Integration of kinetic isotope effect analyses to elucidate ribonuclease mechanism☆

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Abstract
The well-studied mechanism of ribonuclease A is believed to involve concerted general acid–base catalysis by two histidine residues, His12 and His119. The basic features of this mechanism are often cited to explain rate enhancement by both protein and RNA enzymes that catalyze RNA 2′-O-transphosphorylation. Recent kinetic isotope effect analyses and computational studies are providing a more chemically detailed description of the mechanism of RNase A and the rate limiting transition state. Overall, the results support an asynchronous mechanism for both solution and ribonuclease catalyzed reactions in which breakdown of a transient dianionic phosphorane intermediate by 5′-O bond cleavage is rate limiting. Relative to non-enzymatic reactions catalyzed by specific base, a smaller KIE on the 5′-O leaving group and a less negative δc are observed for RNase A catalysis. Quantum mechanical calculations consistent with these data support a model in which electrostatic and H-bonding interactions with the non-bridging oxygens and proton transfer from His119 render departure of the 5′-O less advanced and stabilize charge buildup in the transition state. Both experiment and computation indicate advanced 2′-O–P bond formation in the rate limiting transition state. However, this feature makes it difficult to resolve the chemical steps involved in 2′-O activation. Thus, modeling the transition state for RNase A catalysis underscores those elements of its chemical mechanism that are well resolved, as well as highlighting those where ambiguity remains. This article is part of a Special Issue entitled: Enzyme Transition States from Theory and Experiment.

1. Introduction
Ribonuclease A has long served as an experimental system for exploring the fundamental principles that describe how enzymes achieve catalysis [1]. Extensive structural and biochemical studies have resulted in a model for its mechanism that is widely cited as a foundational example of biological catalysis. In this model, RNase A catalysis of RNA 2′-O-transphosphorylation uses a general acid catalytic mode involving donation of a proton to the 5′-O leaving group by a protonated His119. Activation of the 2′O nucleophile is attributed to His12, which is proposed to act in its neutral form as a general base (Fig. 1). Parallels to this concerted acid/base mechanism have been drawn to many different phosphoryl transfer enzymes including ribozymes [2–4].

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principles of biological catalysis and suggest strategies for engineering novel catalysts. Also, areas of ambiguity identified in the process help to identify limitations to our current understanding of enzyme mechanism.

RNase A is good system to address questions regarding transition state stabilization by enzymes because it is a potent catalyst, providing greater than 10^{10}-fold rate enhancement over the uncatalyzed RNA 2′-O-transphosphorylation reaction at neutral pH [9]. Although spontaneous RNA 2′-O-transphosphorylation is slow, it can be catalyzed by both acid and base. These reactions proceed at experimentally accessible rate constants permitting detailed mechanistic analysis [6,7,10]. Comparison of solution and enzyme mechanisms has permitted a better understanding of the simultaneous application of multiple catalytic modes that defines biological catalysis [1,6–8]. Protein ribonuclease like RNase A contain amino acids in their active sites, such as histidine, glutamine and lysine, capable of acid/base catalysis and electrostatic catalytic modes [8,11]. Similarly, the active sites of small self-cleaving ribozymes contain nucleobases that are proposed to employ general acid/base catalytic modes analogous to His12 and His119 [2,3]. Thus, investigation of the details of RNase A and RNA 2′-O-transphosphorylation can facilitate the comparison of strategies for transition state stabilization across a wide range of biological catalysts.

2. Mechanisms and transition states of solution RNA 2′-O-transphosphorylation

RNA 2′-O-transphosphorylation reactions involve nucleophilic attack of the 2′O on the adjacent phosphonyl group to generate a 2′,3′ cyclic phosphate and a 5′ hydroxyl products (Fig. 2). This reaction is catalyzed by both acid (pathways 1 and 2) and base (pathway 3). At high pH (>9) the rate constant increases log-linear with increasing pH and displays an apparent pK_a at ca. pH 13.4. The observed pH dependence reflects an underlying mechanism involving equilibrium deprotonation of the 2′OH to generate a 2′ oxygen followed by nucleophilic attack [6,10,12]. The rate constant also increases with decreasing pH (pH < 5). The observed pH dependence is believed to reflect ground state protonation of one or more non-bridging oxygens, and reaction via either monoaenic or neutral stepwise reaction channels. Under acidic conditions isomerization products are also observed that result from pseudorotation of singly (I_{monoanion}) or a doubly protonated (I_{neutral}) phosphorane intermediate [6]. The observation that only transphosphorylation products are observed at high pH indicates that the base catalyzed reaction occurs by a concerted mechanism or an asynchronous one in which the diaanion (D_{anion}) phosphorane is too short lived to undergo pseudorotation.

