## Experimental and computational analysis of the transition state for ribonuclease A-catalyzed RNA 2'-O-transphosphorylation

Hong Gu<sup>a</sup>, Shuming Zhang<sup>a</sup>, Kin-Yiu Wong<sup>b,c</sup>, Brian K. Radak<sup>b,d</sup>, Thakshila Dissanayake<sup>b</sup>, Daniel L. Kellerman<sup>a</sup>, Qing Dai<sup>e</sup>, Masaru Miyagi<sup>f</sup>, Vernon E. Anderson<sup>a</sup>, Darrin M. York<sup>b,1</sup>, Joseph A. Piccirilli<sup>e,g,1</sup>, and Michael E. Harris<sup>a,1</sup>

<sup>a</sup>Department of Biochemistry and <sup>f</sup>Center for Proteomics and Bioinformatics, Case Western Reserve University School of Medicine, Cleveland, OH 44106; <sup>b</sup>Center for Integrative Proteomics Research, Biology at the Interface with the Mathematical and Physical Sciences (BioMaPS) Institute for Quantitative Biology and Department of Chemistry and Chemical Biology, Rutgers, The State University of New Jersey, Piscataway, NJ 08854; Departments of <sup>e</sup>Chemistry and <sup>g</sup>Biochemistry and Molecular Biology, University of Chicago, Chicago, IL 60637; <sup>c</sup>Department of Physics, High Performance Cluster Computing Centre, Institute of Computational and Theoretical Studies, Hong Kong Baptist University, Kowloon Tong 852, Hong Kong; and <sup>d</sup>Department of Chemistry, University of Minnesota, Minneapolis, MN 55455

Edited by Philip C. Bevilacqua, The Pennsylvania State University, University Park, PA, and accepted by the Editorial Board June 17, 2013 (received for review September 5, 2012)

Enzymes function by stabilizing reaction transition states; therefore, comparison of the transition states of enzymatic and nonenzymatic model reactions can provide insight into biological catalysis. Catalysis of RNA 2'-O-transphosphorylation by ribonuclease A is proposed to involve electrostatic stabilization and acid/ base catalysis, although the structure of the rate-limiting transition state is uncertain. Here, we describe coordinated kinetic isotope effect (KIE) analyses, molecular dynamics simulations, and quantum mechanical calculations to model the transition state and mechanism of RNase A. Comparison of the <sup>18</sup>O KIEs on the 2'O nucleophile, 5'O leaving group, and nonbridging phosphoryl oxygens for RNase A to values observed for hydronium- or hydroxide-catalyzed reactions indicate a late anionic transition state. Molecular dynamics simulations using an anionic phosphorane transition state mimic suggest that H-bonding by protonated His12 and Lys41 stabilizes the transition state by neutralizing the negative charge on the nonbridging phosphoryl oxygens. Quantum mechanical calculations consistent with the experimental KIEs indicate that expulsion of the 5'O remains an integral feature of the rate-limiting step both on and off the enzyme. Electrostatic interactions with positively charged amino acid site chains (His12/ Lys41), together with proton transfer from His119, render departure of the 5'O less advanced compared with the solution reaction and stabilize charge buildup in the transition state. The ability to obtain a chemically detailed description of 2'-O-transphosphorylation transition states provides an opportunity to advance our understanding of biological catalysis significantly by determining how the catalytic modes and active site environments of phosphoryl transferases influence transition state structure.

**E** nzymes achieve powerful rate enhancements by providing multiple catalytic modes to stabilize reaction transition states, including electrostatic interactions and proton transfer (1). For RNA 2'-O-transphosphorylation reactions, interactions with acid, base, or metal ion catalysts in solution can influence transition state structure (2). Understanding biological catalysis therefore requires knowledge of chemical mechanisms, transition state structure, and transition state interactions for both enzymatic and nonenzymatic RNA cleavage reactions. Comparisons of enzymatic and nonenzymatic catalysis can provide information on which catalytic modes are used by enzymes and whether unique features of the active site environment may alter the transition state charge distribution (3).

RNA undergoes two competing transesterification reactions in solution: isomerization to a 2',5'-phosphodiester and 2'-Otransphosphorylation to yield a cyclic 2',3'-phosphodiester with concomitant release of the 5'O-nucleoside (4, 5). Reactions catalyzed by acid and by buffers yield both isomerization and cleavage products and proceed via a pentacoordinated phosphorane intermediate formed by an attack of the 2'OH at the adjacent phosphorus. The phosphorane intermediate may be neutral ( $I_{neutral}$ ) (Fig. S14) or monoanionic ( $I_{monoanion}$ ) (Fig. S1B). It is not known whether 2'OH deprotonation occurs in a preequilibrium step or if its proton is transferred to water or one of the nonbridging phosphoryl oxygens (NPOs) during nucleophilic attack. Hydroxide catalysis proceeds by equilibrium deprotonation of the 2'OH, followed by nucleophilic attack (Fig. S1C), and only cleavage products are formed (5, 6). Linear free energy relationships (LFERs), stereochemical outcomes, kinetic isotope effects (KIEs), and computational simulations are consistent with either an overall concerted mechanism or transient formation of a dianionic phosphorane intermediate (>pH 9) (4, 5). This complex mechanistic landscape raises the question of which pathway(s) are favored by the active sites of ribonucleases.

