The Free Energy Landscape of Pseudorotation in 3′–5′ and 2′–5′ Linked Nucleic Acids

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Supporting Information

ABSTRACT: The five-membered furanose ring is a central component of the chemical structure of biological nucleic acids. The conformations of the furanose ring can be analytically described using the concept of pseudorotation, and for RNA and DNA they are dominated by the C3′-endo and C2′-endo conformers. While the free energy difference between these two conformers can be inferred from NMR measurements, a free energy landscape of the complete pseudorotation cycle of nucleic acids in solution has remained elusive. Here, we describe a new free energy calculation method for molecular dynamics (MD) simulations using the two pseudorotation parameters directly as the collective variables. To validate our approach, we calculated the free energy surface of ribose pseudorotation in guanosine and 2′-deoxyguanosine. The calculated free energy landscape reveals not only the relative stability of the different pseudorotation conformers, but also the main transition path between the stable conformations. Applying this method to a standard A-form RNA duplex uncovered the expected minimum at the C4′-endo state. However, at a 2′–5′ linkage, the minimum shifts to the C2′-endo conformation. The free energy of the C3′-endo conformation is 3 kcal/mol higher due to a weaker hydrogen bond and a reduced base stacking interaction. Unrestrained MD simulations suggest that the conversion from C4′-endo to C2′-endo and vice versa is on the nanosecond and microsecond time scale, respectively. These calculations suggest that 2′–5′ linkages may enable folded RNAs to sample a wider spectrum of their pseudorotation conformations.
advanced simulation methods such as umbrella sampling and metadynamics. These free energy calculations have provided critical insights into many fundamental chemical or biophysical problems by allowing one to determine not only the population of any given conformer but also the free energy along a transition path that connects two stable conformers.

The ideal CVs for studying the conformational changes of sugar puckers are the two pseudorotation parameters, the phase angle $P$ and the amplitude $\tau_m$, which were first proposed by Altona and Sundaralingam and subsequently generalized by Cremer and Pople. These two parameters analytically describe the complete spectrum of all possible puckered states. Following the Cremer–Pople definition, early theoretical studies calculated the potential energy surface of the phase angle $P$ for several nucleoside analogues in vacuo and revealed that the conversion between $C_2$-endo and $C_3$-endo occurs via the $O_2'$-endo pathway, with an energy barrier of 2–3 kcal/mol. The alternative $O_4'$-exo pathway has a much higher barrier and is thus less favorable. The high computational cost, however, precluded the possibility of applying such ab initio methods to nucleic acids in explicit solvent. For these larger systems typically with no less than tens of thousands of atoms, MD simulation based on empirical force fields is an attractive approach that balances computational cost and accuracy. Indeed, starting with a minimized conformation, benchmark simulations of various ribo- and deoxyribo-nucleosides as well as DNA, RNA, and DNA/RNA hybrid duplexes reproduced the stable sugar pucker modes observed in experiments. The sampling from these unrestrained simulations, however, is often insufficient to identify high free energy barriers or other minima that are separated by such barriers. As such, a complete free energy landscape of the entire pseudorotation cycle has not yet been described for simple nucleosides.

As a step toward elucidating the structural effects of $2'-5'$ linkages upon functional RNAs, and to provide a unified description of the various puckered conformations of nucleic acids, we developed a computational method that directly used $P$ and $\tau_m$ as the CVs to calculate pseudorotation free energy. Compared to previous work that used root-mean-square-displacement as an indirect CV, this new approach overcame the insufficiency of sampling and allowed us to sample the entire two-dimensional pseudorotation cycle. In two nucleoside model systems, the calculation not only accurately predicted the free energy difference between the two main pucker states, $C_2$-endo and $C_3$-endo, but also determined the free energy of key transition intermediates, the $O_4'$-endo and $O_4'$-exo states. Application to a native RNA duplex revealed the $C_3$-endo state as the single dominant free energy minimum. For a $2'-5'$-linked nucleotide in an otherwise identical RNA duplex, our free energy calculation revealed a flattened free energy landscape, in which the $C_2$-endo state is only 3 kcal/mol more stable than the $C_3$-endo conformation. The interconversion between these two states is coupled to a switch of hydrogen bonds between $O_2'-\text{H}$ and the pro-$S'$-oxygen or pro-$R'$-oxygen. The free energy barrier crosses the $C_4'$-exo state, with no intramolecular hydrogen bond between $3'-\text{OH}$ and either nonbridging oxygen of the downstream phosphate.