2.1. Linear free energy relationship analyses

Linear free energy relationship (LFER) analyses provide information on mechanism by analyzing the dependence of reaction rate constants on the pH of the nucleophile and leaving group [13]. To relate these values to transition state bonding, the observed β (β = Δlogk/ΔpK_a) is calibrated against the βEQ which is the effect changing pK_a on reaction equilibria. The parameter α = (β_{LC} or β_{NE}) / βEQ is often used to express the fraction of total charge development in the transition state, which is related to the amount of bond breakage or formation [13]. Quantum chemical calculations provide a molecular level characterization of the structure and bonding in the transition state that can aid in the interpretation of experimental LFER analysis [32]. For reactions involving phosphoryl transfer reactions of ribose hydroxyls, the βEQ is not available, however a value of −1.35 is measured for equilibrium ionization of substituted phenols [14] and a value of −1.56 has been estimated [15]. A β_{LC} of −1.28 is measured for the reaction of a series of uridine 3′ alkyl phosphodiester indicating a late transition state with respect to bond cleavage (α ≈ 0.71) [16]. For reaction of uridine 3′ aryl phosphates, however, a β_{LC} of −0.56 was measured for hydroxypropyl aryl phosphate cyclization (α = 0.44) [17]. These results are consistent with less charge accumulation on the leaving group in the transition state and therefore an earlier transition state for diesters with better aryl leaving groups. A

Fig. 1. The concerted general acid/base mechanism for RNase A catalysis. A scheme for active site interactions involving His12, His119 and Lys41 in the ground state, transition state (TSc) and product state is shown from left to right. Note that nucleophilic addition (2′→O bond formation) and leaving group departure (5′→O bond cleavage) occur in a single transition state. Proton transfer involving the nucleophile and leaving group occur concomitantly with changes in O→P bonding. General base catalysis involves transfer of the 2′O proton (shown in red) to the deprotonated form of His12. General acid catalysis occurs via transfer of a proton (shown in blue) from the protonated form of His119 to the departing 5′. The concerted general acid/base mechanism for RNase A catalysis. A scheme for active site interactions involving His12, His119 and Lys41 in the ground state, transition state (TSc) and product state is shown from left to right. Note that nucleophilic addition (2′→O bond formation) and leaving group departure (5′→O bond cleavage) occur in a single transition state. Proton transfer involving the nucleophile and leaving group occur concomitantly with changes in O→P bonding. General base catalysis involves transfer of the 2′O proton (shown in red) to the deprotonated form of His12. General acid catalysis occurs via transfer of a proton (shown in blue) from the protonated form of His119 to the departing 5′.
breakpoint in the relationship between leaving group $pK_a$ and log $k$ obtained by linear extrapolation of the two data sets is observed at $ca. 12.5$, and this feature has been attributed to a change in mechanism from a concerted to stepwise mechanism [16]. A $\beta_{\text{NUC}}$ of 0.75 for solution RNA transphosphorylation has been measured using a series of 2$'$ substituted analogs [15]. For comparison, the $\beta_{\text{NUC}}$ values measured for oxygen and nitrogen nucleophiles attacking monoesters are in the range of 0.1–0.2 [18]. This result suggests that the 2$'$O nucleophile has a relatively smaller effective charge in the rate limiting transition state for reactions of alkyl phosphate diesters.

2.2. Kinetic isotope effects

The substitution of a stable heavy isotope for a reacting atom can affect both the rate and equilibrium constants for the reaction. Kinetic isotope effects (KIEs) arise primarily due to differences in bond vibrational properties between the ground state and transition state [19]. Differences in the observed KIEs for transphosphorylation reactions with good (aryl) and poor (alkyl) leaving groups also indicate that there are significant differences in rate limiting transition states for the two classes of substrates. Computational studies reveal that these different transition states have distinct kinetic isotope signatures [33].

