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# UNIVERSITY OF ALBERTA

# THE MOLECULAR STRUCTURES OF TWO CYTOTOXIC RIBONUCLEASES OF VERTEBRATES, ONCONASE AND EOSINOPHIL-DERIVED NEUROTOXIN

STEVEN C. MOSIMANN C

# A THESIS SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF DOCTOR OF PHILOSOPHY

DEPARTMENT OF BIOCHEMISTRY

EDMONTON, ALBERTA FALL 1995



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# FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research for acceptance, a thesis entitled "The Molecular Structures of two Cytotoxic Ribonucleases of Vertebrates, Onconase and Eosinophil-Derived Neurotoxin" submitted by Steven C. Mosimann in partial fulfillment of the requirements for the degree of Doctor of Philosophy.

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Date: October 6, 1995

#### ABSTRACT

Onconase<sup>TM</sup> (P-30 Protein; ONC) is a cytotoxic, pyrimidine-specific ribonuclease of Rana pipiens with therapeutic properties in vivo. Orthorhombic single crystals of purified ONC (space group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>) are obtained from an (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> solution at pH 4.5 and grow as rosettes: a = 40.53 (8), b = 69.64 (9), c = 32.52 (8) Å. The crystal structure of ONC has been determined by multiple-isomorphous replacement, using three heavy atom derivatives and refined at 1.7 Å resolution. The comparative molecular model of ONC based upon the known structure of ribonuclease A (RNase A) and the subsequent crystal structure demonstrate ONC can adopt an RNase A-like polypeptide fold. However, there are many large conformational differences in the large loop structures of ONC, the comparative molecular model and RNase A. Unlike RNase A, the N-terminal residue (pyroglutarnyl) of ONC hydrogen bonds to Lys9N<sup>\zeta</sup> at the active site of the enzyme. Recombinant Onconase (rONC) having an N-terminal sequence fMet\_1Glu1Asp2... has been expressed, purified and shown to lack significant cytotoxic and ribonucleolytic activity. The isomorphous rONC structure has been determined and refined at 2.2 Å resolution to examine the structural basis of this inactivity. Large conformational differences in the ONC and rONC structures are confined to the N-terminal residues and only slightly perturb the conformation of the active site residues. Molecular models of ONC and rONC-d(UpG) complexes have been constructed and attribute the lack of ribonucleolytic activity of rONC to differences in the shape and electrostatic nature of the ribose binding site (R2). The modeling implicates Lys33 in the uridine base specificity of Onconase and suggests Ser54, Asn56 and Glu91 are responsible for binding the purine base.

In parallel work, the X-ray crystallographic structure of the related recombinant eosinophil-derived neurotoxin (rEDN) has been determined and refined at 1.83Å resolution. This structure reveals the conformation of several novel insertions unique to

EDN and the presence of a second sulfate anion binding site. The additional anion binding site may correspond to the P-1 phosphate binding subsite for oligoribonucleotide substrates.

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#### LIST OF ABBREVIATIONS AND SYMBOLS

Å Angstrom (10<sup>-10</sup> meters)

B factor Isotropic thermal motion factor in  $Å^2$  (B= $8\pi u^2$ )

B<sub>0</sub>, B<sub>1</sub>, B<sub>2</sub> ... Base binding subsites

CMP, Cp Cytidine 3' monophosphate
CpA Cytidine 3',5' adenylate
CpG Cytidine 3',5' guanylate

Da Dalton (1/12 mass of <sup>12</sup>C nucleus)
d(CpA) Deoxycytidine 3',5' deoxyadenylate
d(UpG) Deoxyuridine 3',5' deoxyguanylate

d(TpTpTpTp) Deoxythymidylate tetramer

EDN Eosinophil-derived neurotoxin

 $F_c$  Calculated structure factor amplitude  $F_o$  Observed structure factor amplitude

