

NIH Public Access

Author Manuscript

Biochemistry. Author manuscript; available in PMC 2010 August 4.

Published in final edited form as:

Biochemistry. 2009 August 4; 48(30): 7160-7168. doi:10.1021/bi900830g.

The Flexibility of a Distant Loop Modulates Active-Site Motion and Product Release in Ribonuclease A

Nicolas Doucet¹, Eric D. Watt¹, and J. Patrick Loria^{1,2,*}

¹ Department of Chemistry, Yale University, New Haven, Connecticut 06520

² Department of Molecular Biophysics and Biochemistry, Yale University, New Haven, Connecticut 06520

Abstract

The role of the flexible loop 1 in protein conformational motion and the dissociation of enzymatic product from Ribonuclease A (RNase A) was investigated by creation of a chimeric enzyme in which a six residue loop 1 from the RNase A homolog, eosinophil cationic protein (ECP) replaced the twelve residue loop in RNase A. The chimera (RNase A_{ECP}) experiences only local perturbations in NMR backbone chemical shifts compared to WT RNase A. Many of the flexible residues that were previously identified in WT as involved in an important conformational change now experience no NMR-detected millisecond motions in the chimera. Likewise, binding of the product analog, 3'-CMP to RNase A_{ECP} results in only minor chemical shift changes in the enzyme similar to what is observed for the H48A mutant of RNase A and in contrast to WT enzyme. For both RNase A_{ECP} and H48A there is a 10-fold decrease in the product release rate constant, k_{off} compared to WT and in agreement with previous studies indicating the importance of flexibility in RNase A in the overall rate-limiting product release step. Together these NMR and biochemical experiments provide additional insight into the mechanism of millisecond motions in the RNase A catalytic cycle.

Keywords

Ribonuclease; NMR relaxation dispersion; protein dynamics; flexibility; loops

Protein motions are an essential aspect of enzyme function. These motions can enable proper positioning of catalytic residues, sequester reactive intermediates, allow access of substrate to the enzyme's active site, and facilitate the release of product(s) (1–10). Ribonuclease A (RNase A) is one example in which the backbone and side chain positions change conformation as the enzyme interacts with substrate, product and transition-state analogs (3,11–28), indicating the close relationship between the chemical nature of the bound ligand and the structure of the protein. Solution NMR relaxation experiments have characterized these motions in RNase A, localizing them to a specific region of the enzyme including the active site and a distant loop (loop 1) as well as some surrounding residues (3,13,21,22,25,27–30). There is increasing evidence for the importance of this motion in RNase A function including the similarity of the timescales for protein motion and the catalytic turnover rate (k_{cat}) (13,29), a similar pH dependence of RNA transphosphorylation and protein motion (13), nearly identical kinetic solvent isotope effects (KSIE) on k_{cat} and on motion (27,28), and the observation that site-

^{*}CORRESPONDING AUTHOR Telephone: (203)-436-4847; Fax (203)-432-6144. (patrick.loria@yale.edu). SUPPORTING INFORMATION AVAILABLE

Experimental results for the differential multiple-quantum relaxation experiments (Figure S1, Table S1) and field dependent R_{ex} values for WT and RNase A_{ECP} (Table S2). This material is available free of charge via the Internet at http://pubs.acs.org/.

directed mutations that decrease k_{cat} also cause a reduction or abolition of protein motion (3, 28).

Many of these motions in RNase A are localized in a distinct area, approximately 20 Å from the active-site histidine residues (His12 and His119), that encompasses loop 1 (Figure 1) and several surrounding residues. In RNase A, this loop extends from residues 14 to 25 and contains the sequence, DSSTSAASSSNY. An important interaction identified in previous work involves His48, which makes a hydrogen bond via its $N^{\epsilon 2}$ atom with the O^{γ} of T17 in loop 1. Mutation of His48 to alanine results in a loss of motion in and surrounding loop 1, a 10-fold decrease in catalytic activity, and a loss of the KSIE on the transphosphorylation of RNA (28). These experiments suggested that the motion of loop 1 was somehow coupled via His48 to the active site and that disruption of these interactions leads to less than optimal enzymatic function. Loop 1 in RNase A assumes different conformations in the various liganded states (29–32) and NMR data indicates that this loop is flexible and moves on the timescale of the rate-limiting product release step (13,29). These experimental data implicate motion of this loop as an essential component of the product release step. Two other interesting observations suggest that the flexibility of this loop is essential for its function in this capacity. First, RNase A structural homologs, eosinophil cationic protein (ECP) and eosinophil derived neurotoxin (EDN) that have low ribonuclease activity have a much shorter loop 1 (Figure 1), which would restrict its motion and thus its ability to access conformations available to loop 1 in RNase A. Also, unlike RNase A, which shows distinct loop 1 conformations in the apo and bound states, there are insignificant differences in the ligated and unligated conformations of ECP and EDN (33,34), suggesting that loop 1 in the eosinophil enzymes is immobile or unresponsive to different ligand structures. However, the details of the role of loop 1 motion in RNase A function remain unclear.