RNase A catalyzes RNA 2'-O-transphosphorylation, and investigation of its structure and function have provided foundational insights into the function of enzyme active site residues in acid/base catalytic modes (7). Although the mechanism is usually depicted as concerted, computational studies suggest the potential for a stepwise mechanism via an anionic phosphorane intermediate (8) (Fig. 1A, top pathway). Molecular dynamics (MD) simulations indicate that the RNase A active site provides electrostatic stabilization of the dianionic phosphorane while positioning His119 to act as a general acid (9). His12 and Lys41 are proposed to interact with the 2'OH and/or one of the NPOs, possibly stabilizing the transition state for nucleophilic attack (TS1) or anionic intermediates via H-bonding or proton transfer to form  $I_{neutral}$  or  $I_{monoanion}$ . However, lack of a strong thio effect renders the latter mode unlikely (10). Several lines of evidence implicate His119 as a general acid to facilitate departure of the 5'O in a stepwise or concerted mechanism [via a product-like transition state (TS2) or a concerted transition state  $(TS_c)$ ]. Mutation of His119 has a large deleterious effect on catalysis of RNA cleavage but only a small effect on the reaction of a uridine-3'-p-nitrophenol phosphate model substrate, consistent with

Author contributions: H.G., V.E.A., D.M.Y., J.A.P., and M.E.H. designed research; H.G., S.Z., K.-Y.W., B.K.R., T.D., D.L.K., V.E.A., and M.E.H. performed research; Q.D. and M.M. contributed new reagents/analytic tools; H.G., S.Z., K.-Y.W., B.K.R., T.D., D.L.K., V.E.A., D.M.Y., J.A.P., and M.E.H. analyzed data; and H.G., S.Z., D.M.Y., J.A.P., and M.E.H. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission. P.C.B. is a guest editor invited by the Editorial Board.

<sup>&</sup>lt;sup>1</sup>To whom correspondence may be addressed. E-mail: york@biomaps.rutgers.edu, jpicciri@uchicago.edu, or meh2@case.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1215086110/-/DCSupplemental.



**Fig. 1.** (*A*) Proposed active site interactions and potential transition states for RNase A. A concerted mechanism with simultaneous nucleophilic attack, leaving group departure, and concomitant proton transfer involving both His12 and His119 is shown in the lowest pathway. The most likely stepwise mechanism involves nucleophilic attack (via TS1) and leaving group departure (via TS2) separated by a dianionic intermediate that may be only short-lived. (*B*) RNA dinucleotide substrate (UpG) reacts via 2'-O-transesterification to yield 2',3'-cyclic uridine monophosphate (cUMP) and guanosine (G) products. Sites of isotopic substitution used to measure the KIEs for the <sup>18</sup>k<sub>NUC</sub>, <sup>18</sup>k<sub>NPO</sub>, and 5'O <sup>18</sup>k<sub>LG</sub> are indicated in red, green, and blue, respectively.

a role in leaving group stabilization (7). However, the more highly reactive *p*-nitrophenol leaving group may result in a change in mechanism compared with the less reactive ribose 5'O, making it difficult to interpret these results without an understanding of the mechanism and rate-limiting transition state. Substitution of His12 and His119 with 4-fluorohistidine ( $pK_a = 3.5$ ) changes the apparent macroscopic  $pK_a$  values determined from analysis of the pH dependence of  $k_{cat}/K_m$  with their proposed roles in acid/ base catalysis (11). However, effects of 4-fluorohistidine substitution on catalysis are only *ca.* 10-fold, suggesting a relatively small Brønsted effect, or that a nonchemical step is partially rate-limiting. Thus, even for this well-studied enzyme, significant ambiguity remains regarding the interactions with the transition state that provide catalysis.

A powerful method to gain insight into mechanism and transition state structure is to determine the KIEs on substrate atoms and to use the results as benchmarks for evaluating computational simulations (12, 13). The effects of stable heavy atom isotope substitution on reaction rate constants arise mainly due to differences in vibrational zero point energies, and their magnitudes provide information on the differences in bonding between the ground state and transition state (14, 15). Leaving group and NPO KIEs for RNase A cleavage of uridine-*m*-nitrobenzyl phosphate were previously reported by Cleland and colleagues (16). However, this study lacked information on nucleophile bonding and involved a model substrate that lacks binding interactions present in authentic RNA substrates.

Here, KIE measurements on enzyme and solution RNA reactions are coordinated with MD simulations and quantum mechanical (QM) calculations to achieve a detailed model of the transition state for enzymatic RNA cleavage. Together, the results reveal a transition state with advanced 2'O-P bond formation and 5'O-P bond cleavage similar to the late anionic transition state for solution reactions catalyzed by a specific base. MD simulations are consistent with protonated His12 and Lys41 stabilizing an anionic, phosphorane-like transition state by Hbonding to the phosphoryl oxygens, whereas protonated His119 is poised to transfer a proton to the 5'O leaving group. Importantly, the KIEs for the RNase A reaction reveal a significant difference in 5'O bonding compared with solution reactions. Supported by QM calculations, we ascribe this difference to less advanced P-O bond cleavage and proton transfer (general acid catalysis) from His119.