The transphosphorylation reaction of hydroxypropyl-p-nitrophenol phosphate shows a nucleophile KIE that is normal (1.0327) while the nucleophile KIE measured for RNA 2$'$O-transphosphorylation is inverse (0.995) [12,20]. The fact that these values are so different suggests immediately that there are significant differences in transition state structure, at least with respect to nucleophile bonding. However, in order to interpret these effects in terms of specific mechanistic differences it is important to understand that observed KIEs reflect O–P bonding as well as all other vibrational modes that change on going from the ground state to the transition state.
Oxygen isotope effects on transphosphorylation can reflect the fractionation due to deprotonation of the hydroxyl group and the kinetic isotope effect on nucleophilic attack or leaving group departure. Fractionation due to deprotonation or protonation can have a large contribution to the observed effect since the equilibrium deprotonation of the 2′-O is estimated to be 1.024 [21]. The full contribution is expected for a mechanism involving equilibrium deprotonation, while a smaller contribution would result from partial transfer in the rate limiting transition state. The former mechanism defines specific base nucleophile activation while the later exemplifies general base catalysis. For example, the observed effect of 0.995 for specific base catalysis cited above changes to 0.984 above pH 13 due to loss of the O−H stretching mode in the ground state [12,22]. The observation of a nucleophile KIE closer to unity at lower pH is interpreted as reflecting a large inverse contribution that is offset by the normal contribution due to the equilibrium isotope effect on 2′-O deprotonation.

The contribution of nucleophile attack to the observed KIE is itself composed of two factors, the temperature-independent factor (TIF) and the temperature-dependent factor (TDF) [23,24]. The TDF is due to differences in bond vibrational modes in the transition state compared to the ground state and reflect differences in zero point vibration- al energies proportional to, and in the direction of equilibrium isotope effects. Inverse nucleophile KIEs are observed for reactions involving oxygen which hydrogen is replaced by a larger chemical group such as carbon introducing new vibrational modes [25,26]. Formation of the O−P bond, therefore, favors the heavier isotope and results in an inverse contribution to the KIE. The TIF reflects the extent to which the labeled atom participates in reaction coordinate motion. While all other modes are at their minima in the transition state, this motion is uniquely and by definition at a maxima at the transition state. As a consequence the TIF will always favor 18O resulting in a normal contribution to the observed KIE proportional to the extent that its motion is involved in the reaction coordinate.

Thus, observed nucleophile KIEs are normal for early transition states and become less normal as O−P bonding in the transition state increases, eventually becoming inverse as nucleophile bonding becomes more complete [27–30]. In this light, the normal (1.023) nucleophile KIE observed for the transphosphorylation reaction of hydroxypyropyl-p-nitrophosphate reflects an early transition state for a substrate with an aryl leaving group. Inverse KIEs reflect mechanisms in which the nucleophile participates very little in reaction coordinate motion, and the TDF dominates the observed KIE. The inverse KIE measured for the RNA cleavage reaction (0.984) therefore can be interpreted as representing advanced O−P bond formation and in a stepwise mechanism essentially becomes the EIE for formation of a dianionic phosphorane intermediate [21,22].

The leaving group KIEs for transphosphorylation reactions with good (nitrophenol) and poor (ribose) leaving groups are also very different consistent with a change in rate limiting transition state. The leaving group KIE for transphosphorylation of hydroxypyropyl-p-nitrophosphate is small (1.005) while a much larger leaving group KIE (1.034) is measured for RNA strand cleavage. The EIE on deprotonation of the alcohol of malate is reported to be 1.032 and the observed KIE proportional to the extent that its motion is involved in the reaction coordinate. For both stepwise mechanisms and concerted asynchronous mechanisms, the transition states can be further designated as either “early” or “late”, depending on the location of the transition state along a reaction coordinate that involves the difference in the leaving group (R₁) and nucleophile (R₂) distances with the reactive phosphorus.

2.3. A general framework for solution transphosphorylation reactions

Computational studies of nucleophilic attack on diesters including cyclization reactions have consistently illustrated the potential for stepwise mechanism for RNA transphosphorylation with a transient monoanionic phosphorane intermediate [32–36]. Recently, density-functional calculations [33] were used to analyze a series of transphosphorylation models with different leaving groups to provide a direct connection between observed Brønsted coefficients and KIEs with the structure and bonding in the transition state (Fig. 3).