In this work we investigate the effects of swapping the short ECP loop 1 into the RNase A enzyme by creating a recombinant chimera (RNase A_{ECP}). NMR dynamics experiments indicate a similar abolition of motion in this region of the enzyme upon loop swapping. The product release step as measured by NMR line shape analysis indicates a significant decrease in the k_{off} value in the chimeric enzyme compared to WT RNase A, implicating the motion of loop 1 in enabling efficient release of product.

MATERIALS AND METHODS

RNase A_{ECP} DNA Construct

Oligonucleotide synthesis was performed by the W. M. Keck Foundation Biotechnology Resource Laboratory (Yale University, New Haven, CT) and by Integrated DNA Technologies (Coralville, IA). The wild-type, 372-bp bovine RNase A gene was codon-optimized for better expression in *E. coli* by GenScript Co. (Piscataway, NJ), generating construct pUC57-RNaseA (CO). The chimeric RNase A_{ECP} gene was PCR-amplified from that construct using the two following terminal primers: NdeI-RNaseA-13ECP26-F (5'-

CACACACATATGAAAGAAACCGCGGCGGCGAAATTTGAACGTCAGCACATGAG CCTGAAT CCACCGCGTTGCAATCAGATGATGATGAAAAGCCGTAATCTG-3') and RACOHINDR (5'-CACACAAAGCTTTTATTACACGCTCGCATCAAAATGCAC-3'). The 99-bp forward primer was designed so as to replace the 12-residue loop 1 of RNase A (D_{14} SSTSAASSSNY₂₅) by the 6-residue loop 1 of ECP (S_{17} LNPPR₂₂ (ECP numbering)). An additional TAA stop codon was also engineered in the reverse primer. The recombinant RNase A_{ECP} gene was digested with *NdeI* and *Hind*III, subcloned into *NdeI/Hind*III-digested pET22b (+) and transformed into *E. coli* XL10-Gold cells (Stratagene, La Jolla, CA). Vector pET22b (+) was obtained from an *NdeI/Hind*III digestion of the pBXR construct (39), which was a gift from Professor Ronald T. Raines (University of Wisconsin, Madison, WI). Cloning of the codon-optimized RNase A_{ECP} gene was designed so as to remove the original pELB leader

sequence present in the pBXR construct. Colonies were individually picked after selection on a Luria-Bertani (LB) medium containing 100 μ g/mL ampicillin, and the sequence of the RNase A_{ECP} gene was confirmed by DNA sequencing. For protein expression purposes, the resulting pET22b(+)-RNase A_{ECP} construct was transformed into *E. coli* BL21 (DE3) cells (Lucigen Co., Middleton, WI).

Protein expression

Isotopically ¹⁵N- or ¹⁵N-¹³C-labeled RNase A_{ECP} was expressed and purified according to the following protocol based on previously published methods (39). All chemicals were obtained from Sigma (St. Louis, MO), except for isotopes, which were obtained from Cambridge Isotope Laboratories (Andover, MA). All cell growths were performed in M9 minimal medium supplemented with MEM vitamins (Invitrogen, Carlsbad, CA), trace metals, 2.5 g/L of ¹⁵NH₄Cl, 4 g/L of ¹³C-glucose (for ¹⁵N-¹³C-labeled enzymes) or ¹²C-glucose (for ¹⁵N-labeled enzymes) and 100 µg/mL ampicillin. Overnight cultures of BL21(DE3)/pET22b(+)-RNase A_{ECP} grown in M9 media were used to inoculate a total of 3 liters of M9 media at 37° C. These were grown until the $A_{600nm} = 0.8$. Protein expression was then induced by the addition of IPTG to a final concentration of 1 mM, the cultures were grown for an additional 3 hours and harvested by centrifugation (4000 rpm, 20 min, 4°C).

