

Coupling of the Guanosine Glycosidic Bond Conformation and the Ribonucleotide Cleavage Reaction: Implications for Barnase Catalysis

Short Title: GpA hydrolysis mechanism

Key words: Potential of mean force calculations; dinucleotides; conformational analysis; RNA-protein interactions; nucleophilic attack

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ABSTRACT

To examine the possible relationship of guanine-dependent GpA conformations with ribonucleotide cleavage, two potential of mean force (PMF) calculations were performed in aqueous solution. In the first calculation, the guanosine glycosidic ($G\chi$) angle was used as the reaction coordinate, and computations were performed on two GpA ionic species: protonated (neutral) or deprotonated (negatively charged) guanosine ribose O_2' . Similar energetic profiles were obtained for both ionic forms, with two minima (*anti* and *syn* $G\chi$). In both simulations the *anti* conformation was more stable than the *syn*, and barriers of ~ 4 kcal/mol for the *anti* \rightarrow *syn* transition were obtained. Structural analysis showed a remarkable sensitivity of the phosphate moiety to $G\chi$ rotation, suggesting a possible connection between $G\chi$ orientation and the mechanism of ribonucleotide cleavage. This hypothesis was confirmed by the second PMF calculations, for which the O_2' -P distance for the deprotonated GpA was used as reaction coordinate. The computations were performed from two selected starting points: the *anti* and *syn* minima determined in the first PMF study of the deprotonated guanosine ribose O_2' . The simulations revealed that the O_2' attack along the *syn* $G\chi$ was more favorable than that along the *anti* $G\chi$: energetically, significantly lower barriers were obtained in the *syn* than in the *anti* conformation for the O-P bond formation; structurally, a lesser O_2' -P initial distance and a better suited orientation for an in-line attack was observed in the *syn* relative to the *anti* conformation. These results are consistent with the barnase-ribonucleotide catalytic interaction, for which a guanine *syn* conformation of the substrate is required to allow the abstraction of the ribose H_2' proton by the general base Glu73, thereby suggesting a coupling between reactive substrate conformation and enzyme structure and mechanism.

INTRODUCTION

Barnase, an endoribonuclease from *Bacillus amyloliquefaciens*, catalyzes the cleavage of single-stranded RNA. The proposed catalytic mechanism¹ is that of a general acid-base, with Glu73 and His102 as the general base and general acid, respectively. The mechanism involves a trans-esterification reaction yielding a 2',3'-cyclic phosphate intermediate. This process requires the abstraction of the ribose H_{2'} proton by Glu73, the formation of a covalent bond between O_{2'} and the phosphorus atom, and a proton transfer from His102 to O_{5'} (see Figure 1 for mechanism details and Figure 2 for the nucleotide nomenclature).¹

Barnase cleaves ribonucleotides after guanosine either specifically or preferentially for dinucleotides and longer substrates, respectively.^{2,3} Remarkably, the catalytic efficiency depends on the length of the nucleotide. It has been shown² that lengthening the substrate from GpX to GpXp, where X represents any nucleoside and G guanosine, produces a 10-fold decrease in K_m and a 100-fold increase in k_{cat}. Furthermore, the orientation of the guanine base relative to the ribose, governed by the guanine glycosyl torsion angle (Gχ), is key in barnase catalysis⁴ (see Figure 2 for torsion angle nomenclature). The rotation of the χ angle is sterically restricted and two conformational states, *syn* and *anti*, are preferred.⁵ In the *anti* conformation the base points away from the sugar moiety whereas in the *syn* conformer the base is positioned over the sugar ring. On the basis of the χ angle distribution found for mononucleoside and mononucleotide crystal structures⁶ and on potential energy calculations⁷ on deoxyribonucleosides, the *anti* and *syn* conformations were defined⁸ as those where χ adopts values ranging between 170-300° and 30-90°, respectively. It has been postulated that Gχ must adopt a *syn* conformation for barnase catalysis to take place.⁴ However, in the [barnase-d(CpGpApC)]₂ crystal structure⁴ (RCSB-PDB code 1BRN), the value of

the $G\chi$ angle is 215° (*anti* region), leading to what seems to be an unproductive conformation, where the 2'-OH, which was modeled into this structure, is pointing away (~ 7 Å separation) from Glu73 side chain.⁴ Interestingly, modeling the bound guanosine in the *syn* conformation, followed by energy minimization of the nucleotide⁴ or both the nucleotide and the enzyme,⁹ positioned the 2'-OH group at a hydrogen bonding distance from Glu73 carboxylate. These derived models were shown to be consistent with the crystal structure of the complex of barnase with its natural product (3'-GMP)¹⁰ which displays the catalytically competent *syn* $G\chi$ conformation, allowing a Glu73–O_{2'} hydrogen bond to form. Likewise, *anti* and *syn* $G\chi$ conformations were observed in a nuclear magnetic resonance (NMR) study¹¹ of a barnase-3'-GMP complex in solution. In the *anti* conformation, the O_{2'}-H points away from Glu73, whereas in the *syn* conformation both moieties are within hydrogen bonding distance.

Nanosecond time scale molecular dynamics (MD) simulations of GpA and GpAp in water¹² and when bound to barnase,¹³ were recently performed using as starting conformation the dinucleotide structure from the barnase-CGAC complex.⁴ Results showed that none of the sampled conformations were in the *syn* range ($0-90^\circ$). It was argued,^{12,13} that the complete absence of an *anti* \rightarrow *syn* transition in both GpA and GpAp during these simulations was likely due to high energy barriers that cannot be overcome at room temperature during the considered time scales. The problem of investigating the *anti/syn* equilibrium in oligonucleotides by MD simulations has been pointed out recently.⁸ It was attributed to the requirement of surmounting conformational barriers of the type involved in the conversions between nucleic acid forms (such as B- and Z-DNA), which are not accessible during current MD time scales. This limitation does not apply to nucleosides however. MD simulations⁸ of the eight (ribo- and deoxyribo-) nucleoside constituents of nucleic acids in aqueous solution were

shown to yield relatively frequent transitions between *anti* and *syn* conformations, with riboguanosine ranking second in the number of observed transitions.

The energy barrier associated with the χ angle rotation governing the *anti* to *syn* transition in both deoxyribonucleosides and deoxyribonucleotides has been recently estimated by quantum mechanical (QM) methods in vacuo.⁷ In this study the malleability of this torsion angle and its dependence on the type of base and sugar stereochemistry was made evident. For each deoxyribonucleoside, four energy surfaces were obtained by varying the value of χ under γ (g^+ or *trans*) and sugar pucker (north or south) combinations. For all deoxyribonucleosides, the conformation of lowest energy was $\chi=anti$, with $\gamma=g^+$ and a south (adenine, guanine, thymine) or north (cytosine) sugar pucker. The puckering mode was stable along the χ variation except for the profiles with $\gamma = trans$ and initiated with a south sugar, where the puckering switched to the north range in the vicinity of $\chi=340^\circ$ or $\chi=50^\circ$ for all four nucleosides. High level QM calculations showed the energy barrier for the *anti* \rightarrow *syn* transition to be in the range of 3.0-9.8 kcal/mol, and following the pathway through $\chi=120^\circ$.⁷

In the present paper, we aim at gaining further insight into the $G\chi$ barrier for GpA and investigate its potential relation to the catalytic mechanism of barnase. To this end, potential of mean force (PMF) calculations are carried out for this dinucleotide, in water. One series of calculations is carried out over the $G\chi$ angle in two ionic forms of the substrate, the protonated guanosine ribose form (with $O_2^{\cdot-}H$) and the deprotonated form ($O_2^{\cdot-}$). The second series of PMF calculations are performed over the $O_2^{\cdot-}P$ distance for the deprotonated GpA form, starting from the *anti* and *syn* $G\chi$ conformations determined for this ionic form in the first PMF calculations. The second PMF analysis aims at characterizing the energy barrier for the first step of the proposed barnase reaction mechanism, in which the deprotonated guanosine ribose $O_2^{\cdot-}$ forms a

covalent bond with the phosphorus atom yielding a 2'-3'-cyclic phosphate intermediate (see above and Figures 1 and 2).

In our PMF analysis, both the 2'-OH and 3'-OH groups were included in the calculations considering the importance of the 2'-OH group in barnase catalysis and the fact that the potential energies for ribonucleotides are very sensitive to the relative orientations of these groups.⁸ Our calculations furthermore included the entirety of the adenosine group in order to incorporate key effects due to the flexibility of this group and its interactions with the guanosine and phosphate moieties. Our study thus goes beyond the analysis of the intrinsic energetics of an isolated χ angle, and aims at estimating the energetics of the $G\chi$ rotation, in both GpA and guanosine-deprotonated O₂' GpA, in water, including and drawing especial attention to cooperative effects that the various moieties of the molecule may exert on one another.