In this framework, diesters with sufficiently reactive leaving groups react via a concerted mechanism that proceeds through a single early transition state, and LFER analysis predicts a βLG value with a small magnitude in correspondence with experimental results. This transition state is characterized by normal nucleophile KIE and a small leaving group KIE as reported for base catalysis of hydroxypyropyl-m-
nitrobenzyl phosphate. For poor leaving groups like the 5′O of RNA, the computationally observed mechanism is stepwise as reported previously. For this class of reactions the rate-controlling transition state is late, leading to a large negative $\beta_{LC}$ value [33]. The KIEs for this mechanism are in line with the inverse nucleophile and large normal leaving group KIEs observed for specific base catalyzed RNA transphosphorylation described, above. The resulting computational framework can assist in identifying and predicting KIEs that provide insight into mechanism and provide a benchmark for interpreting the effects of enzymes such as RNase A.

3. Evidence for acid/base catalytic modes in the active site of RNase A

3.1. Roles of key active site residues

Structures of complexes of RNase A together with cyclic nucleotides, substrate analogs and a transition state analog have played a large role in defining potential active site interactions important for catalysis. His12 and His119 are proximal to the scissile phosphorib group in enzyme-transition state analog uridine vanadate complexes [37,38] and complexes with inactive substrate oligonucleotides, e.g. [39–41]. His119 is consistently near the 5′O leaving group, or a non-bridging phosphoryl oxygen in product complexes. The interactions of His12 are more variable and it interacts with the 2′ position in the 2′ fluorescence substrate and vanadate complexes. However, it interacts with a non-bridging oxygen in the 2′-deoxy substrate analogs. In structures of RNase A with product or deoxy modified substrates, Lys41 is positioned to interact with either the 2′O or a non-bridging oxygen.

Substitution of either His12 or His119 with alanine decreases the value of $k_{cat}/K_m$ for cleavage of a UpA dinucleotide substrate by more than 104-fold [42]. The general acid/base mechanism is predicted to underlie the observed bell-shaped pH-$k_{cat}/K_m$ profile. Association of polynucleotide substrates is rate limiting for $k_{cat}/K_m$ and these kinetics distort the observed kinetic $pK_a$ values from their intrinsic values [43]. However, for cyclic nucleotide and dinucleotide substrates for which the first irreversible step is not binding, but catalysis or product release, the observed reaction $pK_a$ values are near 6 [44–46] consistent with the microscopic $pK_a$ values of the active-site histidine residues determined by NMR spectroscopy [47,48]. An RNase A variant containing a 4-fluorohistidine, which has a $pK_a$ of 4.5, at both His12 and His119 retains pH dependence that is still bell-shaped, but shifted to lower $pK_a$ consistent with both 4-fluorohistidine residues participating in catalysis [49]. However, there is a relatively small decrease in $k_{cat}$ for the modified enzyme, which suggested that the proton transfer steps may not be rate limiting when the enzyme is in the correct protonation state. Subsequent detailed analyses of RNase A kinetics and structural dynamics show that product release coupled with protein loop motion is rate limiting for $k_{cat}$ [50,51].

Mutations of Lys41 affect the reaction rates for RNA, cyclic nucleotide and uridine-3′-p-nitrophenol substrates equivalently, arguing for a common role in stabilizing the transition states for these substrates. Although the solution TS for aryl substrates is different than RNA and cyclic nucleotides, each proceeds by an anionic transition state. The location of Lys41 adjacent to the reactive phosphorib group is consistent with a role in electrostatic stabilization. Based on mutagenesis and modification data, however, the contribution of Lys41 has been argued to donate a hydrogen bond to the transition state during catalysis rather than simply stabilize negative charge [52].

3.2. Mechanistic information from linear free energy relationship analysis

In order to fully appreciate the roles of Lys41, His12 and His119 in transition state stabilization it is necessary to identify the rate limiting transition state for RNase A catalysis and understand its structure. Davis et al. analyzed the dependence of $k_{cat}/K_m$ for RNase A catalyzed reactions of uridine-3′-arylyphosphate esters on leaving group $pK_a$ and obtained a $\beta_{LC}$ value of ca. $–0.2$ [53]. As indicated, above, LFER analysis of uridine-3′-arylyphosphate reactions catalyzed by specific base has a more negative $\beta_{LC}$ value ($–0.52$) indicating less development of negative charge on the leaving group in the reaction catalyzed by RNase A.