Cell lysis and RNase A_{ECP} refolding

The *E. coli* cell pellet was resuspended in osmotic shock lysis buffer (20 mM Tris, 1 mM EDTA, pH 8.0) and the cells were sonicated on ice for 5×40 s intervals using an S-450A ultrasonic cell disruptor from Branson Ultrasonics Co. (Danbury, CT). The sonicated cell pellets were centrifuged for 1 hour at 20,000 *g*. The pellet from this centrifugation step (which contains the insoluble RNase A_{ECP} chimera) was resolubilized in 400 mL of urea solubilization buffer (20 mM Tris, pH 8.0, 1 mM EDTA, 400 mM NaCl, 6 M Urea, 20 mM DTT) and briefly sonicated again to resuspend cell debris. This was allowed to stir overnight at room temperature. Insoluble debris was removed by an additional centrifugation step (4°C, 45 minutes, 20,000 *g*). The soluble proteins were refolded by performing three sequential 24-h dialysis steps at 4° C against a refolding buffer (20 mM Tris, pH 8.0, 100 mM NaCl, 1 mM reduced glutathione, 0.2 mM oxidized glutathione, 0.5 mM PMSF).

Protein concentration and purification

The supernatant from dialysis was collected, centrifuged (4° C, 30 minutes, 20,000 g), filtered through a 0.22 µm membrane filter and concentrated down to ~30–40 mL using an Amicon 402 stirred-cell concentrator and a 3,000 NMWL regenerated cellulose ultrafiltration membrane from Millipore (Billerica, MA). The concentrated supernatant was dialyzed overnight against 20 mM Tris (pH 8.0) and loaded on a 10 mL prepacked anion-exchange HiTrap Q HP column (GE Healthcare, Piscataway, NJ) using an ÄKTA FPLC system preequilibrated with the same buffer. The flow-through was collected and further purified with a 1-mL Mono S 5/50 GL cation-exchange column (GE Healthcare) and RNase A_{ECP} was eluted by a salt gradient using a 20 mM Tris (pH 8.0) buffer containing 1 M NaCl. The elution peak of RNase A_{ECP} was typically observed at a conductivity of 12-14 mS/cm. Fractions containing the pure protein were combined and concentrated to ~350 µl using a 3,000 NMWL Centricon® filter device (Millipore). Finally, the buffer was changed to 5 mM MES-NaOH, 7 mM NaCl, 0.01 % NaN₃, 5 % D₂O, pH 6.4 by performing multiple dilution/concentration steps with the same Centricon filter device. Protein purity was estimated by SDS-PAGE to be higher than 98 % and protein concentration was calculated by UV absorbance using $\varepsilon_{277.5 \text{ nm}} = 9800$ $M^{-1}cm^{-1}$ (40).

Solution NMR Experiments

All NMR experiments were performed at 298 K as calibrated with a standard methanol sample (41) on Varian 11.7 T, 14.1 T and/or 18.8 T NMR instruments equipped with triple-resonance probes and pulsed-field gradients. A 160 µM ¹⁵N-labeled RNase A_{ECP} sample was used to perform all the relaxation experiments and the 3'-CMP titrations experiments. A 410 μ M ¹⁵N-¹³C-labeled RNase A_{ECP} sample was used for all the assignment experiments. Backbone and side chain resonances were assigned by using standard multidimensional experiments (41). The kinetics of the 3'-CMP binding interaction with RNase AECP were measured by titration of 3'-CMP as described (3,29), where ¹⁵N-HSQC spectra were recorded for titration points corresponding to 3'-CMP/RNase A_{ECP} mole ratios of 0, 0.174, 0.393, 0.691, 1.31, 2.71, 6, and 12. NMR line-shape analysis was performed using the software LineShapeKin (version 3.1) developed by Professor Evgenii L. Kovrigin (Medical College of Wisconsin, Milwaukee, WI, http://lineshapekin.net/) (7). Because it is difficult to obtain true estimates of fully formed protein-ligand complexes when binding constants are not very tight, titrations for RNase A_{ECP} and H48A were considered incomplete and titration end points were adjusted manually to optimally fit models to the experimental data. The 3'-CMP titrations were performed as ¹H-¹⁵N sensitivity-enhanced HSQC experiments (42-45) at 11.7 T using spectral widths (points) of 1600 Hz (256) and 7000 Hz (8192) in the t1 and t2 dimensions respectively.