## **Results and Discussion**

Measurement of <sup>18</sup>O KIEs on RNA 2'-O-Transphosphorylation Reactions. KIEs are typically measured by competitive methods (12) in which mixtures of substrates containing either <sup>16</sup>O or <sup>18</sup>O at one of the reactive phosphoryl oxygens are analyzed (Fig. 1*B*). The larger rate constant for either the  ${}^{18}$ O- or  ${}^{16}$ O-containing RNA results in progressive depletion of that isotopologue in the unreacted substrate population. Quantitative analysis of the <sup>18</sup>O/<sup>16</sup>O ratio as a function of reaction progress allows calculation of the KIE, which is expressed as the ratio of the rate constants for the light and heavy isotopologues  $\binom{18}{k} = k_{16}/k_{18}$  (12). KIEs reflect both the extent to which the labeled atom participates in the reaction coordinate and differences in vibrations of the labeled atom in the ground state compared with the transition state (14). Bond breakage, or a "looser" bonding environment in the transition state relative to the ground state results in faster reactivity for the light isotopologue, and thus an  $^{18}k$  that is greater than unity (referred to as a normal KIE). Conversely, bond formation or a "stiffer" bonding environment favors the heavier isotopologue, and the observed KIE is less than unity (referred to as an inverse KIE). The fractionation of <sup>18</sup>O will also depend on proton transfer in both preequilibrium and chemical steps (12), and may offset the contributions of O-P bonding. Additionally, both bonding and bending vibrational modes can influence the observed KIE on the nonbridging oxygens, and these effects may also be offsetting (13). Deciphering the relative contributions of proton transfer, O-P bonding, and other vibrational modes can be difficult; however, the magnitude of observed KIEs provides insight into mechanism and transition state bonding unobtainable by any other means.

Substrate RNA dinucleotides enriched with <sup>18</sup>O at the 2'O, 5'O, and NPO positions were synthesized (Fig. 1*B*) and reacted with acid and base and with RNase A. The fraction of reaction was measured by integration of RP-HPLC chromatograms in which the <sup>18</sup>O/<sup>16</sup>O ratio was determined by MS (6) (*SI Text* and

Table 1. Comparison of  ${}^{18}k_{LG}$ ,  ${}^{18}k_{NUC}$ , and  ${}^{18}k_{NPO}$  KIE values for 2'-O-transphosphorylation catalyzed by acid, base, and RNase A

Catalyst	$^{18}k_{LG}$	<sup>18</sup> k <sub>NUC</sub>	<sup>18</sup> k <sub>NPO</sub>
RNase A, pH 7	1.014(3)	0.994(2)	1.001(1)
RNase A (QM)	1.026	0.998	1.006
Acid, pH 0	1.005(4)	0.990(4)	0.991(1)
Base, pH 12 (7, 8)	1.037(2)	0.996(2)	0.999(1)
Base, pH 14 (8)	1.034(4)	0.984(3)	_
Base, pH 14 (QM)	1.046	0.973	1.002

SDs for measured KIEs are shown in parentheses (Methods).



**Fig. 2.** Determination of KIEs for RNase A-catalyzed RNA 2'-O-transphosphorylation by analysis of the change in the <sup>18</sup>O/<sup>16</sup>O ratio (ln<sup>18</sup>O/<sup>16</sup>O) as a function of reaction progress (F). The <sup>18</sup>O/<sup>16</sup>O ratio determined by MS is plotted relative to reaction progress, and the data are fit as described in *Methods*. Example fits of data used to calculate the <sup>18</sup>k<sub>LG</sub> (A), <sup>18</sup>k<sub>NPO</sub> (B), and <sup>18</sup>k<sub>NUC</sub> (C) are shown. Experimental data points are shown as filled squares. Simulations for curves representing a range of potential KIEs are shown as gray lines.

Fig. S24). Two product ions contain the isotopically enriched positions: a species at m/z 476/478 resulting from loss of the uridine base and a 211/213 fragment composed of ribose

phosphate (Fig. S2B). The KIE is observed as a change in the relative ratios of the product ions containing only <sup>16</sup>O at all oxygens (M) and those containing <sup>18</sup>O at one of the reacting phosphoryl oxygens (M + 2) (Fig. S2C). Fig. 2 shows the determination of KIEs for RNase A-catalyzed 2'-O-transphosphorylation of uridylyl-3'-guanosine [5'-UpG-3' (UpG)] relative to simulations of <sup>18</sup>O/<sup>16</sup>O ratio vs. *F* (the fraction of substrate UpG reacted to form a product) for a range of potential <sup>18</sup>O KIEs.

Transition States of Solution Reactions Catalyzed by Acid and Base. The nucleophile ( ${}^{18}k_{\rm NUC}$ ), leaving group ( ${}^{18}k_{\rm LG}$ ), and NPO  $(^{18}k_{\rm NPO})$  KIE results presented in Table 1 for acid- and basecatalyzed 2'-O-transphosphorylation reactions of UpG, together with previous biophysical and biochemical studies, provide a framework for understanding the RNase A transition state. For solution 2'-O-transphosphorylation, previous experimental and computational data are consistent with the observed  ${}^{18}k_{\rm NUC}$ being a combination of normal contributions from deprotonation and an inverse effect from advanced 2'O-P bond formation (6, 17). Together, these contributions result in the inverse values observed for the stepwise acid-catalyzed reaction (pH 0) and a similar inverse  ${}^{18}k_{\rm NUC}$  for the late anionic transition state for base catalysis (pH 12) [0.990(4) and 0.996(2), respectively, with standard deviations in parentheses] (6, 18–20). These data are consistent with analyses of nucleophile bonding by LFERs of RNA reactions catalyzed by a specific base, which show that the 2' oxyanion formed in a preequilibrium deprotonation step undergoes substantial loss

of negative charge in the transition state (21). The observed  ${}^{18}k_{\rm NPO}$  is an average of the KIEs on the two NPOs because only one of the two positions is enriched non-specifically (20). Nonetheless, a late TS2 for the base-catalyzed reaction is further indicated by a near-unity  ${}^{18}k_{\rm NPO}$ , which indicates a similar bonding environment in the transition state and in the ground state monoanion. In contrast, formation of a protonated phosphorane during acid catalysis is consistent with the observed inverse  ${}^{18}k_{\rm NPO}$  [0.991(1)] as previously reported for acid catalysis of uridine-3'*m*-nitrobenzyl phosphate transphosphorylation (22).