This conclusion is supported by 420 unrestrained MD simulation trajectories in which the nucleotide spontaneously migrates to the more stable $C_2$-endo state from the initial $C_3$-endo conformation. From these simulations, we estimated that the transition from $C_1$-endo to $C_2$-endo occurs in nanoseconds, whereas the reverse transition requires approximately microseconds. Such a rapid interconversion will allow $2'-5'$-linked RNA to swiftly sample various pseudorotation states across the flattened energy landscape, and such an expanded conformational flexibility could allow a single RNA sequence to sample multiple functional conformations.

### MATERIALS AND METHODS

#### Simulation Systems.
Guanosine (rG) and 2'-deoxyguanosine (dG) nucleosides were solvated in $29 \times 29 \times 34 \text{Å}^3$ TIP3P water boxes with a total of $\sim 2.6 \times 10^3$ atoms for each system. The native RNA duplex (5'-CCGCGCGCGG-3') was based on the reported 1.32 Å resolution X-ray crystal structure (PDB code: 4MS9), and the C5-2'-5'-linked RNA duplex (5'-CCGCCGGCGCGG-3', where the asterisk represents a $2'$-5' phosphodiester bond) was based on the reported 1.55 Å resolution structure (PDB code: 4MSB). Both systems were set up as previously described. The final systems contained $\sim 1.2 \times 10^4$ atoms including RNA, water, and ions. Simulation setup and molecular visualizations were performed using VMD.

#### Unrestrained MD Simulations.
A total of 1.4 μs of unrestrained MD simulations (Table S1, Supporting Information) were performed using the program NAMD 2.98 with the CHARMm36 parameter set. All simulations were performed using periodic boundary conditions in the isobaric–isothermal (NPT) ensemble. Langevin dynamics was used to keep the temperature at 298 K with a damping constant of 5 ps$^{-1}$, and a Langevin piston was applied to maintain the pressure at 1 atm. The bonded, nonbonded, and electrostatic interactions were calculated at time steps of 1, 2, and 4 fs, respectively. The switching (cutoff) distance for nonbonded interactions was set at 10 Å. To compute long-range electrostatic interactions, the Particle Mesh Ewald method with a grid density of at least 1 Å$^{-1}$ was used.

To provide a benchmark for the pseudorotation of nucleosides, 300 ns unrestrained simulations were performed for both rG and dG (Table S1, Supporting Information). Both systems were first subjected to 10 000 steps of minimization using the conjugate gradient method. To enhance the sampling as well as estimate the sampling variability, five replica runs were set up using the same minimized initial structure but different initial velocities. Each replica was equilibrated for 10 ns followed by 50 ns production runs. The coordinates of the simulation trajectories at 0.5 ps intervals were used to calculate $p(P,\tau_m)$, the probability of observing a given pseudorotation conformation. The pseudorotation free energy is then calculated by $F(P,\tau_m)! = -k_BT \ln p(P,\tau_m)$.

To study the kinetics of the $C_3$-endo to $C_2$-endo transition of C5 in the C5-2'-5'-linked RNA duplex, 40 distinct conformations sampled at 100 ps intervals were extracted from a 4 ns simulation in which the pucker state of C5 was restrained at the $C_3$-endo conformation. Each configuration was used as the initial coordinate for 20 1-ns unrestrained simulations with different initial velocities. The aggregate simulation time used to study this conformational change was 800 ns (Table S1, Supporting Information).