Relaxation compensated Carr-Purcell-Meiboom-Gill (rcCPMG) experiments (46) were performed at 18.8 T and 14.1 T on ¹⁵N-labeled wild-type and mutant RNases. The NMR relaxation data were typically acquired with spectral widths in the t₁ and t₂ dimensions of 2700 and 10000 Hz with a digital resolution of 0.1 and 0.2 points/Hz respectively. The proton carrier frequency was coincident with the H₂O resonance and the ¹⁵N frequency was placed in the middle of the amide region at 120 ppm. Spin-relaxation rate constants at each CPMG pulse repetition time (τ_{cp}) were acquired in a constant time manner (47) using a total relaxation delay of 40 ms. Transverse Relaxation rates were determined for CPMG delays (τ_{cp}) = 0.625, 0.714 (× 2), 1.0, 1.25, 1.67, 2.0, 2.50 (× 2), 3.33, 5.0, and 10 ms. The individual spectra were acquired as an interleaved three-dimensional experiment in which the two-dimensional planes were extracted and peak intensities determined from a 3 × 3 grid using in-house written software. The CPMG dispersion data were fit with the fast-limit equation (48):

$$R_2(1/\tau_{cp}) = R_2^0 + \varphi_{ex}/k_{ex} [1 - 2\tanh(k_{ex}\tau_{cp}/2)/(k_{ex}\tau_{cp})]$$
(1)

in which $\phi_{ex} = p_A p_B \Delta \omega^2$, $p_{A/B}$ are the equilibrium site populations for the two (A and B) magnetically distinct conformers, $\Delta \omega$ is the chemical shift difference between conformations A and B, k_{ex} is the is the exchange rate constant and is the sum of the forward and reverse rate constants, R_2^0 is the 'exchange-free' transverse relaxation rate and τ_{cp} is the time delay between 180° CMPG pulses. Amino acid residues were considered for further analysis if the difference in measured R_2 ($\Delta R_2(1/\tau_{cp})$) values at fast ($\tau_{cp} = 0.625$ ms) and slow ($\tau_{cp} = 10$ ms) pulsing rates was greater than 1.5 s⁻¹. The parameter α (49) that determines the chemical exchange time regime was estimated from

$$\alpha = \left(\frac{B_{02} + B_{01}}{B_{02} - B_{01}}\right) \left(\frac{R_{ex2} - R_{ex1}}{R_{ex2} + R_{ex1}}\right) \tag{2}$$

in which $B_{01(2)}$ refers to the static magnetic field strength, 14.1(18.8) T respectively and $R_{ex1(2)}$ is the exchange contribution due to conformational exchange and is determined from fits of equation (1) to the CPMG dispersion data.

Differential multiple-quantum relaxation experiments were performed as described by Kloiber and Konrat (50) in an identical manner to earlier work on RNase A (28). Peak intensities for a 10 ms relaxation delay (T_c) were measured for two experiments, a reference (I_{auto}) and a cross-relaxation (I_{cross}) experiment. The intensities of each experiment represent the decay of multiple quantum coherence ($2I_xS_x$) and the conversion of $2I_xS_x$ to $2I_yS_y$ respectively. The conformational exchange modulated isotropic chemical shifts differentially affects these terms. The ratio of peak intensities for the relaxation of $2I_xS_x$ (I_{cross}) and relaxation of $2I_xS_x$ to I_yS_y (I_{auto}) were used to determine the differential multiple quantum relaxation rate ΔR_{MQ} from:

$$I_{cross}/I_{auto} = \tanh\left(\Delta R_{MQ}T_c/2\right)$$
(3)

RESULTS

Chemical shift assignments

Both H48A and RNase A_{ECP} are well-folded enzymes as evidenced by the chemical shift dispersion in Figure 2A–C. Upon mutation to form the H48A enzyme and the RNase A_{ECP} chimera, ¹H and ¹⁵N chemical shifts are perturbed for several similar regions in these proteins (Figure 2). For H48A, the largest changes in chemical shifts are located in loop 1 and in β -sheet 4, with more minor effects observed for the $\beta 1/\alpha 3$ region and $\beta 5$ residues (Figure 2). For the loop 1 chimera, the regions that are most significantly affected are $\beta 1/\alpha 3$ and $\alpha 2$, with more minor perturbations occurring in $\beta 4$ and $\beta 5$ (Figure 1). Not surprisingly, the residues Met13, Asp14, and Ser15, which are just N-terminal to the site of loop swapping display the largest changes in chemical shifts in RNase A_{ECP} (Figure 2).

3'-CMP Ligand Interaction

To further investigate the kinetics of the rate-limiting product release step of this reaction, we performed a steady-state NMR line shape analysis (3) by following the titration of the product analog 3'-CMP in WT RNase A, RNase A_{ECP} and in the previously-characterized point mutant H48A (28). In addition to following the various chemical shift effects observed upon ligand binding, this technique allows for the calculation of the dissociation constant (K_d) and the dissociation rate constant (k_{off}) of 3'-CMP from RNase A.