The <sup>18</sup> $k_{LG}$  for acid catalysis is small [1.005(4)], consistent with models that propose rate-limiting phosphorane breakdown with concomitant transfer of a proton from a phosphoryl oxygen to the 5'O (22) (Fig. S14). Base-catalyzed RNA 2'-O-transphosphorylation is sensitive to the pK<sub>a</sub> of the leaving group, with a  $\beta_{LG}$  of -1.28 indicating that the alkoxide character of the 5'O is advanced (23), which is consistent with the observed large normal <sup>18</sup> $k_{LG}$  value [1.037(2)]. This value is near the expected maximum (*ca.* 1.04) for breakage of an O-P bond (13). Recent QM simulations of the base-catalyzed reaction identify a representative late transition state for breakdown of a transient dianionic phosphorane (TS2<sub>B</sub> in Fig. S1C) that reproduces the observed inverse <sup>18</sup> $k_{NUC}$ , large normal <sup>18</sup> $k_{LG}$ , and near-unity <sup>18</sup> $k_{NPO}$  (6, 20). For a rate-limiting TS1, a normal <sup>18</sup> $k_{NUC}$  and a small normal <sup>18</sup> $k_{LG}$  are predicted (20). For a concerted mechanism a less inverse <sup>18</sup> $k_{NUC}$ , as well as a normal or inverse <sup>18</sup> $k_{NPO}$ , is observed depending on the associative or dissociative character of TS<sub>c</sub> (2, 17, 24).

**RNase A Kinetics and Substrate Binding Commitments.** KIEs measured by competitive methods are effects on  $k_{cat}/K_m$ ; therefore, substrate binding steps may interfere with the KIE measurements for RNase A. If substrate dissociation is slow relative to chemistry, resulting in a "commitment" of the bound substrate to undergo chemical transformation, the commitment generates an attenuation of the observed KIEs relative to their intrinsic values (12). Several lines of evidence support the assertion that observed KIEs on the RNase A reaction approximate intrinsic effects on the chemical step (*SI Text*). Foremost, the KIEs for



Fig. 3. Structure of RNase A transition state mimic derived from a 50-ns MD simulation using the assisted model building with energy refinement (AMBER) FF10 force field in TIP4PEw water (1-ns average). The dianionic transition state is stabilized by interactions between His12, Lys41, and His119 and the transition state. In this model, His119 is poised for general acid catalysis, whereas His12, in its protonated form (after accepting a proton from the nucleophile), provides electrostatic stabilization to the transition state.

RNase A in Table 1 are nonzero, indicating that the chemical step is at least partially rate-limiting (6, 22). The leaving group KIE for RNase A is similar to the value measured by Cleland and colleagues (16) for the slow model substrate uridine-3'*m*-nitrobenzyl phosphate. UpG is also a slow substrate with *ca*. 40-fold lower  $k_{cat}/K_m$  compared with an optimal uridylyl-3'-adenosine [5'-UpA-3' (UpA)] dinucleotide substrate (Fig. S3). The  $k_{\text{cat}}/K_{\text{m}}$  for UpG is similar in magnitude to that for uridine 3'-pnitrophenol phosphate substrate, which has been argued to react with a rate-limiting step subsequent to binding based on lack of sensitivity to solvent viscosity (25). The rate of UpG cleavage by RNase A under  $k_{cat}/K_m$  conditions is also insensitive to viscosity (Fig. S4). The pH dependence of  $k_{cat}/K_m$  for the reaction of the UpG (Fig. S5) is bell-shaped, as expected, and shows apparent macroscopic pK<sub>a</sub> values (5.2  $\pm$  0.2 and 6.3  $\pm$  0.1) that are similar to the microscopic pKa values of His12 and His119 determined by NMR (5.7 and 6.3) (26), which is also consistent with minimal binding commitments under conditions where KIEs were measured.

KIE Analysis of RNase A-Catalyzed RNA 2'-O-Transesterification. The inverse <sup>18</sup>k<sub>NUC</sub> [0.994(2)] for RNase A suggests that 2'O-P bond formation is advanced in the rate-limiting transition state. A similar inverse  ${}^{18}k_{\rm NUC}$  is observed for the stepwise mechanism for acid catalysis in which the phosphorane is sufficiently stable to undergo pseudorotation (Table 1). The base-catalyzed solution reaction also has an inverse  ${}^{18}k_{\rm NUC}$  [0.996(2)], which becomes more inverse [0.984(3)] above the pK<sub>a</sub> of the 2'OH consistent with loss of the contribution from 2'O deprotonation (6). QM simulations of the base-catalyzed reaction that are consistent with the KIE data indicate a late anionic transition state (TS2 in Fig. S1) and suggest the formation of a transient dianionic phosphorane (20, 27). Interpretation of advanced 2'O-P bonding in the RNase A transition state assumes that the measured  ${}^{18}k_{\rm NUC}$  includes a large inverse contribution 2'O-P bonding that is partially offset by a normal equilibrium isotope effect (EIE) from alcohol deprotonation (28). The relative magnitudes of these contributions are difficult to assess, and combinations of less advanced 2'O-P bonding and partial proton transfer in the transition state (general base catalysis) cannot be entirely excluded.