#### Umbrella Sampling Using Pseudorotation Coordinates.
We implemented $P$ and $\tau_m$ as the CV for umbrella sampling and metadynamics in a modified version of NAMD 2.98 using the Cremer–Pople pseudorotation definition, from which the gradient of $P$ and $\tau_m$ can be computed analytically. The derivation of these equations, together with a brief introduction to the Cremer–Pople definition, is provided in the Supporting Information. In the Results and Discussion section, the $P$ and $\tau_m$ values calculated according to the Cremer–Pople definition are converted to the Altona–Sundaralingam definition that is more widely used in the literature using the following equations:

\[
p^{\text{SP}}(\psi) = p^{\text{CP}}(\psi) + 90^\circ \tag{1}
\]

\[
\tau_m^{\text{AS}}(\psi) = \tau_m^{\text{SP}}(\psi) \times 102.5(\psi/\text{Å}) \tag{2}
\]

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The umbrella sampling calculation of nucleoside pseudorotation comprises five replicas of simulations at 298 K with a biasing harmonic potential centered on \( P \) (varying successively from \(-180 \) to \(180^\circ \) every \(10^\circ \) with a force constant of 0.007 kcal mol\(^{-1}\) deg\(^{-2}\)) and \( \tau_m \) (varying successively from 0.2 to 0.55 Å every 0.05 Å with a force constant of 280 kcal mol\(^{-1}\) Å\(^{-2}\)). Each simulation was 400 ps, and the distribution of \( (P, \tau_m) \) from the last 280 ps trajectory was used as the input to reconstruct the unbiased free energy surface. The weighted histogram analysis method (WHAM)\(^{11}\) with Bayesian bootstrapping\(^{12}\) was applied to generate 200 bootstrapped free energy surfaces from 1440 histograms of the five replica runs (Table S1, Supporting Information). A reliable free energy reconstruction requires significant overlap between the histograms from adjacent simulation windows. Indeed, extensive overlap was observed for both \( rG \) (Figure S3, Supporting Information) and \( dG \) (Figure S4, Supporting Information). The average and standard deviation of the free energy was calculated on the basis of these 200 bootstrapped free energy surfaces. The source code for the analyses was developed on the basis of the one-dimensional WHAM code written by David Minh,\(^{10}\) and has been deposited to simtk.org.

For \( CS \) in the native RNA duplex, the umbrella sampling calculation focused on the “east half” of the pseudorotation space, with biasing potential centered on \( P \) (varying successively from \(-90 \) to \(90^\circ \) every \(10^\circ \) with a force constant of 0.008 kcal mol\(^{-1}\) deg\(^{-2}\)) and \( \tau_m \) (varying successively from 0.15 to 0.50 Å every 0.05 Å with a force constant of 320 kcal mol\(^{-1}\) Å\(^{-2}\)). For \( CS \) in \( CS-2'-S' \)-linked RNA duplex, a similar setup was used except \( P \) was sampled at every \(7.5^\circ \) and \( \tau_m \) was sampled from 0.10 to 0.50 Å with 0.05 Å intervals. Each simulation was performed for 800 ps, and the last 500 ps was used for free energy calculation. As above, the distributions from adjacent simulation windows overlap extensively (Figure S5, Supporting Information) for native RNA duplex; Figure S6, Supporting Information, for \( CS-2'-S' \)-linked RNA duplex). For each duplex, five replicas were performed, and the free energy surface was reconstructed using the same protocol as for free nucleosides.

## RESULTS AND DISCUSSION

### Free Energy Landscape of Pseudorotation in Guanosine

We first calculated the free energy surface defined by \( P \) and \( \tau_m \) for guanosine (\( rG \)) in solution by directly using \( P \) and \( \tau_m \) as the collective variables in umbrella sampling (see Materials and Methods for computational details). The resulting free energy surface represents the intrinsic properties of guanosine pseudorotation in the absence of other structural constraints imposed by secondary or tertiary structures of RNA, and therefore, it serves as an important reference for understanding nucleic acid pseudorotation. Previous NMR\(^{36}\) and crystallographic\(^{37}\) studies provide critical experimental data for testing the accuracy and validity of our computational method.