COMPUTATIONAL METHODS

Building the System

To study the behavior of GpA in aqueous solution we used the hybrid quantum mechanical and molecular mechanical (QM/MM) methodology¹⁴⁻¹⁶ as implemented in the DYNAMO¹⁷ software. This allowed us to combine the flexibility of QM methods for describing chemical reactions with the computational efficiency of MM methods. Starting configurations were taken from the crystal structure of the [barnase-d(CpGpApC)]₂ complex⁴ (RCSB-PDB code 1BRN). The dinucleotide was solvated by adding a cubic box of 31.4 Å side dimensions, containing 995 water molecules. In each simulation, the QM subsystem comprised the dinucleotide protonated (66 atoms) or deprotonated (65 atoms) that was described by means of the semiempirical AM1 method,¹⁸ while the water molecules were modeled using the TIP3P force field.¹⁹ All molecular dynamics simulations were done at room temperature (300 K) and the Velocity-Verlet-Langevin algorithm was used with a time step of 1 fs. The canonical thermodynamic ensemble (NVT) was used for all calculations and periodic boundary conditions were applied. A switched outer cut-off radius of 12 Å was employed for MM-MM and QM-MM interactions.

PMF Calculations of $G\chi$ Rotation for GpA in Water

The first goal of the present study was to evaluate the free energy barrier for the $G\chi$ rotation in GpA in aqueous solution (see Figure 2 for atom and torsion angle notations). To this end, PMF calculations were performed taking $G\chi$ as the reaction coordinate. To analyze whether the energy profile depends on the protonation state of the guanosine O_{2'} atom, computations were performed for the protonated (O_{2'}-H) and deprotonated (O_{2'}⁻) forms of the GpA molecule. In addition and importantly, the latter

calculations provided us with the structures for the study of the nucleophilic attack by the negatively charged O_2^- on the P atom, representing the step in barnase catalysis following the proton transfer from O_2^- -H to Glu73 (see Figure 1).

First, 100 ps of QM/MM molecular dynamics (MD) simulations were run on the full system for both the protonated and deprotonated GpA dinucleotides. Thereafter, a minimization step was performed and a minimum (*anti* conformation) for each GpA protonation form was obtained as the starting point for the PMF calculations. The PMFs were obtained using the umbrella sampling method.^{20,21} The added umbrella potential is a function of the reaction coordinate, χ (the guanosine glycosidic angle). In this study we took a harmonic form:

$$v(\chi) = \frac{1}{2}K(\chi - \chi^0)^2$$

where K is the force constant for the potential and χ^0 is a constant reference value for the reaction coordinate. Then, a series of molecular dynamics simulations were performed with different reference values of the reaction coordinate, χ^0 in the umbrella potential. The aim was to sample the complete range of values of χ . In practice, we found that using a force constant on the reaction coordinate of $1.5 \text{ kJ mol}^{-1} \text{ rad}^{-2}$ for the umbrella potential and reference values that differed by 0.6 degrees between simulations was sufficient to allow an adequate overlapping between consecutive simulation windows. We run 600 simulation windows for each protonated form of GpA, each consisting in 10 ps of equilibration and 10 ps of production. The values of the reaction coordinate were saved in order to obtain the histograms corresponding to each simulation, which were finally pieced together using the weighted histogram analysis method (WHAM)²² to obtain the full probability distributions function.

PMF Calculations of Guanosine Ribose O₂⁻ Attack on the P Atom for GpA in Water

The second goal of the present study was to examine the relationship between the *anti* and *syn* G χ conformations and the putative catalytic reaction mechanism (see Introduction). In particular, we investigated the second step of the enzymatic reaction, namely, the attack by the negatively charged guanosine ribose O₂⁻ on the P atom (see Figures 1 and 2) in the deprotonated *anti* and *syn* conformers identified in the PMF calculations described above. This involved performing PMF calculations as a function of the distance between the O₂⁻ and P atoms, using the same procedure described above. In these calculations, the force constant applied as constraint along the reaction coordinate was 2500 kJ mol⁻¹ Å⁻². At each simulation window the reaction coordinate was increased by 0.03 Å. Starting from the *syn* conformation a total of 60 simulation windows were required to cover the whole range of interest of the reaction coordinate, while 100 simulations were needed when using the *anti* conformation as starting point. Each window consisted, as before, in 10 ps of equilibration and 10 ps of production using 1 fs time step.

Structural and Statistical Analyses

To assess representative conformations for the minima and maxima of the PMF energy profiles, the mean and standard error of the mean (SEM) of each of the dihedral angles from the set of structures (100) saved in each of the corresponding windows were calculated. When the scatter plot of any of these angles relative to G χ (in the first PMF study) or to the guanosine O₂⁻-P distance (in the second PMF study) suggested the presence of more than one conformation, a K-means cluster analysis was performed. The clustering was done using the nearest centroid sorting method on the basis of

Euclidean distances computed from the nucleotide torsion angle values.²³ Since our aim was to find the minimum number of clusters able to characterize the main conformational families sampled in our systems, the number of clusters was fixed to the minimum number that was able to explain the observed angle distributions (see Ref. 12 for details). To determine which angles were responsible for the conformational differences between the initial and the final states of the PMF studies, the Student's t-test and the ANalysis Of VAriance (ANOVA) procedure, including the *post hoc* Tukey's test, were performed. *P*-values lower than 0.05 were considered statistically significant.

RESULTS

PMF for the Rotation of $G\chi$ in solvated GpA

Figure 3 shows the PMF profiles of the $G\chi$ torsion angle for the $O_2\text{-H}$ guanosine ribose (protonated) and O_2^- guanosine ribose (deprotonated) forms of GpA obtained in this study. Each of the profiles displays two minima and two maxima. For the protonated form, the $G\chi$ values are -88.8° , 21.0° , 71.5° , and 140.1° , on average, for the left minimum, left maximum, right minimum, and right maximum, respectively. For the deprotonated form the corresponding values are -80.4° , 44.9° , 63.7° , and 127.4° (see Figure 3, and Tables I and II). According to a recent conformational assignment for the χ angle,⁸ the two left minima correspond to *anti* conformers ($170\text{-}300^\circ$) whereas the two right minima correspond to *syn* conformers ($30\text{-}90^\circ$). Following the Klyne-Prelog angle notation,⁵ the two left maxima are found within the *+synperiplanar* (*+sp*: 0° to 30°) and the *+synclinal* (*+sc*: 30° to 90°) regions for the protonated and deprotonated forms, respectively. The two right maxima involve guanine-ribose orientations fitting the *+anticlinal* (*+ac*: 90° to 150°) conformations. The latter conformations correspond to the global maxima in our simulations, indicating that the *anti* \rightarrow *syn* $G\chi$ transitions in both protonation forms of GpA would prefer a pathway through 0° rather than through 180° .

We see furthermore that both GpA ionic forms display similar energy profiles, with lower energy minima for the *anti* conformations (on the left) than for the *syn* conformations (on the right) (Figure 3). These findings are in agreement with structural data from X-ray diffraction and NMR experiments as well as with theoretical QM studies, where a preference for *anti* χ conformations was observed (see Foloppe et al.,⁷ and references therein). The barriers for the *anti* \rightarrow *syn* $G\chi$ transition (through $\chi=0^\circ$) computed here are 4.0 and 3.9 kcal/mol for the protonated and deprotonated forms, respectively. These values are at the lower end of the range computed by QM methods

in vacuo (3.0-9.8 kcal/mol).⁷ It is noteworthy that the latter analysis⁷ indicated that the χ *anti* \rightarrow *syn* transition occurs preferably through 120° rather than through 0° as found here. However, MD simulations in aqueous solution performed by some of the authors of the same study revealed⁸ that, for purines, the *anti/syn* conversion takes place preferentially via $\chi = 0^\circ$, in agreement with our results. Finally, the value of ~ 4 kcal/mol that we compute for the *anti* \rightarrow *syn* $G\chi$ barriers is well above thermal energy at room temperature. It is therefore not too surprising that this transition was not observed in previous MD simulations.¹²

The conformational flexibility within each of the stationary states found in our $G\chi$ PMF study was examined by plotting the fluctuation of the GpA dihedral angles (for both protonation forms) as a function of the guanosine glycosidic angle taken as the reaction coordinate (Figure 4). For most of the sampled states the measured fluctuations were indicative of a uniform conformational population. In a number of states however several conformational families could be detected. Conformations were characterized by the mean and SEM values of the dihedral angles. Depending on whether one or more conformational populations were present, these values were computed either for the full ensemble of conformations or for subgroups of the sampled conformations, derived by a k-means cluster analysis (Tables I and II for the protonated and deprotonated GpA species, respectively). Statistical analysis of the computed values indicated that most torsion angles displayed significantly different behavior in the *anti* versus *syn* conformers (see Methods), suggesting in turn that *anti* \rightarrow *syn* transition is accompanied by a global conformational change of the nucleotide.