A series of two-dimensional HSQC experiments were used to follow the ¹H-¹⁵N chemical shifts observed for each enzyme upon progressive increases in [3'-CMP] (Figure 3 and Figure 4). As shown by the chemical shift variations observed in Figure 3, binding of 3'-CMP to WT alters the chemical environment of a number of residues throughout the enzyme, mainly in regions encompassing residues 10-15, 40-50, 80-90, 95-102, and 119-123. The protein-wide average chemical shift changes per amino acid residue upon binding of 3'-CMP are 0.071, 0.053, and 0.040 ppm for WT, RNase A_{ECP} , and H48A respectively. Many of the residues displaying the largest chemical shift changes ($\Delta > 0.15$ ppm) between the apo (mole ratio, 3'-CMP:RNase A = 0) and 3'-CMP bound (mole ratio, 3'-CMP:RNase A = 12) forms for WT are either in direct contact or in close proximity to the ligand (Lys7, Gln11, Lys41, Val43, Asn44, His119, Asp121 and Ala122) (51). A number of residues located farther away from the CMP binding site also display chemical shift variations greater than 0.15 ppm in WT, including Phe46, Asp83, Arg85, Glu86, Thr100 and Gln101, all of which are located in the adjacent β 1sheet that separates the active site cavity from loop 1. In contrast to WT, the chemical shift differences observed between the unbound and the 3'-CMP-bound forms of both RNase A_{FCP} and mutant H48A are significantly smaller (Figure 3). For example, the only residues displaying a $\Delta > 0.15$ ppm in these two mutant enzymes are Val43, Asp83, His119, Asp121 for RNase A_{ECP} and Val43, Asn44 and Glu86 for H48A, demonstrating that mutation of either loop 1 or His48 alters the mode of ligand interaction with RNase A.

The extent of the attenuation of the chemical shift changes upon binding of 3'-CMP is illustrated in Figure 4. The magnitude of the chemical shift changes is greatly diminished in both RNase A_{ECP} and H48A, yet the direction of the changes in chemical shift for both is similar to WT RNase A.

Among all the residues with chemical shifts in the 3'-CMP titration experiment, only a restricted set (Gln11, His12, Arg39, Lys41, Phe46, Ser75, Tyr97 and Ser123) showed significant shifts and were also sufficiently resolved for line shape quantitation in all three enzymes (Figure 5). Whether NMR line shape analysis was independently performed on individual residues or simultaneously fitted using a global, two-state model, there were few variations in the calculated binding constants, indicating that all residues likely report on the same binding event. 3'-CMP binding values are therefore reported for a global fit of residues and are presented in Table 1.

The 3'-CMP dissociation constant (K_d) for WT RNase A determined from these NMR titrations is 74 ± 2 µM. This value is indicative of tight binding between 3'-CMP and WT RNase A and is in good agreement with dissociation constants previously determined for this ligand (11, 18,53). In contrast, the resulting 3'-CMP dissociation constants (K_d) for RNase A_{ECP} and mutant H48A are respectively 300 ± 16µM and 895 ± 50 µM. Replacement of the distal 12residue loop 1 of RNase A by the shorter 6-residue loop 1 of ECP in the RNase A_{ECP} chimera reduces binding affinity for 3'-CMP ~4-fold relative to WT. Interestingly, despite the fact that it is more than 18 Å away from the catalytic center and that it is in direct contact with loop 1 (Figure 1), mutation at position His48 reduces the binding affinity to a greater extent (~12-fold relative to WT) than does swapping the entire loop 1 sequence.

The product release (k_{off}) is the rate-limiting step of the reaction in WT RNase A (29,54–56). In previous work it was demonstrated that k_{off} and the conformational exchange rate constant k_{ex} were equivalent (28) and in combination with other experimental evidence suggested that motion in RNase A, particularly loop 1, was part of the rate-limiting step (3,13,27–29). As shown in Table 1, the NMR line shape analysis of the 3'-CMP titration series yields a WT k_{off} value of ~1200 s⁻¹, a result in good agreement with previous calculations (3). Replacement of loop 1 or mutation of H48 in RNase A has a significant effect on the rate of product release, as k_{off} values for 3'-CMP are decreased ~10-fold in both RNase A_{ECP} ($k_{off} = 96 \pm 9 \text{ s}^{-1}$) and mutant H48A ($k_{off} = 142 \pm 19 \text{ s}^{-1}$) relative to WT (Table 1).