The small  $^{18}k_{NPO}$  for the RNase A reaction indicates that the NPOs experience a similar bonding environment in both the ground state and transition state, which is also consistent with

a late product-like transition state. An  ${}^{18}k_{\rm NPO}$  near unity is inconsistent with protonation of an NPO. In this case, an inverse  ${}^{18}k_{\rm NPO}$  would be predicted similar to that observed for acid catalysis [0.991(1)]. Together with the lack of a thio effect on RNase A (10), these results provide strong evidence against protonation. Rather, the small normal  ${}^{18}k_{\rm NPO}$  we observe suggests that the NPO atoms gain anionic character in the transition state.

The normal  ${}^{18}k_{LG}$  for RNase A indicates breaking of the 5'O-P bond in the rate-limiting transition state similar to catalysis by a specific base in solution (6). However, the  ${}^{18}k_{LG}$  for the enzyme reaction is significantly less than the value observed for the solution reaction [1.014(3) vs. 1.037(2)], indicating a stiffer bonding environment for the 5'O in the enzyme active site. Alternatively, the smaller  ${}^{18}k_{LG}$  for RNase A could result from a small binding commitment, although available experimental evidence suggests this is unlikely (SI Text). In this case, the intrinsic  ${}^{18}k_{NUC}$  would be more inverse than the solution reaction, which seems unlikely because the measured value of ca. 0.994 is already characteristic of advanced 2'O-P bonding. The predicted large normal  ${}^{18}k_{LG}$ would indicate advanced 5'O-P bond cleavage with a minimal effect of enzyme interactions in mitigating the unfavorable accumulation of negative charge. If the effects accurately reflect intrinsic KIEs on the chemical step, the difference in  ${}^{18}k_{LG}$  could reflect less 5'O-P bond cleavage in the RNase A transition state. It is likely that proton transfer to the 5'O would result in a stiffer bonding environment, given the large inverse EIE in the direction of deprotonation of ca. 0.976 (28) and the proximity of His119.

MD Simulations and QM Calculations to Model the Mechanism and Transition State. To gain further insight into the mechanistic implications of the KIE measurements, we performed MD simulations and QM calculations that build on previous work for RNase A (9) and nonenzymatic models (20, 27). Based on the KIE results supporting an anionic transition state with advanced 2'O-P bonding, we first performed MD simulations of RNase A with a dianionic phosphorane transition state mimic as described previously (9). Because transfer of the 2' proton is considered to be complete in this model, His12 and His119 are protonated to characterize the electrostatic stabilization of the anionic transition state. The results summarized in Fig. 3 are derived from a 50-ns MD simulation and suggest that a protonated His12, formed in the general base step of a stepwise mechanism, can further provide electrostatic stabilization of an anionic phosphorane intermediate/transition state. Throughout the simulation, the imidazolium side chain of His119 forms a strong hydrogen bond with



transition state model based on hydrogen bonding in the active site of

the RNase A (B).

Fig. 4. Transition state structures for the nonenzymatic dianionic transition state model for the specific base reaction (A) and for the enzymatic

the 5'O (Fig. 3). Our findings suggest that RNase A can support a mechanism involving a transient anionic phosphorane intermediate or phosphorane-like transition state like the one implicated by theoretical and experimental studies of specific base catalysis (20, 27). Moreover, the MD results provide a basis for construction of a fully QM active site model from which transition state structures can be determined and KIEs can be estimated.

KIEs were calculated for both enzymatic and nonenzymatic reaction models using density-functional theory and a continuum model to treat solvation effects (Methods). The nonenzymatic model corresponds to alkaline conditions and assumes that the deprotonation of the 2'OH occurs in a prior step that is not ratecontrolling. In the enzymatic model, the 2'OH is assumed to be protontated in the ground state; however, whether deprotonation occurs in a preequilibrium step or concomitant with nucleophilic attack is not known. The enzymatic reaction model was based on the MD simulation structures and consisted of the hypothesized dianionic phosphorane stabilized by hydrogen bonds donated from two histidine side chains (protonated imidazoles). Results for the nonenzymatic reaction model were similar to our previous results using an ethylene glycol phosphate methyl ester (20, 27) in which nucleophile bonding is advanced (1.8 Å compared with 1.7 Å for an O-P bond). Here, we included a more realistic ribose-like sugar ring and ethoxide leaving group, and obtained KIE results in better agreement with experimental data (Fig. 4 and Table 1). Comparison of the calculated KIEs for the nonenzymatic QM model with the measured values for the solution reaction catalyzed by a specific base (calculated/ measured) reveal an inverse  ${}^{18}k_{\rm NUC}$  (0.973/0.984), large normal  ${}^{18}k_{\rm LG}$  (1.046/1.034), and near-unity  ${}^{18}k_{\rm NPO}$  (1.002/0.999).

Alternative mechanistic models involving protonation of an NPO or the 5'O alone and 5'O protonation in the presence of Hbonding to an NPO were evaluated (Table S1 and Fig. S6). The calculated and measured KIE values are in good agreement for the enzymatic reaction model involving proton transfer from His119 to the 5'O, together with H-bonding between the active site and the NPO atoms [ $^{18}k_{\rm NUC}$  (0.998/0.994),  $^{18}k_{\rm LG}$  (1.026/ 1.014), and  ${}^{18}k_{\rm NPO}$  (1.006/1.001) (calculated/measured)] (Table 1). Importantly, the calculated  ${}^{18}k_{\rm LG}$  for the base-catalyzed reaction simulation (1.048) is notably larger than the calculated  ${}^{18}k_{\text{LG}}$  for the enzymatic model (1.026), consistent with the observed experimental trend (1.034 vs. 1.017). In the simulations, the P-5'O bond length is considerably shorter for the RNase A transition state than that for the base-catalyzed reaction (1.95 Å vs. 2.3 Å) (Fig. 4) and it retains a higher degree of covalent bond character. Moreover, proton transfer from the general acid (His119) further creates a stiffer bonding environment for stabilizing the leaving group.