Mutations of His119 has a relatively small effect on the rate constant for transphosphorylation of the aryl diester uridine-3′-p-nitrophenol, while it decreases the rate constant for cleavage of RNA substrates by 103-fold. Because the $pK_a$ of the nitrophenol is much lower than ribose ($ca. 7 \text{ versus} 14$), it is argued that for the uridine-3′-p-nitrophenol substrate, the function of His119 as a general acid is no longer required [8,9] and therefore its mutation has little effect. As detailed above, aryl phosphate reactions appear to proceed by a different mechanism relative to RNA in solution, proceeding by an early transition state with little P-O bond cleavage. If His119 does not interact with this transition state, then it follows that the observation of a lower $\beta_{LC}$ value for RNase A reflects an overall change in transition state structure, rather than the influence of general acid catalysis. A much larger $\beta_{LC}$ value of $–1.28$ is observed for non-enzymatic reactions of alkyl diesters such as RNA as described, above [54]. However, the $\beta_{LC}$ for RNase A reaction for substrates in this $pK_a$ range has not been reported, but such analyses could help resolve the extent to which His119 influences the accumulation of negative charge on the 5′O leaving group.

Corresponding analyses of $\beta_{nuc}$ for RNase A and the potential role of His12 in altering this parameter relative to solution reactions are not available. A 2′-deoxy-2′-thio-UpA substrate was reported to bind the active site of RNase A, but is apparently not a substrate [8]. The rate of thiolate attack on the adjacent phosphodiester bond is estimated to be 107-fold slower than that of the corresponding alkoxide, although at neutral pH there is a much higher concentration of the thiolate, due to its lower $pK_a$ ($–8 \text{ versus} –13$) resulting in only a 25-fold slower observed rate constant for the solution reaction [55]. The inactivity of RNase A toward this substrate is not understood, but could be an issue related to reaction geometry, or due to stable non-catalytic interactions that form due to the presence of a full negative charge at the 2′ position in the ground state. A $\beta_{nuc}$ of 0.75 for solution RNA transphosphorylation has been measured using a series of 2′ substituted analogs [15]. This result suggests that the 2′O nucleophile has an effective charge of only $–0.25$ in the rate limiting transition state for the spontaneous reaction. A comparative determination of $\beta_{nuc}$ for RNase A is not available, but may be feasible if the requisite analogs do not otherwise interfere with enzyme function.

3.3. Transition state by analysis using heavy atom ($^{18}O$) isotope effects

In principle, measurement of heavy atom KIEs on the RNase A catalyzed RNA 2′-O-transphosphorylation can provide a greater extent of mechanistic clarity. This optimism is based in part on the fact that KIEs involve the smallest possible perturbation of substrate structure that can be used to determine reaction mechanism. Moreover, the work of W. W. Cleland and colleagues on KIEs of phosphoryl transfer reactions including RNase A catalysis provides a firm foundation for interpreting mechanistic detail [56–59]. These classic studies involved application of a remote label method in which the heavy oxygen isotope was monitored indirectly using a $^{15}N$ label in the nitrophenol leaving group. The required isotope ratio measurements were made to high precision using an isotope ratio mass spectrometer. They further measured RNase A catalyzed transphosphorylation of uridine-m-nitrobenzyl phosphate (UmNB) substrate by the same remote label method. The $pK_a$ of the m-nitrobenzyl alcohol leaving group is close to that of a natural substrate nucleotide ribose 5′O leaving group and is expected to react with a similar transition state [59].

The results obtained for RNase A transphosphorylation of uridine-m-nitrobenzyl phosphate (UmNB) by Cleland and colleagues [59] are shown in Fig. 4. These values are compared to KIEs determined for RNA 2′-O-transphosphorylation reactions using whole molecule mass spectrometry by Gu et al., 2013. In this alternative approach the isotope...
The observed and calculated KIEs for the 2′O nucleophile (18KNUC), 5′O leaving group (18KLG), and non-bridging phosphoryl oxygens (18KPO) are shown adjacent to the corresponding atoms in the transition state models. KIE values obtained for RNA and for uridine-m-nitrobenzylphosphate (UmNB) are labeled accordingly. Note that the computational results were obtained using a simplified ribose-3′-etholphosphate model.

KIE values agree best for a model involving H-bonding between the active site and non-bridging oxygen atoms along with proton transfer from His119 to the 5′O as in the classic mechanistic model [59] [18KNUC (0.998/0.994), 18KLG (1.026/1.014), and 18KPO (1.006/1.001) (calculated/measured)] (Fig. 4). The calculated 5′O isotope effect for the base-catalyzed reaction simulation (1.048) is larger than the value 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) 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.