In addition we examined the conformations sampled by the *anti* and *syn* $G\chi$ states of the deprotonated GpA for their propensity to favorably position the guanosine O_{2'}, the central P, and the leaving adenosine O_{5'} atoms for the nucleophilic attack on the

phosphorus by the O_{2'} group. Both shorter distances between O_{2'} and P atoms and wider O_{2'}-P-O_{5'} angles were found in the *syn* relative to the *anti* G χ conformation (Figure 4B and Table II), suggesting that the ribonucleotide cleavage reaction is favored in the *syn* conformation. This is illustrated in Figure 5, which shows representatives of the *anti* and *syn* molecular structures (4.322 and 3.769 Å, for the *anti* and *syn* conformations, respectively) featuring O_{2'}-P distances closest to the mean values (4.32 and 3.77 Å, Table II) observed in the sampled conformations. Likewise, the corresponding O_{2'}-P-O_{5'} angle in these structures takes the values of 81.15° and 90.10°, for the *anti* and *syn* conformers, respectively (also close to the measured mean values of 85.1° and 93.8°, Table II), and is thus more open in the *syn* than in the *anti* conformer. To further explore this hypothesis, PMF calculations along the O_{2'}-P distance were performed for the deprotonated GpA molecule.

PMF Calculations of the Guanosine Ribose O_{2'} Attack on the P Atom in Solvated GpA

Figure 6 shows the PMF profiles of the O_{2'} nucleophilic attack on the P atom for deprotonated-O_{2'} GpA with the *anti* and *syn* G χ conformations. The starting conformations were taken from the minima of the G χ PMF profile of the deprotonated GpA form in Figure 3, and all PMF calculations were done in presence of explicit solvent. In good agreement with our first PMF calculations, the initial state for the *syn* conformation is 3.4 kcal/mol higher in energy than that of the *anti* conformation, taken as a reference. The final state is a pentacovalent intermediate with an O_{2'}-P covalent bond.

The most striking differences between the PMF profiles of the *syn* and *anti* GpA conformers are in the height of the energy barrier separating the initial and final states.

This barrier is 14.6 kcal/mol for the *anti* but only 2.7 kcal/mol for the *syn* conformation. These large differences likely reflect fundamental differences in the conformational and energetic landscape of the process when the latter takes place while the nucleotide adopts the *anti* or *syn* $G\chi$ conformations (see Ref. 24 for a discussion on the stereochemistry of enzymatic reactions involving the phosphate group and a recent theoretical study on the nucleophilic attack on phosphate diesters²⁵). In the following we investigate some of these differences.

Tables III and IV list the mean and SEM values of torsion angles, guanosine $O_{2'}$ -P distances, and guanosine $O_{2'}$ -P-adenosine $O_{5'}$ angles for the collections of 100 structures generated in each of the windows corresponding to the stationary states (the two minima and the maximum connecting both states) of the considered PMF. Our results reveal that the $O_{2'}$ attack starting from the *syn* conformation is structurally more favorable than that starting from the *anti* conformation. First, the initial distance between the two interacting atoms ($O_{2'}$ and P) is shorter for the *syn* conformation (3.55 Å versus 4.29 Å). Second, the $O_{2'}$ in the *syn* conformation is more favorably oriented for the progression of the reaction: the $O_{2'}$ -P- $O_{5'}$ angle equals 130.9°, 154.6°, and 147.0° for the initial minimum, TS, and pentacovalent intermediate, respectively, whereas smaller values for this angle (84.9°, 69.2°, and 82.0) are observed for the same states in the *anti* $G\chi$ conformation.

Comparisons between the final (pentacovalent) and the initial (either *anti* or *syn*) states of the PMF profile showed statistical significance for all the selected parameters (dihedral angles and the $O_{2'}$ -P- $O_{5'}$ angle; Tables III and IV), indicating that formation of the pentacovalent species involves conformational rearrangements throughout the entire dinucleotide structure. Furthermore, the pentacovalent intermediates resulting from *anti* and *syn* $G\chi$ conformations are structurally different as Tables III and IV show; in

particular, the dihedral angles ($G\epsilon$, $G\zeta$, $A\alpha$, and $A\beta$) involving the central P atom show distinct values. Figure 7 displays the representative molecular structures for the pentacovalent intermediates of the *anti* and *syn* $G\chi$ conformational families. We see that within the *syn* $G\chi$ conformation, the attacking $O_{2'}$ and leaving $O_{5'}$ groups are situated approximately at the apical positions of a trigonal bipyramid, thereby fitting an in-line mechanism for ribonucleotide cleavage. In contrast, the attack from the *anti* $G\chi$ conformation resembles an adjacent mechanism, for which the nucleophile enters on the same side as the leaving group. It is worth noting that the *syn* minimum obtained in the PMF and described in Table IV is different from that reported in Table II obtained from the conformational PMF. In particular, the values of the $O_{2'}$ -P distance and the $O_{2'}$ -P- $O_{5'}$ angle show that the present *syn* minimum is slightly more advanced for the nucleophilic attack on the phosphorus atom. This difference is not unexpected as the conformational and reaction PMFs are obtained using different reaction coordinates, corresponding to the integration over different regions of the configurational space. In addition, the behavior of the PMF presented in Figure 6 shows that the free energy landscape can be quite flat for large $O_{2'}$ -P distances in the *syn* conformation and thus no important free energy differences are expected between the two reported (Tables II and IV) *syn* minima.

Taken together, the energy differences between the *anti* and *syn* minima of deprotonated- $O_{2'}$ GpA and the barriers for both the $G\chi$ rotation and the $O_{2'}$ attack on the phosphorus atom in both the *anti* or *syn* conformations (Figures 3 and 6) lead us to postulate that the nucleophilic attack on the phosphorus atom by the $O_{2'}$ takes place when the GpA molecule adopts the *syn* conformation. With our analysis confirming that the *anti* conformation of the dinucleotide is the predominant molecular species in solution, and hence likely to be that of the substrate of the enzymatic reaction, we

conclude that an *anti* \rightarrow *syn* transition of the glycosidic angle must occur prior to the nucleophilic attack.

DISCUSSION AND CONCLUDING REMARKS

The general acid-base mechanism¹ proposed for barnase catalysis, with Glu73 acting as the general base, together with structural data from nucleotide-barnase complexes^{4,9-11} suggest that a *syn* conformation for the $G\chi$ angle is required in order to allow the abstraction of the ribose $H_{2'}$ proton by Glu73, a key step of barnase catalysis. The work described here provides insight into the link between this requirement and the intrinsic conformational properties of the solvated nucleotide and, by the same token, allows us to gauge the role that the enzyme scaffold may play. The first PMF calculations for GpA in water as a function of the $G\chi$ torsion angle allowed us to characterize the influence of this angle on the GpA conformational landscape. In addition it also provided us with the initial structures for studying the nucleophilic attack by the ribose $O_{2'}$ on the P atom. Our second series of calculations, which simulated the $O_{2'}$ -P covalent bond formation, showed that the *syn* $G\chi$ conformation favors the in-line attack of the nucleophile. Since the nucleophile is believed to be produced through proton abstraction by the catalytic residue Glu73, our latter finding thus provide a rationalization for the relative orientation of this residue in the barnase active site.

It is noteworthy to relate the conclusions from our PMF calculations in water with information derived from the crystal structure of the barnase-d(CGAC)] complex,⁴ where the guanine base adopts an *anti* conformation, and from a modeled barnase-5'3'(AAGAAp)-O-methyl ester complex,⁹ in which $G\chi$ was rotated to achieve a *syn* conformation. In the crystal structure, the guanosine $O_{2'}$ atom, which was added to the coordinates of the bound deoxynucleotide, was found to be positioned ~ 7 Å away from the carboxylate of Glu73, too far to enable proton abstraction by the latter. On the other hand, in the modeled complex,⁹ the corresponding groups were positioned at a hydrogen

bonding distance, optimal for proton abstraction (Figure 8). What more, in both the crystal and modeled complexes, the GpA molecule displays conformational properties in good agreement with those described in this work. Thus, for the *syn* guanosine (Figure 8 right), the $O_{2'}$ (following proton abstraction by Glu73) is correctly oriented towards the P- $O_{5'}$ bond for an in-line attack. This is clearly not the case for the guanosine in the *anti* conformation (Figure 8 left). It is thus quite remarkable to see that two seemingly distinct processes, the initiation of catalysis through the abstraction of the $O_{2'}$ proton and the conformational transitions towards the pentacovalent intermediate, are both fostered by the $G\chi$ *syn* conformation of the nucleotide and not the *anti* state.