The calculated free energy surface of \( rG \) pseudorotation is depicted in Figure 2A. The lowest free energy state corresponds to the \( C_2'-endo \) conformation with \( P \) from 160 to 180° and \( \tau_m \) between 35 and 42° (Figure 2D). This agrees well with the one conformer observed in the crystal structure of guanosine dihydrate (\( P = 161.4^\circ \), \( \tau_m = 36.2^\circ \)).\(^{2,37}\) The other conformer adopts a \( C_2'-exo \) conformation (\( P = 139.2^\circ \), \( \tau_m = 44.3^\circ \)) that is different from many other purine nucleosides or their derivatives.\(^{2} \) This unusual \( C_2'-exo \) conformation is likely due to crystal packing. Our free energy calculation also located a second minimum around \( P = 20^\circ \) that corresponds to the \( C_2'-endo \) conformation (Figure 2B). The free energy difference between these two states is 1 kcal/mol, in excellent agreement with 0.8 kcal/mol derived from previous NMR measurements based on \( ^{3}J_{H_1'-H_2'} \).\(^{36}\) For \( S'-\)guanosine monophosphate, our
previous NMR measurements yield a free energy difference of 0.5 kcal/mol,\(^\text{38}\) suggesting that the equilibrium between these two states is largely unaffected by the C\(_3\) exocyclic substituents.

The calculated free energy landscape also predicts the free energy barriers of the transitions between the two minima and the corresponding pseudorotation states. The main transition pathway, via an O\(_4\)-\textit{endo} intermediate, has a barrier of 2 kcal/mol, which can be attributed to the eclipsed O\(_2\)–C\(_3\)–C\(_4\)–O\(_3\) torsion angle (Figure 2C). The height of this barrier is low enough to be sampled by sufficiently long unrestrained MD simulations and serves as an additional benchmark for our free energy calculations. Five independent unrestrained simulations (60 ns each) were performed. During the 250 ns aggregate production time, 848 transitions between C\(_2\)-\textit{endo} and C\(_3\)-\textit{endo} were observed, providing extensive sampling over the “east” half of the pseudorotation cycle. The calculated free energy surface from these unrestrained simulations (Figure S8, Supporting Information) agrees well with the one from umbrella sampling, with an identical 2 kcal/mol free energy barrier. Detailed analyses of these unrestrained simulations regarding the nucleobase orientation and the intramolecular S'-OH–N3 hydrogen bond\(^\text{23,39}\) during the pseudorotation cycle are provided in the Supporting Information. The agreement between umbrella sampling results and independent computational as well as experimental data suggests that our implementation of pseudorotation collective variables allows the accurate calculation of the phase angle, amplitude and the energetics of pseudorotation.

Umbrella sampling can accurately determine the free energies of high free energy conformations, which are otherwise poorly represented or even absent, in unrestrained simulations. Previous ab initio calculations of the conformations of a nucleoside analogue predicted that the O\(_4\)-\textit{exo} conformer is such an unstable intermediate.\(^\text{22}\) This state, together with its adjacent C\(_1\)-\textit{endo} and C\(_4\)-\textit{endo} states, were not observed in our 300-ns unrestrained simulations (Figure S8, Supporting Information) nor in a previous 50-ns benchmark simulation with a slightly different force field.\(^\text{28}\) By employing umbrella sampling with pseudorotation parameters as the collective variables, we could sample these “rare” high energy states and determine their free energies. The O\(_4\)-\textit{exo} state has the highest free energy of 9.5 kcal/mol, and the origin of this high barrier is illustrated in Figure 2E: in addition to the eclipsed O\(_2\)–C\(_3\)–C\(_4\)–O\(_3\) torsion angle, the nucleobase and C\(_5\)' exocyclic substituents are also in steric conflict in the O\(_4\)-\textit{exo} conformation. The calculated two-dimensional free energy surface also revealed that the O\(_4\)-\textit{exo} state prefers a smaller \(r_m\) (Figure 2A) because a more puckered conformation will reduce the C\(_1\)–C\(_4\) distance and aggravate the steric conflict.