Systematic and detailed QM/MM studies of RNase A mechanism have also been reported by Weare and colleagues [62]. This investigation also supports a stepwise mechanism although with two key features that differ from the model described, above [Fig. 5]. In the alternative mechanism, the transition state for nucleophilic addition (TS1) is observed to be rate limiting. However, the observation of inverse nucleophile and normal leaving group KIEs similar to solution reactions unambiguously supports a late, product-like transition state dominated by leaving group departure (TS2). The second difference regards the structure of the phosphorane intermediate. In the alternative mechanism, the proton initially coordinating the O2′ migrates first to His12 and then to the nonbridging O1P resulting in a triester mechanism with a monoanionic intermediate. However, simple interpretation of weak thio effects on RNase A and the observation of a small nonbridging oxygen isotope effect is that such triester mechanism is unlikely [63].

4. Conclusions and future directions

Experiment and computation together indicate that RNase A stabilizes a product-like transition state that is very similar to the anionic solution mechanism catalyzed by specific base in solution [12,34,36,60]. Nonetheless, the enzyme significantly alters leaving group bonding attributable to general acid catalysis from His119. The overall asynchrony of the mechanism that is inherent in both the computational and experimental perspectives, however, complicates experimentally evaluating modes of nucleophilic activation. Similarly, the specific catalytic interactions between the active site and the non-bridging oxygens remain difficult to resolve with precision.

Because the interpretation of LFER and KIE is necessarily grounded in transition state theory, assumptions or interpretations of available
mechanistic data must be made in order to frame the results in terms of overall transition state structure [64]. The current model framework for RNA transphosphorylation predicts the solution mechanism to be stepwise and that the transition state for leaving group departure (TS2) is rate limiting. It is worth noting that Cleland and Cook demonstrated using adenosine and cytosine deaminases that if proton transfer and changes in $^{18}$O bonding occur in the same step, then the $^{18}$O KIEs will be the same in both H$_2$O and D$_2$O [65]. For a mechanism in which rate constants for both formation and breakdown of an intermediate are kinetically significant, then the observed heavy atom KIE will be decreased in proportion to the change in proton fractionation. For phosphate groups the equilibrium deuterium isotope effect on protonation is large (3–5) [26], and thus differences in the observed nucleophile and leaving group KIEs may be detectable such an experiment could provide an additional incisive test of concerted/stepwise mechanism.

As described above, the asynchronous mechanism for solution and RNase A reactions makes it difficult to experimentally interrogate the chemical details of nucleophilic activation because this step is not rate limiting. An attractive strategy, given the emerging framework for solution reactions, would be to adjust the reactivity of the leaving group to make nucleophilic attack rate-limiting. In such a mechanism a normal nucleophile KIE would be predicted, but its value would be reduced by the degree of loss of the 2$'_{-}$O–H bond. In general, the normal contribution to the nucleophile KIE from deprotonation is diminished by the degree of O–P bonding in the transition state. Therefore, in a general base mechanism in which deprotonation and nucleophilic attack occur in the same step, a more inverse KIE is expected. Complete loss of the 2$'_{-}$O–H stretching mode prior to nucleophilic attack in a specific base mechanism results in a large normal contribution to the observed nucleophile KIE of ca. 1.027 based on observed effects on phenols and quantum calculations. Mutation of His12 would be predicted to result in a change to a specific base mechanism with a concomitant increase in the observed nucleophile KIE.

In sum, KIE analyses of RNA transphosphorylation have significant power to distinguish between alternative mechanisms, however, they cannot always be interpreted simply. Thus, additional benchmarks must be sought and integration of theory and experiment continued.

Fig. 5. Stepwise model for RNase A catalysis. An alternative to the concerted general acid/base model consistent with experimental data and indicated by computation involves transfer of the 2$'$O proton (red) to His12 followed by formation of the 2$'$O–P bond. The resulting intermediate is shown as a dianion in which an H-bond between His12 and the substrate phosphoryl group is maintained. However, computational studies have also suggested the potential for formation of a protonated phosphorane in which this proton resides on the phosphoryl oxygen [62]. The rate limiting transition state involves departure of the 5$'$O leaving group and transfer of a proton from His119 (blue).

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