¹⁵N Backbone Dynamics

A relaxation-compensated CPMG dispersion experiment was performed on apo ¹⁵N-labeled RNase A_{ECP} for comparison with WT and H48A to investigate the effects of the loop 1 swap on the important millisecond motions of the backbone amide groups. As evidenced by the characteristic difference in R_2 values measured at increasing values of the CPMG 180° pulse delays (τ_{cp}), a number of residues display conformational exchange throughout the structure of RNase A_{ECP} (Figure 6). Among the well-resolved ¹H-¹⁵N resonances that show good signal-to-noise values, twelve residues experience a conformational exchange process with a ΔR_2 (1/ τ_{cp}) value > 1.5 s⁻¹ (Val43, Cys58, Asn62, Ala64, Cys65, Lys66, Gln69, Thr70, Asn71, Cys72, Asp121 and Ser123). A few of those residues are amenable to quantitative analysis (Ala64, Cys65, Gln69 and Asn71) and are shown in Figure 6B. These dynamic positions are mainly clustered in a surface loop adjacent to the active site of RNase A (loop 4), which contains the highly conserved P0 and B2 subsite residues K66 and N71, respectively (57). Interestingly, those residues also display similar dispersion profiles both in WT RNase A and in the previously characterized mutant H48A (Figure 6B) (28).

Comparison with WT dynamics also shows a number of residues displaying loss of CPMG relaxation dispersion, indicative of a loss of μ s-ms protein motion for RNase A_{ECP}. Among

Page 7

Ser80, Thr82, Asp83, Thr100 and Gln101) show no NMR evidence of any conformational exchange motion in RNase A_{FCP} (Figure 6B). Surprisingly, every residue that displays a flat dispersion profile in RNase A_{FCP} also exhibits the same flat profile in the H48A mutant (Figure 6B), indicating that both alterations to RNase A result in the loss of a similar conformational exchange process. The residues common to both RNase AECP and WT that experience measurable CPMG dispersion curves (Cys65, Gln69, Asn71, and Asp121; Figure 6B) also exist in similar exchange time regimes as estimated from the α parameter (49), with the exception of Asn71 (Supporting Information). Residues 65, 69, and 121 all have α values greater than 1.2 indicating conformational exchange that is fast on the chemical shift time scale. For Asn71 in RNase $A_{ECP} \alpha = 0.7 \pm 0.1$ and is 1.6 ± 0.1 WT RNase A. The lower value of α for Asn71 is most likely due to underestimation of Rex at 18.8 T (Supporting Information).

The flat ¹⁵N dispersion curves for both H48A and RNase A_{ECP} indicate that in these two proteins the amide nitrogen atoms are not moving, do not experience significant $\Delta \omega$ values, and/or exists in a highly skewed equilibrium (> 0.995:0.005). To address this further, differential multiple quantum relaxation rates (ΔR_{MO}) (50) were measured (Supporting Information). The ΔR_{MO} values depend on chemical shift differences experienced by both ¹⁵N and ¹H and can therefore provide an additional assessment of the presence of millisecond motions. For all assignable residues, the ΔR_{MO} rates mirror those of the ¹⁵N single quantum relaxation dispersion measurents. Residues that have measurable ¹⁵N-single quantum dispersion curves also have non-zero differential multiple quantum relaxation rates. Therefore from the perspectives of the ¹H and the ¹⁵N nuclei Lys37, Lys41, Phe46 Val47, Thr82, Asp83, Thr100 and Gln101 are no longer flexible on the µs-ms timescale in H48A and RNase A_{ECP} (Supporting Information).

DISCUSSION

Loops are typically tethered to the main body of the protein via rigid secondary structure elements and therefore can be envisioned as structural units moving on protein hinges. This simplified view of loop function often explains a great deal regarding their biochemical role (8,58,59). This inherent flexibility of loop regions has been harnessed to provide proteins and enzymes with a mechanism to alter ligand-binding specificity (60-67). In an extreme example, a loop distant from the protease active site modulates the differential specificity between trypsin and chymotrypsin (63). Loop regions in RNase A have also been the subject of loop-swapping experiments. Benner and coworkers exchanged three loops from the RNase homolog Angiogenin into bovine pancreatic RNase A (60). Loop residues 63-74 contact the purine portion of the substrate. When these residues from Angiogenin were placed into RNase A, the resulting enzyme had purine substrate specificity resembling that of Angiogenin, not that of RNase A. Unlike loop 1 in ECP, which is six residues long, loop 1 in Angiogenin is nine residues long. Exchange of the longer Angiogenin loop 1 into RNase A had a modest 2-fold effect on catalytic activity (60). Here we show that the flexibility offered by the longer loop 1 in RNase A is closely associated with the product release step despite not directly interacting with substrate or product.