The results reported here are comparable to those of previous studies directed at determining the influence of active site environments on transition state structure. Herschlag and colleagues (29, 30) reported a large negative  $\beta_{LG}$ , as well as a large leaving group KIE for both alkaline phosphatase (AP) and nonenzymatic monoester hydrolysis. These results are consistent with a dissociative transition state both on and off the enzyme. However, a more inverse nonbridging oxygen KIE is observed for AP and is consistent with active site metal coordination in the transition state (29). In solution, acid-catalyzed hydrolysis of nucleoside glycosidic bonds occurs via dissociative transition states with considerable oxycarbonium ion character in the ribose ring (31, 32). Human and malarial purine nucleoside phosphorylases proceed through similarly dissociative transition states (33); however, more associative transition states are observed for NAD<sup>+</sup> hydrolysis by both diphtheria toxin and human thymidine phosphorylase (34, 35). Nonenzymatic peptide bond formation occurs with equilibrium deprotonation and nucleophilic attack of a neutral amine and loss of a second proton from nitrogen on breakdown of a zwitterionic intermediate (36). Recent KIE and computational results by Hiller et al. (37) suggest that in the ribosome active site, the second deprotonation occurs concomitant with C-N bond formation. These examples illustrate that interactions with enzyme active site residues can subtly alter the transition state in a manner reflecting, in part, the nature of catalytic modes that are used.

In the present study, coordinated theory and experiment provide a chemically detailed comparison of transition state structures for RNA 2'-O-transphosphorylation in solution and in the active site of RNase A. The results reveal that both reactions can proceed via late anionic transition states with complete loss of the 2'O proton and advanced 2'O-P bonding. Nonetheless, the active site environment provides a combination of interactions (general acid and electrostatic catalysis), and thereby influences transition state structure. Electrostatic interactions with positively charged amino acid site chains (His12/Lys41), together with proton transfer from His119, render departure of the 5'O less advanced compared with the solution reaction and stabilize charge buildup in the transition state. Expulsion of the 5'O remains an integral feature of the rate-limiting step both on and off the enzyme, reflecting principles that are fundamental to RNA 2'-O-transphosphorylation reactions. In addition to mechanistic insights, the transition state structure reported here could aid inhibitor design through the construction of stable molecules that mimic its charge distribution. The ability to obtain a chemically detailed description of 2'-O-transphosphorylation transition states paves the way for a new frontier in understanding biological catalysis: defining how active site environments of phosphoryl transferases and substrate reactivity influence transition state structure.

## Methods

**Determination of** <sup>18</sup>**O KIEs on RNA** 2'-**O**-**Transphosphorylation Reactions.** The reaction kinetics and KIEs on the <sup>18</sup> $k_{\text{NUC}}$ , <sup>18</sup> $k_{\text{LG}}$ , and <sup>18</sup> $k_{\text{NPO}}$  for UpG to form 2',3'-cyclic uridine monophosphate and guanosine were analyzed essentially as described by Harris et al. (6) (*SI Text* and Fig. S2B). Briefly, KIEs were determined by analyzing the change in the <sup>16</sup>O/<sup>18</sup>O ratio of an RNA substrate of defined sequence that has been site-specifically enriched with <sup>18</sup>O at the 2'O, 5'O (6, 20, 38), or NPO position as a function of reaction progress. The KIE values determined in Fig. 2 and reported in Table 1 were calculated from plots of ln(<sup>18</sup>O/<sup>16</sup>O) vs. *F* by fitting to ln(<sup>18</sup>O/<sup>16</sup>O) = (1/<sup>18</sup>k - 1)ln(1 – *F*) + ln(*R*<sub>0</sub>), where <sup>18</sup>k is the isotope effect, *F* is the fraction of substrate consumed as determined from integration of HPLC chromatograms, and *R*<sub>0</sub> is the initial <sup>18</sup>O/<sup>16</sup>O ratio in the unreacted substrate. Fitting errors and comparison of fits of the data to the equation, above, to simulations at different isotope effects at the same *R*<sub>0</sub> in Fig. 2 show the precision to be in the range of 0.2–0.5%.

MD Simulation and QM Calculation Methods. Isothermal-isobaric (300 K, 1 atm) MD simulations of RNase A bound to a dinucleotide cytidylyl-3'-adenosine [5'-CpA-3' (CpA)] having the 3',5'-phosphodiester linkage replaced by a pentacovalent phosphorane 2'-O-transphosphorylation transition-state mimic (Fig. 3) were performed for 50 ns, starting from the X-ray crystal structure complexed with cytidilyl-3'-5'-adenosine (Protein Data Bank ID code 1RPG) (39). His12, His119, and Lys41 were in their protonated (+1) states. The assisted model building with energy refinement (AMBER) FF10 force field (40) was used to describe the RNase A protein, along with a set of specific force-field parameters to mimic the late transition state of the dinucleotide (41) and the TIP4PEw model for water (42). The software not (just) another molecular dynamics program (NAMD) was used for all MD simulations (43). For the QM calculations on the nonenzymatic and enzymatic models (Fig. 4), the structures of reactants and the transition states in solution were obtained through full geometry optimization using densityfunctional theory with solvation effects included through the polarizable continuum model (44) and with specialized atomic radii and smooth analytical gradients for treating the solvent effects (45-48). The hybrid B3LYP exchange-correlation functional was used together with the 6-31++G(d,p) basis set. Vibrational frequency analyses were performed to confirm the nature of all stationary points (minima and transition states). KIEs were then calculated from the Bigeleisen equation (14) at 37 °C based on the density functional theory (DFT) potential energy surface. The software package Gaussian 09 was used for all QM calculations (49).