\textbf{Free Energy Landscape of Pseudorotation in 2',Deoxyguanosine.} To elucidate how the absence of a 2'-hydroxyl group affects nucleoside pseudorotation, we computed the pseudorotation free energy surface of 2'-deoxyguanosine (dg) in solution. The free energy landscape of dg is generally similar to that of rG but with noticeable alterations (Figure 3A). The absence of the 2'-hydroxyl group releases the constraint imposed by the eclipsed O\(_2\)–C\(_3\)–C\(_4\)–O\(_3\) torsion angle, stabilizing both the O\(_4\)-\textit{exo} (Figure 3E) and O\(_4\)-\textit{endo} states by approximately 2.5 kcal/mol. Consequently, the free energy of the former drops to 7 kcal/mol, whereas the latter merges into a broad basin that includes the C\(_2\)-\textit{endo} and adjacent C\(_1\)-\textit{exo} states. The peak of the main transition path shifts to the C\(_4\)-\textit{exo} state (Figure 3C), with a free energy barrier of 1.5 kcal/mol.

This broad basin is 1.0 kcal/mol more stable than the C\(_1\)-\textit{endo} state (Figure 3B), and the free energy difference again agrees well with previous NMR results\(^\text{38}\) as well as with our unrestrained MD simulations (Figure S9, Supporting Information).

\textbf{Free Energy Landscape of Pseudorotation in a Native RNA Duplex.} To delineate how distinct backbone connectivities can influence the pseudorotation free energy landscape of RNA, it is critical to compare two regioisomers of a nucleotide at the same position within a duplex to avoid potential interference by positional and sequence effects. Our previous crystallographic studies provided one such opportunity through the high-resolution structures of a native S'-CCGGCGCCGG-3' RNA duplex and a regioisomeric S'-CCGGC*GCGGG-3' RNA duplex (C5–2'-S'-linked RNA), where the asterisk represents a 2'-S'-phosphodiester bond.\(^\text{11}\) Here, we first calculated the pseudorotation free energy landscape of C5 within the native RNA duplex using umbrella sampling to cover the “east half” of the pseudorotation cycle, including both C\(_2\)-\textit{endo} and C\(_3\)-\textit{endo} states as well as the more preferable O\(_4\)-\textit{endo} transition pathway. The resulting free energy landscape (Figure 4A) has a deep minimum corresponding to the C\(_3\)-\textit{endo} conformation, with compact stacking of planar base pairs (Figure 4B). The other minimum matches the C\(_2\)-\textit{endo} conformation and is significantly less stable by 6 kcal/mol.
specifically labeling ribose with $^{13}$C (ref 40) and so far has not been performed on an A-form duplex. Therefore, there is no direct experimental data to compare with our calculated barrier. In the GCAA tetraloop, the measured sugar puckering barrier varies from 10 to 18 kcal/mol depending on the local structural context.40 Our calculated value falls within the same range and provides a testable prediction that may stimulate further experimental investigations.