The work presented here, and in particular the results on the lower energy barrier for reaching the pentacovalent intermediate in the *syn* versus the *anti* conformation clearly concern only the solvated unbound GpA molecule. They hence provide no information on the GpA conformer (*syn* or *anti*) that may bind to the enzyme active site, or in the case that the *anti* conformation is the one to bind, what the barrier for the *anti* to *syn* transition might be for the bound dinucleotide. It should be mentioned however, that the only known case of GpA bound to barnase adopting the $G\chi$ *anti* conformation is that of the [barnase-d(CGAC)] crystal structure,⁴ where the ligand is a deoxynucleotide inhibitor that cannot be cleaved by the enzyme.

Overall therefore, our calculations and the above considerations suggest that there exists a link between $G\chi$ torsion angle adopting a *syn* conformation and the requirement to have a negatively charged $O_{2'}$ in order to carry out the in-line attack on the phosphorus atom. This in turn suggests that the active site of barnase might have been shaped by evolution in a way in which abstraction of $H_{2'}$ and in-line orientation of $O_{2'}$ were coupled. This idea is in agreement with the concept of enzyme

preorganization.²⁶ According to this concept, the catalytic power of an enzyme is mainly due to transition state stabilization (TSS), achieved through the electrostatic environment provided by its active site. The greater similarity between the transition state (TS) and the catalytic conformation than between the TS and the noncatalytic conformations^{27,28} suggests that TSS involves the destabilization of the noncatalytic conformations (*anti* $G\chi$ conformation in barnase catalysis), thereby providing an additional rate enhancement of the enzyme catalyzed reaction relative to its uncatalyzed counterpart in solution.²⁹

Coupling between a specific substrate conformation and molecular reactivity and the connection of this property with enzyme structure and mechanism may illustrate the functioning of enzymes as finest machines, evolved to optimize all the processes that favor the productive pathway of the reactions they catalyze. To explore further these ideas, studies of the ribonucleotide cleavage in the barnase environment are in progress.

ACKNOWLEDGEMENTS

V.M. thanks the DGI for projects DGI CTQ2006-15447-C02-01/BQU, UJI-BANCAIXA foundation for project P1·1B2005-13 and Generalitat Valenciana for project GV06/152. I.T. is indebted to Ministerio Educación y Ciencia for project CTQ2006-15447-CO2-02 and Generalitat Valenciana for project GV06-021. J.G. is grateful to Ministerio de Educación y Ciencia for project SAF2004-06134.

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Figure legends

Figure 1. Schematic representation of the first two steps of ribonucleotide cleavage mechanism by barnase. Step 1: Glu73 picks up a proton from ribose O_{2'} (see Figure 2 for atom notation) and the negatively charged O_{2'} attacks on and makes a covalent bond with the P atom. Step 2: The positively charged His102 gives a proton to O_{5'} and the O_{5'}-P bond breaks.

Figure 2. Notation scheme for atoms and torsion angles in GpA. Torsion angles are defined as: $\alpha = \text{O}_{3'}-\text{P}-\text{O}_{5'}-\text{C}_{5'}$, $\beta = \text{P}-\text{O}_{5'}-\text{C}_{5'}-\text{C}_{4'}$, $\gamma = \text{O}_{5'}-\text{C}_{5'}-\text{C}_{4'}-\text{C}_{3'}$, $\delta = \text{C}_{5'}-\text{C}_{4'}-\text{C}_{3'}-\text{O}_{3'}$, $\varepsilon = \text{C}_{4'}-\text{C}_{3'}-\text{O}_{3'}-\text{P}$, $\zeta = \text{C}_{3'}-\text{O}_{3'}-\text{P}-\text{O}_{5'}$, $\chi = \text{O}_{4'}-\text{C}_{1'}-\text{N}_9-\text{C}_4$. For a given X angle, GX and AX stand for guanosine and adenosine X angles, respectively.

Figure 3. PMF profiles of the G χ rotation for two GpA ionic forms in water: protonated (guanosine ribose O_{2'}-H) GpA (green line) and deprotonated (guanosine ribose O_{2'}) GpA (red line).

Figure 4. Scatter plots of G χ with the rest of dihedral angles for the stationary states (minima and maxima) found in Figure 3. A) Protonated GpA. B) Deprotonated GpA. For the G χ , the (-180° to +180°) range was chosen; for the other angles, the former or the (0° to 360°) ranges were selected; nevertheless, the polar nature of angles should be taken into account when looking at the plots. In Figure B (deprotonated GpA), the guanosine O_{2'}-P distance and the guanosine O_{2'}-P-adenosine O_{5'} angle are also examined.

Figure 5. Molecular structures of deprotonated GpA representative members of the *anti* and *syn* stationary states obtained in the PMF profile of the $G\chi$ rotation displayed in Figure 3. These structures have been selected because they are those with the guanosine $O_{2'}$ -P distances closest to their corresponding conformational group mean values (Table II). The *syn* conformation favors the $O_{2'}$ attack on the P atom relative to the *anti* conformation: $O_{2'}$ -P distances of 3.769 and 4.322 Å and $O_{2'}$ -P- $O_{5'}$ angles of 90.10° and 81.15° for the *syn* and *anti* conformations, respectively. The guanosine ribose $O_{2'}$ and the P atoms are connected by a dotted line.

Figure 6. PMF profiles (red: *anti* $G\chi$ and green: *syn* $G\chi$) of the $O_{2'}$ attack on the P atom for deprotonated (guanosine ribose $O_{2'}$) GpA. The *anti* and *syn* $G\chi$ starting points were taken from previously determined minima (Figure 3): the *syn* is higher in energy than the *anti*, being the latter taken as the reference (0 kcal/mol).

Figure 7. Molecular structures representative of the pentacovalent intermediates obtained in the PMF profiles of the $O_{2'}$ attack displayed in Figure 6 from either *anti* or *syn* $G\chi$ conformations. The structure for the $G\chi$ *syn* conformation (on the right), in contrast to the structure for the *anti* $G\chi$ conformation (on the left), displays a trigonal bipyramid with the entering and leaving groups in apical positions.

Figure 8. Molecular structures showing the conformational arrangement of the GpA fragment for barnase-d(CGAC) crystal⁴ (left) and barnase-5'3'(AAGAAp)-*O*-methyl ester modeled⁹ (right) complexes. For the crystal structure the ribose $O_{2'}$ atoms were modeled because deoxyribonucleotides need to be used in experimental studies to avoid enzyme catalysis. In the crystal complex the guanosine displays an *anti* conformation

($G\chi=215^\circ$) whereas in the modeled complex the guanosine displays a *syn* conformation ($G\chi=95^\circ$). In the *anti* conformation, the guanosine $O_{2'}$ is far away ($\sim 7 \text{ \AA}$) from the oxygens of Glu73 whereas in the *syn* conformation, a hydrogen bonding distance between both moieties is found. In the *syn* guanosine conformation the $O_{2'}$ is correctly positioned for an in-line attack on the P atom (once its proton is abstracted by Glu73) in contrast to the *anti* guanosine conformation.

TABLE I. $G\chi$ PMF study: Torsion and pseudorotation angles for protonated ($O_2\text{-H}$) GpA.