ACKNOWLEDGMENTS. We gratefully acknowledge the encouragement, advice, and seminal contributions of Dr. W. W. Cleland. We thank Dr. Dan Herschlag for providing advice on mechanistic interpretations and comments on the manuscript. This work was supported by the Case Western Reserve University Center for Proteomics and Bioinformatics Center and by National Institutes of Health (NIH) Grant GM096000 (to M.E.H.), NIH Grant AI081987 (to J.A.P.), Hong Kong Baptist University (HKBU) startup and Faculty Research Grant funds (38-40-088 and 40-49-495) to K.-Y.W.,

- Jencks WP (1969) Catalysis in Chemistry and Enzymology, McGraw–Hill Series in Advanced Chemistry (McGraw–Hill, New York).
- Cleland WW, Hengge AC (1995) Mechanisms of phosphoryl and acyl transfer. FASEB J 9(15):1585–1594.
- Lassila JK, Zalatan JG, Herschlag D (2011) Biological phosphoryl-transfer reactions: Understanding mechanism and catalysis. Annu Rev Biochem 80:669–702.
- Perreault DM, Anslyn EV (1997) Unifying the current data on the mechanism of cleavage—Transesterification of RNA. Angew Chem Int Ed Engl 36(5):432–450.
- Oivanen M, Kuusela S, Lönnberg H (1998) Kinetics and mechanism for the cleavage and isomerization of phosphodiester bonds of RNA by Bronsted acids and bases. *Chem Rev* 98(3):961–990.
- Harris ME, et al. (2010) Kinetic isotope effects for RNA cleavage by 2'-O- transphosphorylation: Nucleophilic activation by specific base. J Am Chem Soc 132(33): 11613–11621.
- 7. Raines RT (1998) Ribonuclease A. Chem Rev 98(3):1045-1066.
- Elsässer B, Valiev M, Weare JH (2009) A dianionic phosphorane intermediate and transition states in an associative A(N)+D(N) mechanism for the ribonucleaseA hydrolysis reaction. J Am Chem Soc 131(11):3869–3871.
- Formoso E, Matxain JM, Lopez X, York DM (2010) Molecular dynamics simulation of bovine pancreatic ribonuclease A-CpA and transition state-like complexes. J Phys Chem B 114(21):7371–7382.
- Herschlag D (1994) Ribonuclease revisited: Catalysis via the classical general acid-base mechanism or a triester-like mechanism? J Am Chem Soc 116(26):11631–11635.
- 11. Jackson DY, et al. (1994) A designed peptide ligase for total synthesis of ribonuclease A with unnatural catalytic residues. *Science* 266(5183):243–247.
- 12. Cleland WW (2005) The use of isotope effects to determine enzyme mechanisms. Arch Biochem Biophys 433(1):2–12.
- Hengge AC (2002) Isotope effects in the study of phosphoryl and sulfuryl transfer reactions. Acc Chem Res 35(2):105–112.
- 14. Bigeleisen J, Wolfsberg M (1958) Theoretical and experimental aspects of isotope effects in chemical kinetics. Adv Chem Phys 1(1):15–76.
- O'Leary MH (1977) Studies of enzyme reaction mechanisms by means of heavy atom isotope effects. Isotope Effects on Enzyme Catalyzed Reactions, eds Cleland WW, O'Leary MH, Northrop DB (University Park Press, Baltimore), pp 233–251.
- Sowa GA, Hengge AC, Cleland WW (1997) 18O isotope effects support a concerted mechanism for ribonuclease A. J Am Chem Soc 119:2319–2320.
- Humphry T, et al. (2008) Altered transition state for the reaction of an RNA model catalyzed by a dinuclear zinc(II) catalyst. J Am Chem Soc 130(52):17858–17866.
- Humphry T, Forconi M, Williams NH, Hengge AC (2004) Altered mechanisms of reactions of phosphate esters bridging a dinuclear metal center. J Am Chem Soc 126 (38):11864–11869.
- Cassano AG, Anderson VE, Harris ME (2002) Evidence for direct attack by hydroxide in phosphodiester hydrolysis. J Am Chem Soc 124(37):10964–10965.
- Wong KY, et al. (2012) Characterization of the reaction path and transition states for RNA transphosphorylation models from theory and experiment. Angew Chem Int Ed Engl 51(3):647–651.
- Ye JD, Li NS, Dai Q, Piccirilli JA (2007) The mechanism of RNA strand scission: An experimental measure of the Brønsted coefficient, beta nuc. Angew Chem Int Ed Engl 46(20):3714–3717.
- 22. Gerratana B, Sowa GA, Cleland WW (2000) Characterization of the transition state structures and mechanisms for the isomerization and cleavage reactions of 3'-mnitrobenzyl phosphate. J Am Chem Soc 122(51):12615–12621.
- Lonnberg H, Stromberg R, Williams A (2004) Compelling evidence for a stepwise mechanism of the alkaline cyclisation of uridine 3'-phosphate esters. Org Biomol Chem 2(15):2165–2167.
- 24. Cleland WW (1990) Secondary 18O isotope effects as a tool for studying reactions of phosphate mono-, di-, and triesters. FASEB J 4(11):2899–2905.
- Thompson JE, et al. (1995) Limits to Catalysis by Ribonuclease A. *Bioorg Chem* 23(4): 471–481.

