structures is often computed to
313 evaluate the accuracy of predicted tertiary structures. Although the RMSD calculation
314 does not provide detailed information on local structural features such as base pairing
315 and base stacking, it gives a straightforward measure of the overall structure predic-
316 tion. Recently, we have developed an approach to evaluate the statistical significance
317 of RNA 3D structure prediction with a given RMSD for different lengths (Hajdin et al.
318 2010). Alternatively, Parisien et al. (2009) have proposed new metrics to account for
319 both local and global structural information during structural comparison. However,
320 their calculation requires the atomic structure of the prediction. To evaluate the overall
321 3D fold of our coarse-grained models, we computed the RMSD to compare our
322 predictions with experimental structures. We found that for RNA molecules with
323 nucleotide length < 50 nt, the RMSD of predicted structures are less than 6 Å.
324 Predictions of longer RNAs exhibit larger RMSD due to the highly flexible nature
325 of RNA molecules. Among the 153 sequences simulated, 84% of the predicted tertiary
326 structures have an RMSD of < 4 Å with respect to the experimentally derived native
327 RNA structure. The benchmark results highlight the predictive power of the
328 DMD–RNA methodology, at least for small RNA molecules.

329 Three out of 153 RNA molecules studied are longer than 65 nucleotides, where
330 the DMD–RNA method cannot be applied to predict the native secondary and
331 tertiary structure from sequence alone. The challenges to predict large RNA folding
332 *ab initio* arise from the exponentially increasing size of the conformational space
333 and inaccuracies in the force field. Therefore, it is important to develop new
334 approaches to predict the 3D fold of large RNA molecules.

335 **9.4 Automated RNA Structure Determination Using** 336 **Experimental Constraints**

337 RNA structural information including secondary structure and some tertiary
338 interactions can often be derived experimentally and computationally prior to the
339 determination of high-resolution 3D structure. Accurate RNA secondary structures
340 can be obtained from comparative sequence analysis (Gutell et al. 2002; Michel and
341 Westhof 1990) and experimentally constrained prediction (Deigan et al. 2009a).

SHAPE chemistry (selective 2'-hydroxyl acylation analyzed by primer extension) 342 was recently shown to be a powerful approach for analyzing secondary structure at 343 single nucleotide resolution for RNAs of any length (Merino et al. 2005; Wilkinson 344 et al. 2006). SHAPE exploits the discovery that the 2'-OH group in unconstrained or 345 flexible nucleotides reacts preferentially with hydroxyl-selective electrophilic 346 reagents. In contrast, nucleotides constrained by base-pairing or tertiary 347 interactions are unreactive. The resulting reactivity information can be used, in 348 concert with a secondary structure prediction algorithm, to obtain accurate second- 349 ary structures (Deigan et al. 2009b; Mathews et al. 2004; Mortimer and Weeks 350 2007; Wang et al. 2008; Wilkinson et al. 2008). Long-range interactions of RNA 351 molecules can also be inferred by biochemical and bioinformatic methods, such as 352 dimethyl sulfate (DMS) modification (Jan and Sarnow 2002; Flor et al. 1989), 353 hydroxyl radical protection (Murphy and Cech 1994), mutational analysis 354 (Kanamori and Nakashima 2001; De la Pena et al. 2003; Khvorova et al. 2003; 355 Murphy and Cech 1994; Wang et al. 1995), and sequence covariation (Cannone 356 et al. 2002). Therefore, we propose to incorporate experimentally determined 357 secondary and tertiary structure information into DMD simulations to reconstruct 358 a conformational ensemble that is consistent with experimental measurements. 359

In general, existing programs for modeling complex RNAs use either computa- 360 tionally intensive all-atom reconstruction, which limits their applications to small 361 RNAs, or overly simplified models that omit key structural details. Other challenges in 362 many current approaches are requirements for high levels of expert user intervention 363 or comparative sequence information and the reliance on chemical intuition derived 364 from preexisting information on tertiary interactions [reviewed in (Shapiro et al. 365 2007)]. Here, we developed an approach for accurate de novo determination of 366 RNA tertiary fold that does not require expert user intervention nor impose heavy 367 computational requirements, and that is efficient for large RNAs (Fig. 9.4). The 368 approach takes an input list of base pairs and distance constraints between specific 369 pairs of nucleotides and outputs a structural ensemble that is consistent with the input 370 constraints. Starting from the extended conformation, we performed DMD 371 simulations with biased potential for base-pairing constraints. Iterative DMD optimi- 372 zation was performed until all base pairs formed. After base-pair formation was 373 confirmed, long-range interaction constraints were added for DMD simulated 374 annealing simulations. At the end of each simulated annealing simulation, we devised 375 filters to evaluate the simulation results, including radius of gyration and/or number of 376 satisfied long-range constraints. We performed iterative annealing simulations until 377 all filters were satisfied and, after constructing the structural ensemble from simulation 378 trajectories, performed cluster analysis to identify representative structures. In all 379 DMD simulations, only serial computation (instead of replica exchange) was used, 380 which also reduced the computational requirement. 381

We tested the automated structure refinement method on tRNA^{asp} (Gherghe et al. 382 2009). Base pairing from the X-ray crystallography structure was consistent with the 383 SHAPE-derived secondary structures. Long-range distance constraints were deter- 384 mined using a site-directed footprinting experiment. An Fe(II)-EDTA moiety was 385 tethered specifically to RNA using the site-selective intercalation reagent 386