Torsion and pseudorotation angles	Minimum (<i>anti</i>)	TS (<i>anti</i> \rightarrow <i>syn</i>)	Minimum (<i>syn</i>)	Absolute maximum
$G\chi$	-88.8 ± 0.1	21.0 ± 0.1	71.5 ± 0.1	140.1 ± 0.1
$G\gamma$	$192.1 \pm 0.9^*$	295.9 ± 1.1	277.9 ± 0.9	167.2 ± 1.4
$G\delta$	141.6 ± 0.7	127.0 ± 0.6	140.0 ± 0.6	126.3 ± 0.6
Gp	194.1 ± 2.2	N=10 83.8 ± 13.9 N=90 246.1 ± 3.2	188.9 ± 1.8	290.6 ± 3.1
$G\epsilon$	$237.2 \pm 0.7^*$	279.6 ± 0.8	277.6 ± 0.8	202.2 ± 1.0
$G\zeta$	$215.4 \pm 1.0^*$	176.8 ± 0.6	116.9 ± 1.2	108.8 ± 1.3
$A\chi$	$82.4 \pm 1.0^*$	59.6 ± 0.9	88.9 ± 1.3	63.8 ± 1.1
$A\alpha$	$98.8 \pm 1.0^*$	297.3 ± 1.0	153.3 ± 2.6	182.3 ± 2.0
$A\beta$	$73.4 \pm 1.2^*$	173.8 ± 1.0	156.9 ± 2.1	155.0 ± 1.3
$A\gamma$	$181.1 \pm 0.8^*$	188.5 ± 0.8	279.3 ± 1.8	193.8 ± 1.5
$A\delta$	125.4 ± 0.7	119.7 ± 0.7	124.6 ± 0.5	100.8 ± 0.7
Ap	N=38 $213.1 \pm 6.5^{A,B}$ N=62 121.5 ± 3.6^B	115.8 ± 2.2	A N=88 281.6 ± 3.2 B N=12 114.0 ± 13.9	N=7 327.5 ± 7.1 N=93 45.3 ± 2.3

Mean \pm standard errors of torsion angles of 100 structures saved in each of the windows corresponding to the stationary states of $G\chi$ PMF energy profile (Figure 3). A K-means cluster analysis was used when scatter plots (Fig. 4A) suggested that more than one conformational family was present. Gp and Ap correspond to the pseudorotation angles for guanosine and adenosine riboses, respectively. Comparisons between *anti* and *syn* conformations for each of the angles were performed by Student's t-test or the Tukey's *post hoc* test if two or more than two conformational populations were involved, respectively. *P*-values lower than 0.05 are indicated by * and considered statistically significant. If the *syn* conformation presents two subpopulations (A,B) then A and/or B in the *anti* conformation indicates statistical significance with A and/or B *syn* conformation, respectively.

TABLE II. $G\chi$ PMF study: Torsion and pseudorotation angles, guanosine O_2 -P distance, and guanosine O_2 -P- O_5 adenosine angle for deprotonated (O_2^-) GpA.

Variables	Minimum (<i>anti</i>)	TS (<i>anti</i> \rightarrow <i>syn</i>)	Minimum (<i>syn</i>)	Absolute maximum
$G\chi$	-80.4 \pm 0.1	44.9 \pm 0.1	63.7 \pm 0.1	127.4 \pm 0.1
$G\gamma$	46.6 \pm 1.2*	173.2 \pm 0.8	182.4 \pm 1.7	178.2 \pm 1.1
$G\delta$	143.8 \pm 0.6*	141.7 \pm 0.6	136.8 \pm 0.7	125.2 \pm 0.6
$G\rho$	204.9 \pm 1.5*	195.7 \pm 1.3	187.8 \pm 2.2	277.5 \pm 1.8
$G\varepsilon$	291.0 \pm 1.0*	246.3 \pm 1.0	232.1 \pm 0.8	240.5 \pm 1.0
$G\zeta$	197.5 \pm 1.6	209.0 \pm 1.4	194.4 \pm 0.8	178.7 \pm 0.9
$A\chi$	150.9 \pm 1.1*	262.3 \pm 0.9	260.9 \pm 0.8	276.1 \pm 0.8
$A\alpha$	N=44 220.3 \pm 2.3* N=56 121.1 \pm 2.5*	143.1 \pm 1.2	143.2 \pm 1.2	70.3 \pm 1.2
$A\beta$	N=23 63.0 \pm 1.5* N=33 28.1 \pm 3.9* N=44 76.3 \pm 2.5*	89.7 \pm 1.6	100.5 \pm 1.0	105.4 \pm 0.9
$A\gamma$	169.5 \pm 3.1*	178.7 \pm 0.9	177.6 \pm 0.9	194.2 \pm 0.8
$A\delta$	120.5 \pm 0.8	126.0 \pm 0.8	122.6 \pm 0.8	138.5 \pm 0.9
$A\rho$	N=23 308.5 \pm 3.7 ^{A,B} N=69 262.9 \pm 2.2 ^B N=3 81.2 \pm 7.3 ^{A,B} N=5 12.0 \pm 3.0 ^A	N=20 277.7 \pm 6.5 N=72 196.5 \pm 3.6 N=8 64.5 \pm 11.6	A N=92 263.8 \pm 3.5 B N=8 23.3 \pm 6.3	241.1 \pm 2.6
Guanosine O_2 -P distance	4.32 \pm 0.01*	3.96 \pm 0.01	3.77 \pm 0.01	3.97 \pm 0.02
Guanosine O_2 -P- adenosine O_5 angle	85.1 \pm 0.5*	85.6 \pm 0.8	93.8 \pm 0.6	98.7 \pm 0.4

Mean \pm standard errors of torsion angles, guanosine O_2 -P distances, and guanosine O_2 -P- O_5 adenosine angles of 100 structures saved in each of the windows corresponding to the stationary states of $G\chi$ PMF energy profile (Figure 3). A K-means cluster analysis was used when scatter plots (Fig. 4B) suggested that more than one conformational family was present. $G\rho$ and $A\rho$ correspond to the pseudorotation angles for guanosine and adenosine riboses, respectively. Comparisons between *anti* and *syn* conformations for each of the variables were performed by Student's t-test or Tukey's *post hoc* test if two or more than two populations were involved, respectively. *P*-values lower than 0.05 are indicated by * and considered statistically significant. If the *syn* conformation presents two subpopulations (A,B) then A and/or B in the *anti* conformation indicates statistical significance with A and/or B *syn* conformation, respectively.

TABLE III. Guanosine O_{2'}→P PMF study under *anti* G χ conformation: Torsion and pseudorotation angles, guanosine O_{2'}-P distance, and guanosine O_{2'}-P-adenosine O_{5'} angle for deprotonated (O_{2'}⁻) GpA.

Variables	Minimum (<i>anti</i>)	TS (<i>anti</i> →pentacovalent)	Minimum (pentacovalent from <i>anti</i>)
G χ	-64.3±0.7*	-59.1±0.8	-72.6±0.9
G γ	81.7±1.5*	68.8±1.6	67.2±1.2
G δ	140.1±0.6*	135.6±0.8	115.5±0.7
G ρ	212.0±2.0*	217.5±2.8	-2.6±7.5
G ϵ	289.0±0.9*	181.7±0.9	106.2±0.7
G ζ	181.3±1.0*	268.7±1.3	-8.7±2.2
A χ	182.9±0.8*	168.5±1.1	189.2±1.0
A α	111.5±0.9*	243.1±0.8	124.8±2.0
A β	58.5±1.2*	288.2±0.9	66.2±2.0
A γ	151.0±*	233.9±0.6	142.1±1.5
A δ	144.4±0.9*	147.8±0.9	126.2±1.4
A ρ	227.6±1.2*	240.0±1.5	-82.7±4.6
Guanosine O _{2'} -P distance	4.29±0.00	2.78±0.00	1.74±0.00
Guanosine O _{2'} -P-O _{5'} adenosine angle	84.9±0.4*	69.2±0.4	82.0±0.3

Mean \pm standard errors of torsion angles, guanosine O_{2'}-P distances, and guanosine O_{2'}-P-adenosine O_{5'} angles of 100 structures saved in each of the windows corresponding to the stationary states of the guanosine O_{2'} attack on the P atom from a PMF energy profile (Figure 6). G ρ and A ρ correspond to the pseudorotation angles for guanosine and adenosine riboses, respectively. Comparisons between Minimum *anti* and Minimum pentacovalent conformations for each of the variables were performed by Student's t-test. *P*-values lower than 0.05 are indicated by * and considered statistically significant.

TABLE IV. Guanosine O_{2'}→P PMF study under *syn* G χ conformation: Torsion and pseudorotation angles, guanosine O_{2'}-P distance, and guanosine O_{2'}-P-adenosine O_{5'} angle for deprotonated (O_{2'}⁻) GpA.