We observe an important similarity between the millisecond dynamical behavior of RNase A_{ECP} and the previsouly characterized mutant H48A (28). Both mutants display flexible residues involved in conformational exchange at similar atomic positions throughout the enzyme (Figure 6 and Figure 2 of ref. (68)), and both enzymes display two independent clusters of flexible residues that are affected in a similar manner when His48 or loop 1 are mutated. The first cluster is close to the active site of RNase A and is centered around the flexible loop 4 (Figure 6, red spheres). This cluster is insensitive to solvent isotope effects in H48A and displays WT-like motional behavior in both mutants. The second cluster of residues, centered about the adjacent β 1-sheet (formed by β 1, β 4 and β 5 strands) (Figure 6A, blue spheres), experiences an important kinetic solvent isotope effect in WT RNase A, which is absent in H48A. In addition, this second group of residues display significantly reduced dynamics in both RNase A_{ECP} and in H48A noted by the lack of CPMG dispersion. The fact that His48 is hydrogen bonded to loop 1 through T17 (Figure 1B) and that both mutants show a similar motional behavior upon replacement of either one of these structural elements suggests that both loop 1 and His48 are dynamically linked by a network of coupled motions on the catalytically relevant millisecond time scale. It is reasonable to assume that these concerted motions are at least partly propagated through Thr17 (Figure 1B), which was shown to display conformational exchange in WT but not in H48A (68).

The fact that mutation of His48 or swapping of loop 1 affects the dynamics of many residues of the β 1-sheet in a similar fashion relative to WT (*e.g.* Thr82, Asp83, Thr100, Gln101) (Figure 6) also demonstrates that the β 1-sheet, His48 and loop 1 are all part of a global concerted dynamical event that propagates between the β 1, β 4 and β 5 strands through His48 to loop 1 on the time scale of catalytic turnover. Mutating either His48 or loop 1 results in a similar loss of motion in the β 1-sheet, further demonstrating the importance of His48 in acting as a relay of the dynamical changes from loop 1 to the the β 1-sheet, and/or vice versa. Threonine 17 and Thr82 form hydrogen bonds to His48 and may be an important mechanism of connectivity of loop 1 dynamics to the active site of RNase A. Interestingly, we also observe a significant correlation between residues of the second cluster (Figure 6) and many residues showing altered dynamics upon 3'-CMP binding.

Whether the millisecond dynamics of RNase A_{ECP} and H48A are retained with respect to WT (as with residues Gln69 and Asn71) or significantly reduced (as with residues Thr82, Asp83 and Gln101), these two mutant enzymes are affected in the same way at the same amino acid positions. This result further supports the existence of a network of coupled motions in RNase A that would propagate dynamics from loop 1 to His48 and other residues of the adjacent β 1-sheet, extending all the way to the ligand in the active site of the enzyme. Additional experiments are currently underway to further address this hypothesis.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

ND acknowledges a postdoctoral fellowship from the Fonds Québécois de la Recherche sur la Nature et les Technologies (FQRNT) from the Government of Québec. EDW acknowledges NIH training grant 5T32GM0082383. JPL acknowledges NSF awards MCB-0236966 and MCB-0744161.

The authors thank Professor Evgenii L. Kovrigin (Medical College of Wisconsin, Milwaukee, WI) for helpful discussions. This manuscript is dedicated to Professor Fred Richards.

ABBREVIATIONS

RNase A (or WT)

wild-type ribonuclease A

ECP

eosinophil cationic protein

EDN

eosinophil-derived neurotoxin

RNase A_{ECP} (or chimera)
RNase A with loop 1 from ECP

H48A

RNase A (H48A)

3'-CMP

cytidine-3'-monophosphate

rcCPMG

relaxation compensated Carr-Purcell-Meiboom-Gill

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Figure 1.

Structural comparison of bovine ribonuclease A (RNase A) and human Eosinophil Cationic Protein (ECP). A) Superposition of RNase A (blue structure, PDB code 1FS3 (35)) and ECP (red structure, PDB code 1DYT (33)). H12 highlights the position of the active site relative to the position of His48 and loop 1 in RNase A. B) Zoomed view of a few atomic interactions between His48 and loop 1 in RNase A. Note the longer loop 1 in RNase A

(D₁₄SSTSAASSSNY₂₅) relative to the shorter loop 1 of ECP (S₁₇LNPPR₂₂, ECP numbering). C) Primary sequence alignment of RNase A, ECP and EDN. Active site positions Gln11, His12, Lys41, His119 and Asp121 are marked with stars while cysteine residues forming the strictly conserved four disulfide bridges are represented by arrows. Loop 1 and position 48 are boxed. Alignment was performed with TCoffee Expresso (36) using PDB coordinates 7RSA (32), 1DYT (37) and 1GQV (38).



Figure 2.