and NIH Grant GM064288 (to D.M.Y.). D.L.K. was supported by NIH Training Grant T32-GM008056. This work was partially supported by the computing resources of the Minnesota Supercomputing Institute, by the High Performance Cluster Computing Centre and the Information Technology Office at HKBU (sciblade and jiraiya), and by the National Science Foundation through TeraGrid resources provided by Ranger, Texas Advanced Computing Center, and Kraken at the National Institute for Computational Sciences under Grant TG-CHE100072.

- Quirk DJ, Park C, Thompson JE, Raines RT (1998) His...Asp catalytic dyad of ribonuclease A: Conformational stability of the wild-type, D121N, D121A, and H119A enzymes. *Biochemistry* 37(51):17958–17964.
- 27. Radak BK, Harris ME, York DM (2013) Molecular simulations of RNA 2'-O-transesterification reaction models in solution. J Phys Chem B 117(1):94–103.
- Rishavy MA, Cleland WW (1999) 13C, 15N and 18O equilibrium isotope effects and fractionation factors. Can J Chem 77(5-6):967–977.
- Zalatan JG, et al. (2007) Kinetic isotope effects for alkaline phosphatase reactions: Implications for the role of active-site metal ions in catalysis. J Am Chem Soc 129(31): 9789–9798.
- Catrina I, et al. (2007) Probing the origin of the compromised catalysis of E. coli alkaline phosphatase in its promiscuous sulfatase reaction. J Am Chem Soc 129(17): 5760–5765.
- Mentch F, Parkin DW, Schramm VL (1987) Transition-state structures for N-glycoside hydrolysis of AMP by acid and by AMP nucleosidase in the presence and absence of allosteric activator. *Biochemistry* 26(3):921–930.
- Berti PJ, Schramm VL (1997) Transition State Structure of the Solvolytic Hydrolysis of NAD<sup>+</sup>. J Am Chem Soc 119(50):12069–12078.
- Taylor Ringia EA, Schramm VL (2005) Transition states and inhibitors of the purine nucleoside phosphorylase family. Curr Top Med Chem 5(13):1237–1258.
- Berti PJ, Blanke SR, Schramm VL (1997) Transition State Structure for the Hydrolysis of NAD Catalyzed by Diphtheria Toxin. J Am Chem Soc 119(50):12079–12088.
- Schwartz PA, Vetticatt MJ, Schramm VL (2010) Transition state analysis of thymidine hydrolysis by human thymidine phosphorylase. J Am Chem Soc 132(38):13425–13433.
- Leung EK, Suslov N, Tuttle N, Sengupta R, Piccirilli JA (2011) The mechanism of peptidyl transfer catalysis by the ribosome. *Annu Rev Biochem* 80:527–555.
- Hiller DA, Singh V, Zhong M, Strobel SA (2011) A two-step chemical mechanism for ribosome-catalysed peptide bond formation. *Nature* 476(7359):236–239.
- Dai Q, Frederiksen JK, Anderson VE, Harris ME, Piccirilli JA (2008) Efficient synthesis of [2'-180]uridine and its incorporation into oligonucleotides: A new tool for mechanistic study of nucleotidyl transfer reactions by isotope effect analysis. J Org Chem 73 (1):309–311.
- Zegers I, et al. (1994) The structures of RNase A complexed with 3'-CMP and d(CpA): Active site conformation and conserved water molecules. Protein Sci 3(12):2322–2339.
- Hornak V, et al. (2006) Comparison of multiple Amber force fields and development of improved protein backbone parameters. *Proteins* 65(3):712–725.
- Lee TS, Giambaşu G, Harris ME, York DM (2011) Characterization of the Structure and Dynamics of the HDV Ribozyme at Different Stages Along the Reaction Path. J Phys Chem Lett 2(20):2538–2543.
- Horn HW, et al. (2004) Development of an improved four-site water model for biomolecular simulations: TIP4P-Ew. J Chem Phys 120(20):9665–9678.
- Phillips JC, et al. (2005) Scalable molecular dynamics with NAMD. J Comput Chem 26 (16):1781–1802.
- Cossi M, Rega N, Scalmani G, Barone V (2003) Energies, structures, and electronic properties of molecules in solution with the C-PCM solvation model. J Comput Chem 24(6):669–681.
- Khandogin J, Gregersen BA, Thiel W, York DM (2005) Smooth solvation method for dorbital semiempirical calculations of biological reactions. 1. Implementation. J Phys Chem B 109(19):9799–9809.
- Scalmani G, Frisch MJ (2010) Continuous surface charge polarizable continuum models of solvation. I. General formalism. J Chem Phys 132(11):114110.
- York DM, Karplus M (1999) A smooth solvation potential based on the conductor-like screening model. J Phys Chem A 103(50):11060–11079.
- Gregersen BA, York DM (2005) High-order discretization schemes for biochemical applications of boundary element solvation and variational electrostatic projection methods. J Chem Phys 122(19):194110.
- Frisch MJ, et al. (2009) Gaussian 09, Revision B.01 (Wallingford, CT). Available at www. gaussian.com. Accessed July 9, 2013.