Variables	Minimum (<i>syn</i>)	TS (<i>syn</i> →pentacovalent)	Minimum (pentacovalent from <i>syn</i>)
G χ	71.0±0.9*	96.5±1.0	74.9±0.9
G γ	38.2±2.4*	59.1±2.8	47.8±1.9
G δ	124.2±0.6*	141.4±0.6	121.9±0.9
G ρ	155.0±3.0*	184.1±2.0	125.4±3.0
G ϵ	219.2±0.6*	183.4±0.6	139.0±1.5
G ζ	144.8±0.8*	136.4±0.8	180.2±1.7
A χ	83.3±1.5*	78.3±1.4	77.2±0.8
A α	108.4±1.1*	113.4±1.1	144.1±2.0
A β	144.4±1.1*	146.6±1.0	153.6±1.3
A γ	177.7±0.8*	189.1±0.9	189.6±0.9
A δ	110.0±1.0*	109.2±0.8	113.2±0.8
A ρ	18.0±5.0*	-23.9±3.9	-26.6±5.1
Guanosine O _{2'} -P distance	3.55±0.00	2.69±0.00	1.80±0.00
Guanosine O _{2'} -P-O _{5'} adenosine angle	130.9±0.4*	154.6±0.4	147.0±1.1

Mean ± standard errors of torsion angles, guanosine O_{2'}-P distances, and guanosine O_{2'}-P-O_{5'} adenosine angles of 100 structures saved in each of the windows corresponding to the stationary states of the guanosine O_{2'} attack on the P atom from a PMF energy profile (Figure 6). G ρ and A ρ correspond to the pseudorotation angles for guanosine and adenosine riboses, respectively. Comparisons between Minimum *anti* and Minimum pentacovalent conformations for each of the variables were performed by Student's t-test. *P*-values lower than 0.05 are indicated by * and considered statistically significant.