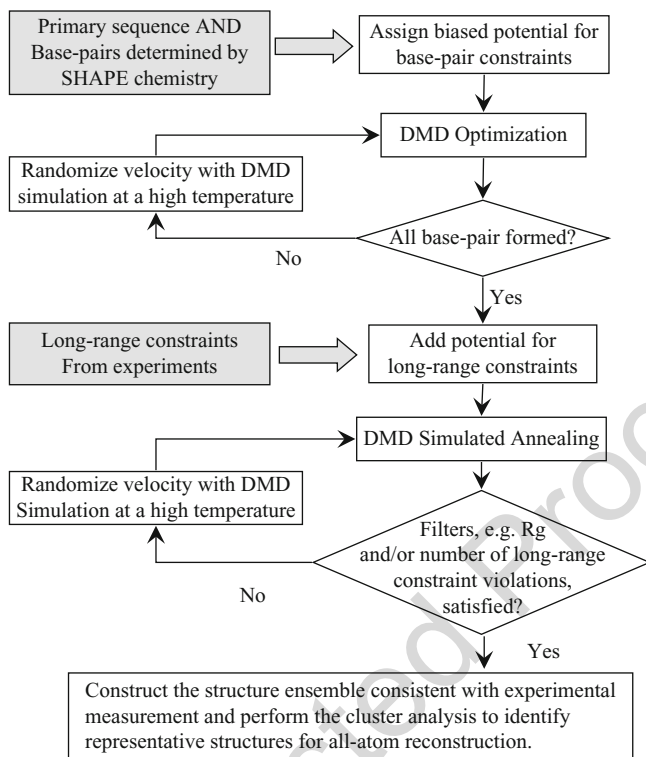
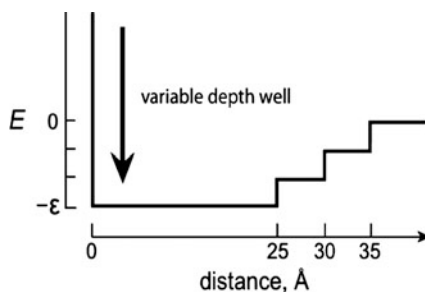


Fig. 9.4 Flowchart of the DMD-RNA structure determination method using experimentally derived structural information

387 methidiumpropyl-EDTA (MPE) (Hertzberg and Dervan 1982). MPE preferentially
 388 intercalates at CpG steps in RNA at sites adjacent to a single-nucleotide bulge (White
 389 and Draper 1987; White and Draper 1989), which can be introduced by mutations in
 390 helical regions. To apply the cleavage information to bias DMD simulations, we
 391 developed a generic approach to interpret each cleavage event as a distance constraint
 392 (Fig. 9.5). The interaction potential features a “soft” energy wall at 25 Å, with smaller
 393 energy bonuses extending out to 35 Å (Fig. 9.5). The 25-Å barrier corresponds to the
 394 distance cutoff within which the nucleotides exhibit strong cleavage and beyond
 395 which the nucleotides have weak cleavage. The interaction strength is assigned
 396 according to the cleavage intensity [$E \propto \ln(I/\langle I \rangle)$]. This approach has two
 397 advantages: (1) no user input is required to decide whether a given cleavage is
 398 significant or not and (2) structure refinement is highly tolerant of measurement errors
 399 inherent in any hydroxyl radical footprinting experiment. By using this structure
 400 determination approach (Fig. 9.5), we were able to refine the structure of tRNA^{asp} to
 401 6.4 Å RMSD relative to the crystal structure (Gherghe et al. 2009).

402 Recently, we applied the structure refinement methodology on four RNAs:
 403 domain III of the cricket paralysis virus internal ribosome entry site (CrPV)

Fig. 9.5 Potential function used to convert experimental cleavage information into DMD potential energy constraints



(49 nts), a full-length hammerhead ribozyme from *S. mansoni* (HHR) (67 nts), 404
S. cerevisiae tRNA^{Asp} (75 nts), and the P546 domain of the *T. thermophila* group 405
 I intron (P546) (158 nts). Each of these RNAs has a complex three-dimensional fold, 406
 involving more than simple intrahelix interactions. Prior to publication of the high- 407
 resolution structures (Cate et al. 1996; Costantino et al. 2008; Martick and Scott 2006; 408
 Westhof et al. 1988), significant biochemical or bioinformatic data describing tertiary 409
 interactions were available for each RNA. The secondary structure was also known to 410
 high accuracy in each case. Only this prior information was used during DMD 411
 refinement. In all cases, we were able to generate a low-RMSD structure. The 412
 RMSD between the predicted structure and the native state for the CrPV, HHR, 413
 tRNA^{Asp}, and P546 RNAs are 3.6, 5.4, 6.4, and 11.3 Å, respectively (Lavender et al. 414
 2010). Calculations were performed on a Linux workstation (Intel Pentium 4 proces- 415
 sor, 3.2 GHz) and the CPU times ranged from 18 (CrPV, 49 nts) to 42 h (P546, 416
 158 nts). Therefore, the combination of efficient DMD simulations and sufficient 417
 biochemical experiments can accurately determine RNA structure of arbitrary length. 418

9.5 Conclusions

419

We have developed a multiscale RNA modeling approach to model 3D structure 420
 and dynamics of RNAs having a wide range of lengths. We use a coarse-grained 421
 representation of the RNA to efficiently model the conformational space. For short 422
 RNA molecules (<50 nt), we are able to capture the folded state from the sequence 423
 alone. The availability of replica-exchange simulation trajectories at multiple 424
 temperatures allows for the characterization of folding thermodynamics as well 425
 as capture of the final folded state. To efficiently sample the exponentially increas- 426
 ing conformational space of large RNA molecules, we devised an automated 427
 modeling approach to determine large and complex RNA structures using experi- 428
 mentally derived structural information. A benchmark study (Lavender et al. 2010) 429
 highlights the application of combining DMD simulation and experimental struc- 430
 tural information to yield native-like models for the diverse universe of functionally 431
 important RNAs whose structures cannot be characterized by conventional 432
 methods. 433

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