Chemical shift changes caused by mutation. ¹H-¹⁵N HSQC spectra for (A) wild-type, (B) RNase A_{ECP}, and (C) H48A enzymes. The spectra were acquired at 14.1 T and 298K, pH = 6.4. In (D) chemical shift differences for WT vs. H48A (blue) and WT vs. RNase A_{ECP} are shown as a function of amino acid sequence. Gray boxed areas depict protein regions with larger than average chemical shift disturbances. (Δ (ppm) = [($\Delta\delta^2_{HN} + \Delta\delta^2_N/25$)/2]^{1/2}) in which δ is the difference in chemical shift between the two proteins for ^NH and ^HN nuclei (52).

Doucet et al.



Figure 3.

¹H-¹⁵N chemical shift differences upon 3'-CMP binding to WT RNase A, RNase A_{ECP}, and H48A. (A) ¹H-¹⁵N chemical shift differences Δ (ppm) mapped on the primary sequence of WT, RNase A_{ECP}, and H48A. The value of Δ was calculated between a ligand-free ¹H-¹⁵N HSQC (mole ratio, 3'-CMP/Enzyme = 0) and a 3'-CMP-bound ¹H-¹⁵N HSQC (mole ratio, 3'-CMP/Enzyme = 12) according to the following equation: (Δ (ppm) = [($\Delta\delta^2$ _{HN} + $\Delta\delta^2$ _N/25)/2]^{1/2}) (52). (B) Mapping of Δ (ppm) on the three-dimensional structure of RNase A (PDB code 1RPF (51)). The white to red gradient corresponds to Δ values ranging from 0 ppm (white) to > 0.15 ppm (dark red). (C) Same view as in (B) showing residues with Δ > 0.15 ppm as green spheres. 3'-CMP is shown as orange sticks representation.



Figure 4.

Kinetics of 3'-CMP binding to WT RNase A, RNase A_{ECP} and mutant H48A. Titration of 3'-CMP is followed by monitoring the changes in backbone chemical shifts observed in ¹H-¹⁵N-HSQC spectra for WT RNase A, RNase A_{ECP} and mutant H48A. Titration points are shown for residues Gln11, His12, Phe46 and Asp121 for [3'-CMP]:[RNase A] mole ratios of 0 (red) 0.174 (orange), 0.393 (yellow), 0.691 (green), 1.31 (blue), 2.71 (purple), 6 (magenta), and 12 (cyan). Arrows indicate the direction of resonance shift with increasing [3'-CMP]. The magnitude of 3'-CMP binding-induced chemical shifts is smaller in mutant H48A than in RNase A_{ECP} .



Figure 5.

3'-CMP titration of RNase A. The titration curves of residue His12 are shown for WT, RNase A_{ECP} and H48A. Fitted line shapes (red) are superimposed on the experimental NMR data (blue).



Figure 6.

Conformational dynamics in RNase A. (A) Residues in RNase A_{ECP} undergoing conformational exchange with a ΔR_2 (1/ τ_{cp}) greater than 1.5 s⁻¹ are plotted on the structure of RNase A (PDB code 7RSA (32)). Red spheres = similar rate constants (k_{ex}) in WT and RNase A_{ECP} ; Blue spheres = Decreased k_{ex} values in RNase A_{ECP} relative to WT. H12 and H48 are shown as yellow sticks and loop 1 is shown in cyan. B) CPMG dispersion curves for WT RNase A (black circles), RNase A_{ECP} (red squares) and mutant H48A (blue diamonds). ¹⁵N relaxation dispersion curves are shown for two residues displaying similar conformational exchange in the three enzymes (Gln69 and Asn71), and three residues displaying significantly affected k_{ex} in both RNase A_{ECP} and mutant H48A (Thr82, Asp83 and Gln101). Y115 is represented

as a motionless control. Fitted lines to data points are from single-field fits at 14.1 T using equation (1).

Table 1 3'-CMP binding constants for WT RNase A, RNase A_{ECP} and mutant H48A.

Enzyme	<i>K</i> _d (μM)	$k_{\rm off}({ m s}^{-1})$	$k_{on} \left(M^{-1} s^{-1} \right)^a$
WT Rnase A	74 ± 2	1155 ± 120	$1.56 \times 10^7 \pm 1.67 \times 10^6$
RNase A _{ECP}	300 ± 16	96 ± 9	$3.20 \times 10^5 \pm 3.45 \times 10^4$
RNase A (H48A)	895 ± 50	142 ± 19	$1.59 \times 10^5 \pm 2.30 \times 10^4$

 a k_{on} determined from k_{off}/K_d.