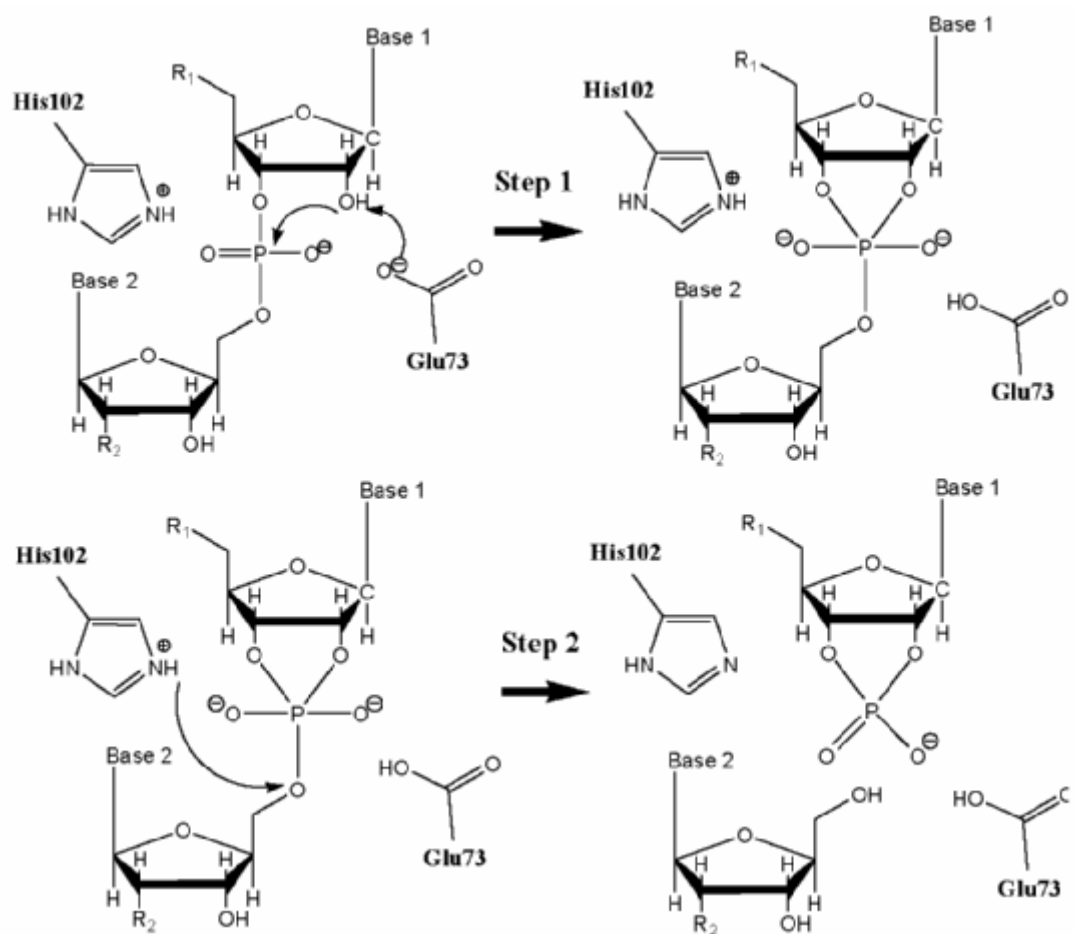


Fig. 1

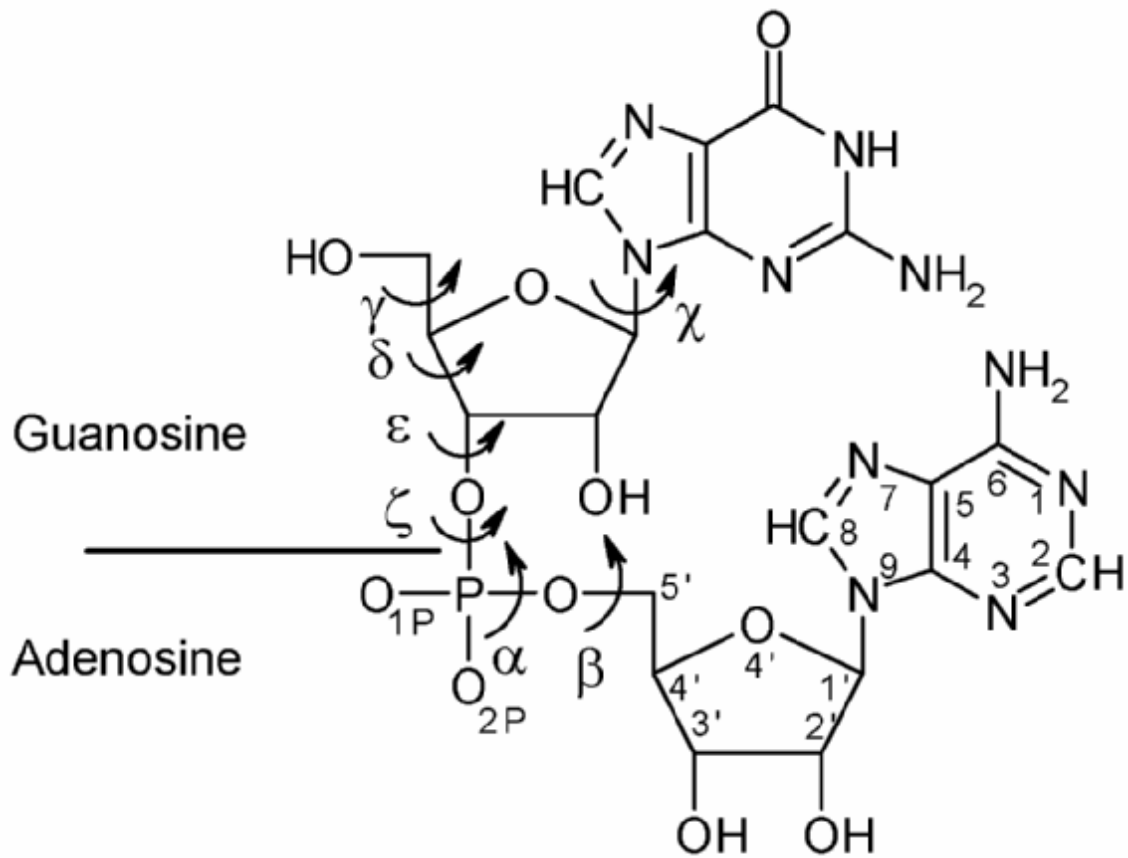


Fig. 2

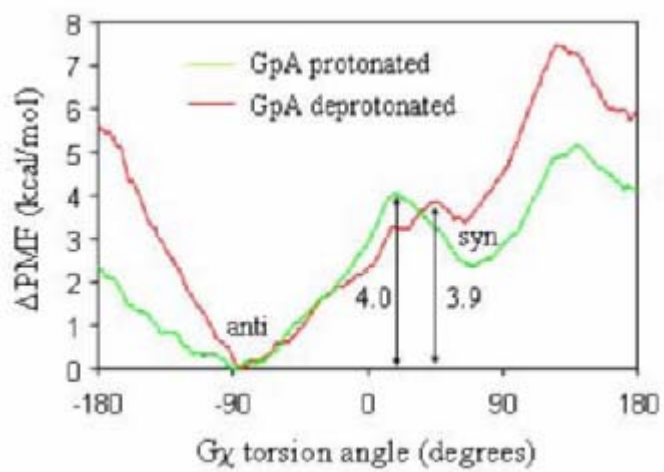


Fig. 3

a

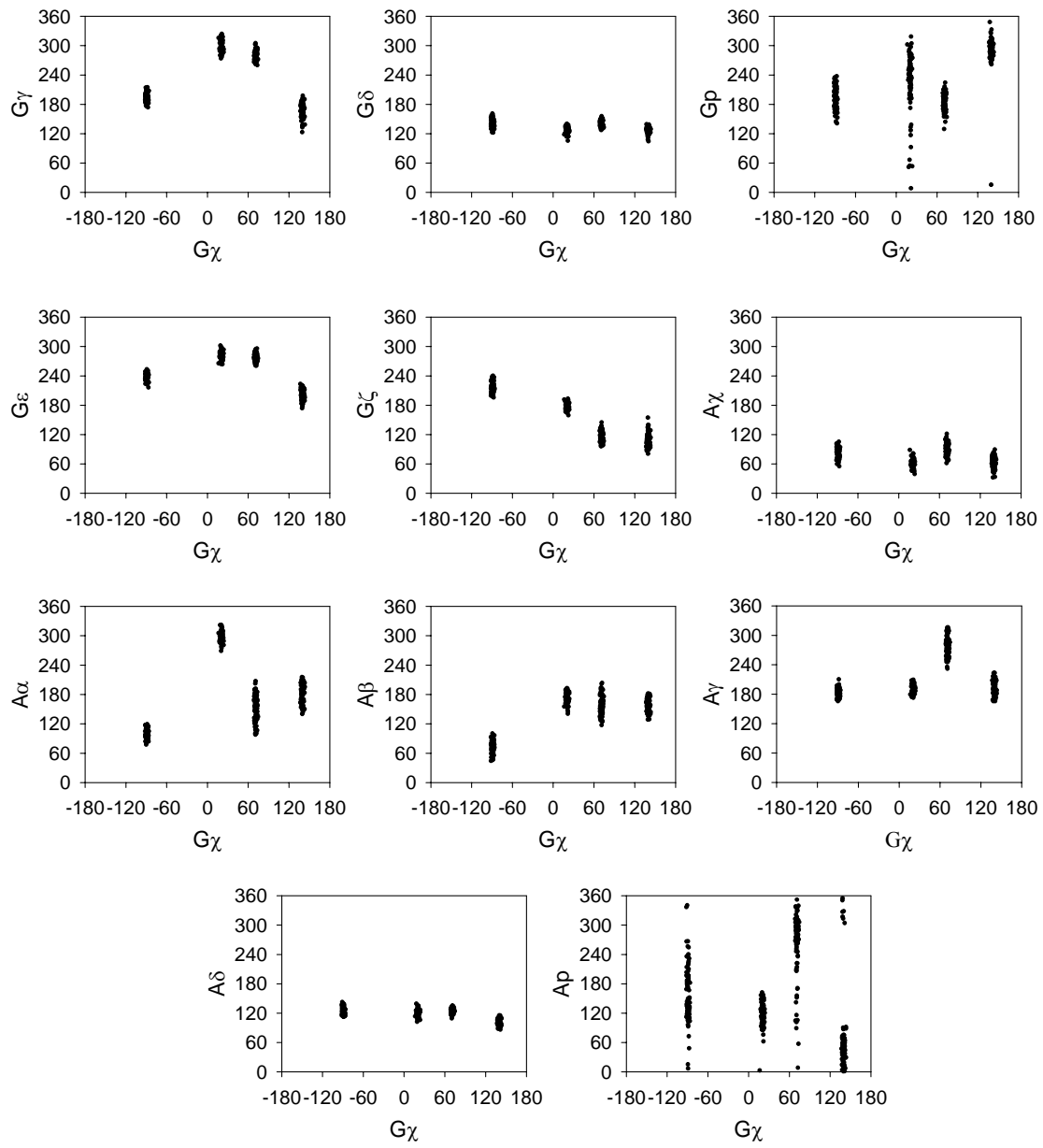


Fig. 4

b

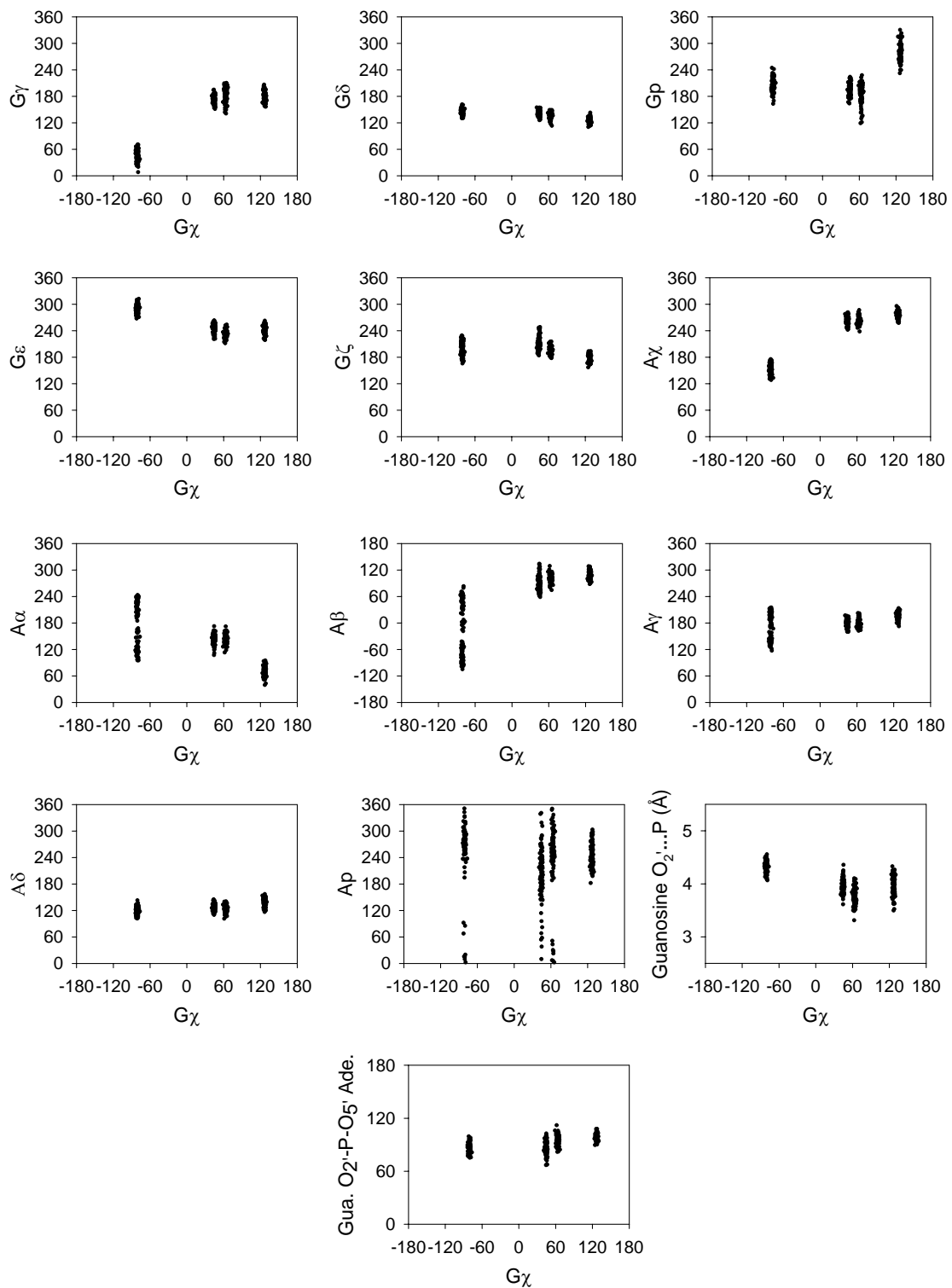