**Free Energy Landscape of Pseudorotation of Nucleotides with 2′−5′ Linkages.** To elucidate the structural effect of 2′−5′ linkages on the pseudorotation of nucleotides in an RNA duplex, we performed a pseudorotation free energy calculation for 2′−5′-linked C5 in an RNA that is regioisomeric to the native RNA duplex described above. As shown in Figure SA, the most stable conformation of the resulting free energy surface corresponded to the C2-end state (Figure SD), with $P$ between 150 and 180° and $\tau_m$ around 35°, in excellent agreement with the crystallographic result ($P = 160 \pm 6^\circ$, $\tau_m = 36 \pm 2^\circ$).11 Previous NMR studies also showed that the C2-end state is preferred in homogeneous 2′−5′-linked RNA duplexes41 and branched trinucleotides containing 2′−5′-linkages,42 suggesting that such a preference may be an intrinsic property of 2′−5′-linked RNAs. The free energy calculation also reveals that the other minimum corresponds to the C3-end conformation ($P$ around 10°, $\tau_m$ around 35°, Figure SB), with a slightly higher free energy of only 3 kcal/mol, which could be overcome with favorable molecular interactions. Indeed, the calculated pseudorotation parameters agree well with a C3-end conformation ($P = 16.7^\circ$, $\tau_m = 40.8^\circ$) that we serendipitously captured in G3 of an RNA duplex with identical sequence but three 2′−5′-linkages11 (5′-CCG*GC*GC*CGG-3′), where the asterisks represent the 2′−5′ phosphodiester bonds, hereafter referred to as “triple 2′−5′-linked RNA”. Compared to the native duplex, which has a deep free energy minimum at the C3-end state, a 2′−5′ linkage profoundly flattens the pseudorotation free energy landscape and shifts the minimum to the C3-end state.

To understand what destabilized the C3-end state, we performed a 50-ns MD simulation in which the ribose of C5 was restrained in the C3-end state ($P = 15 \pm 5^\circ$, Table S2, Supporting Information). A conformational ensemble of the C5-2′−5′-linked RNA duplex with C5 in the C3-end pucker state was generated using 20 000 frames of the last 40-ns trajectory sampled at 2 ps intervals, and was subsequently used to calculate structural parameters including pseudorotation, base pair, and base-pair step parameters. Most of the parameters are indistinguishable from those of the C2-end conformation,11 although the C5 G6:G6 C5 base-pair step in the C3-end state has a smaller overlap area (Tables S2 and S3, Supporting Information). This may weaken the base-stacking and partly explain the instability of the C3-end state.

The MD simulation also confirmed that the conformational shift from C2-end to C3-end concomitantly altered the hydrogen bond between the 3′-OH and the downstream phosphate from O5′-H···pro-Sp-oxygen to O5′-H···pro-Rp-oxygen (Figure SB), a result that was initially inferred on the basis of the oxygen−oxygen distance in the crystal structures.11 A close examination of the conformational ensemble from our restrained simulations further revealed that this hydrogen bond in the C3-end state is weaker than its counterpart in the C2-end state (Figure S13, Supporting Information), with a longer average O−O distance (2.94 Å in C3-end versus 2.86 Å in C2-end), and a larger deviation from the desired in-line
conformation (O–H–O angle is 136 ± 20° in C₃-endo versus 152 ± 12° in C₅-endo). Therefore, the relative instability of the C₅-endo state can be attributed to both a weaker hydrogen bond and weaker stacking, although we cannot rule out other possibilities. According to our free energy calculation, the peak of the C₂-endo to C₅-endo transition path is at the C₃-exo state (P is from 40 to 60°, and τₘ is around 30°), and the corresponding free energy barrier is approximately 6 kcal/mol (Figure S5C). In this C₃-exo state, the intramolecular hydrogen bond between 3'-OH and either nonbridging oxygen of the downstream phosphate is broken (Figure S5C). A similar conformation is observed in C7 of the triple 2'−5'-linked RNA crystal structure, albeit with a slightly larger amplitude (P = 42.8°, τₘ = 42.8°). In this case the 3'-OH group forms a hydrogen bond with a nearby water molecule instead of phosphate (Figure S14, Supporting Information). This supports the notion that in the C₃-exo state there is no intramolecular hydrogen bond between the 3'-OH and the downstream phosphate.