Fig. 4

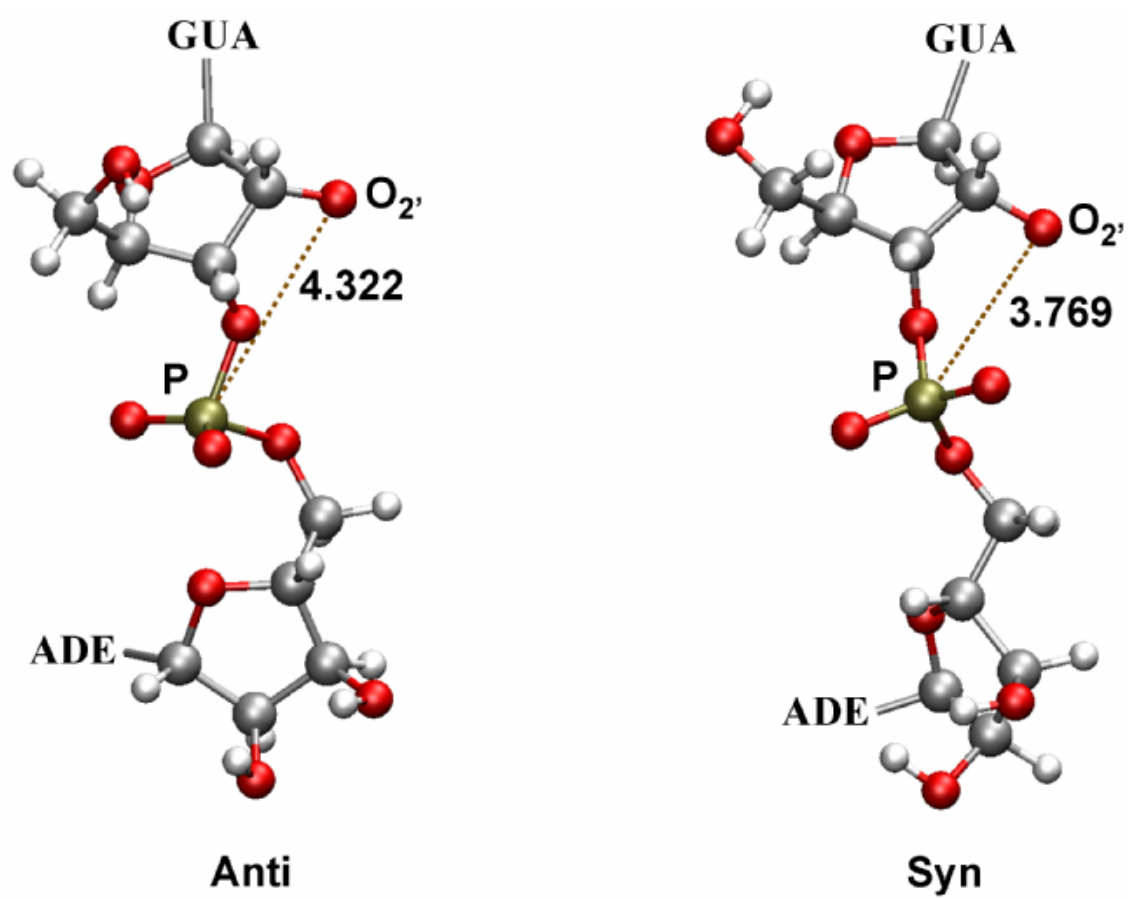
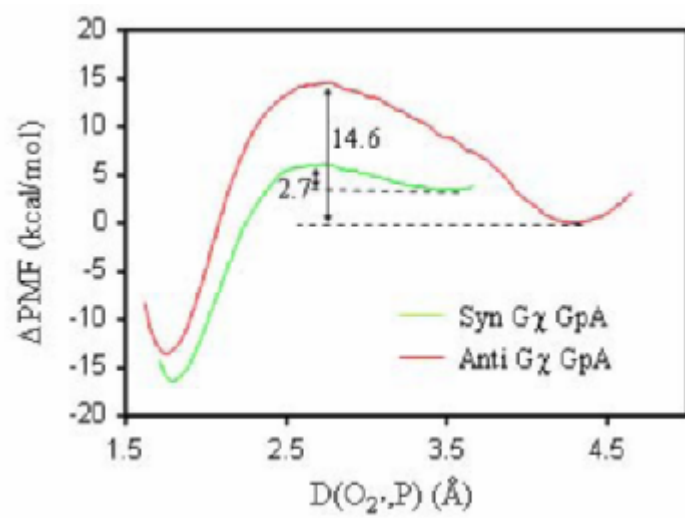
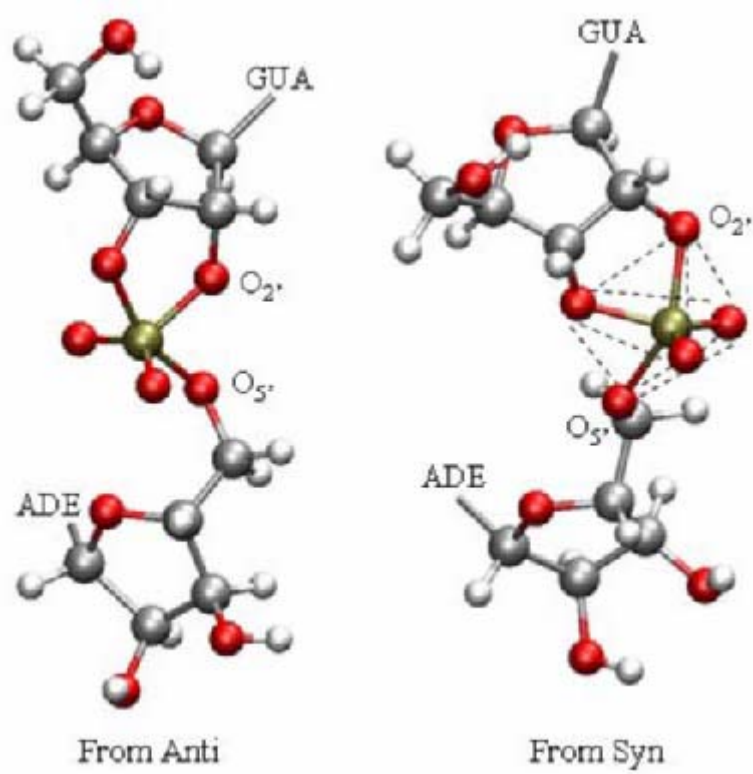
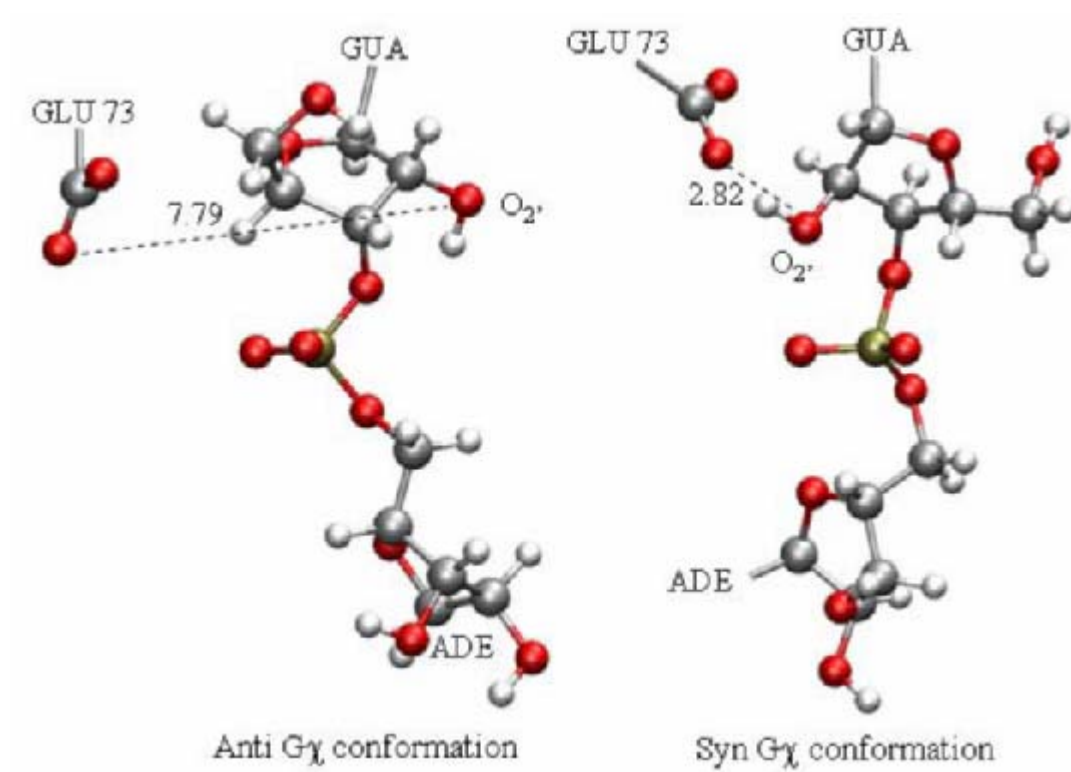


Fig. 5

**Fig. 6**

**Fig. 7**

**Fig. 8**