**Estimation of Transition Rates.** To study the kinetics of the interconversion between the C₂-endo and C₅-endo conformations of a 2'−5'-linked nucleotide in an RNA duplex, 800 1-ns unrestrained simulations were carried out from 40 distinct initial coordinate sets with C5 in the C₄-endo conformation. Among these, 420 trajectories (52.5%) successfully reached the more stable C₅-endo conformation with 144° < P < 180°. To further extract kinetic details of these spontaneous conformational changes from C₃-endo to C₅-endo, we focused on 5 ps trajectories immediately prior to forming a stable C₅-endo state. Analysis of these trajectories confirmed that during the spontaneous transition from the C₃-endo to C₅-endo state, the breaking of the O₃−H···pro-RP oxygen hydrogen bond precedes the formation of the O₃−H···pro-SP oxygen hydrogen bond, and both hydrogen bonds are absent in the C₅-exo state (Figure 6). We also noticed that in the initial stage of these spontaneous transitions, the heavy atom distance of the O₃−H···pro-RP oxygen hydrogen bond is already significantly elongated to approximately 3.4 Å, even though CS remains in the C₅-endo state (Figure 6). This suggests that the transition would first occur among a subpopulation of C₃-endo states with weakened O₃−H···pro-RP oxygen hydrogen bonds. These unrestrained simulations also suggest that t₁/₂ of the transition is approximately 1 ns. Given that the C₅-endo state is about 3 kcal/mol more stable than the C₃-endo state, t₁/₂ of the reverse transition (C₅-endo to C₃-endo) should be approximately 150 ns. It should be noted that estimating transition rates from unrestrained MD simulations is still a great challenge in computational chemistry. Indeed, the error of the calculated

![Figure 5. Pseudorotation free energy landscape of a 2'−5'-linked nucleotide in an RNA duplex.](image)
binding and unbinding rates of benzamidine to trypsin was about an order of magnitude, even though the extensive sampling accurately predicted the ligand binding mode and the binding free energy. Therefore, the rates from these calculations should be interpreted cautiously: they only suggest that the transition rates between these two conformations can occur on roughly the ns and µs time scale, respectively. Nevertheless, these calculations suggest a rapid interconversion between the C3′-endo and C3′-endo states of a 2′-5′-linked nucleotide in an RNA duplex.

### CONCLUSION

The reported free energy calculations and unrestrained MD simulations provide important insights into the thermodynamic and kinetic properties of native RNA duplexes and those containing 2′-5′ linkages. Our study highlights a flattened free energy landscape for pseudorotation in 2′-5′-linked nucleotides, which can switch rapidly between the C2′-endo and C3′-endo conformations. This is in contrast with a single dominant C3′-endo minimum found in the native RNA duplex. Therefore, in addition to lowering the melting temperature, the presence of 2′-5′ linkages may expand the conformational space accessible for an RNA duplex. Mechanistically, our calculations demonstrate that hydrogen bonding between the 3′-hydroxyl group and the downstream phosphate serves as a molecular switch for this backbone conformational change.

Our study establishes the feasibility of an atomic-level description of the pseudorotation free energy for free nucleosides, as well as nucleotides in RNA duplexes in solution. Even though some intermediates are too unstable to be sampled efficiently with conventional MD simulation techniques, we were able to gain insights into these structures by implementing a set of collective variables for use in conjunction with advanced free energy calculation methods. The methodology we developed in this study may be further applied to investigate other processes in which pseudorotation plays a significant role, such as nonenzymatic primer extension reactions. Furthermore, applications to other synthetic nucleic acid systems (e.g., threose nucleic acids) may lead to a fuller understanding of the chemical etiology of nucleic acid structure.

### ASSOCIATED CONTENT

#### Supporting Information

Supporting text, Figures S1–S16, Tables S1–S5, and the complete reference 29. This material is available free of charge via the Internet at http://pubs.acs.org/.

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**Notes**
The authors declare no competing financial interest.

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