I X-ray scattering intensity

iC<sub>50</sub> Concentration required for 50% inhibition

kV Electrical potential in kilovolts mA Electrical current in milliamperes

f<sub>Met</sub> Formyl methionine

 $\begin{array}{ll} \text{MIR} & \text{Multiple isomorphous replacement} \\ \text{M}_{r} & \text{Molecular weight expressed in Daltons} \end{array}$ 

ONC Onconase, P-30 Protein Phosphate binding subsites

P-30 Protein Onconase

Pyr Pyroglutamyl, 5-oxo-2-pyrollidine carboxylate

 $R_0, R_1, R_2 \dots$  Ribose binding subsites

R factor Crystallographic residual minimized during refinement

 $(R=\Sigma||F_o|-|F_c||/\Sigma|F_o|)$ 

 $R_m$  Merging R factor  $(R_m = \sum_{hkl} \sum |I_{ave} - I_{obs}|/\sum |I_{ave}|)$ 

rEDN Recombinant eosinophil-derived neurotoxin rms, rmsd Root mean square, root mean square deviation

RNase Ribonuclease

rONC Recombinant Onconase

Up Uridine 3' monophosphate

Vm Specific volume in Å<sup>3</sup>/Dalton

 $\lambda$  Wavelength of electromagnetic radiation

σ Standard deviation

#### CHAPTER 1

#### INTRODUCTION

Ribonucleases (RNases) can be cytotoxins. In bacteria, fungi, higher plants, amphibians and mammals, there exist ribonucleases that specifically bind to and enter the cytosol of a wide variety of target cells where they degrade RNA species, ultimately killing the target cell. The bacterial cytotoxins, colicin E3 and cloacin DF13, function in interstrain competition and target ribosomal RNA (rRNA: DeGraaf & Oudega, 1986). The fungal toxin α-sarcin specifically inactivates eucaryotic ribosomes and may be a virulence factor for Aspergillus (Chan et al., 1990). The S2 glycoproteins of Nicotiana alata function in the prevention of self-pollination and degrades the rRNA of incompatible pollen tubes (Murfett et al., 1994). The cytotoxic RNases of amphibians and mammals are homologous proteins belonging to the pyrimidine-specific RNase superfamily of vertebrates. They include P-30 Protein (Onconase<sup>TM</sup>, ONC) and sialic-acid binding lectins of frog eggs, eosinophil-derived neurotoxin (EDN), eosinophil cationic protein (ECP) and bovine seminal RNase. At present, we do not know how the vertebrate enzymes recognize and enter target cells nor do we know the precise intracellular target of their RNase activity.

The cytotoxic members of the RNase superfamily of vertebrates are of medical interest. The amphibian enzymes have potential therapeutic applications and favorable immunogenic properties (Mikulski et al., unpublished results). Both EDN and ECP have diagnostic applications and play roles in many human pathologies (Youle et al., 1993 and references therein). As a consequence, the ability of these enzymes to specifically bind and enter susceptible cells and their individual ribonucleolytic activities are of interest. In addition, within the RNase superfamily of vertebrates some members are cytotoxic and other are not. Comparison of cytotoxic and non-cytotoxic members of the superfamily offer the possibility of identifying functionally important differences in these enzymes. For

example, which residues contribute to the individual ribonucleolytic specificities of the different enzymes? Which residues or segments of polypeptide chain adopt significantly different conformations in these enzymes? Which residues or segments are responsible for their different biological functions? To address these issues and questions, we determined and analyzed three-dimensional structures of ONC, recombinant ONC (rONC) and recombinant EDN (rEDN). The long term goals of these collaborative projects include the molecular engineering of cytotoxins with enhanced or altered specificity and activity and the development of specific inhibitors of EDN and ECP.

# Pyrimidine-specific Ribonuclease Superfamily of Vertebrates

The pyrimidine specific ribonuclease (RNase) superfamily of vertebrates is referred to as the RNase superfamily, pancreatic RNase superfamily and the pyrimidine-specific mammalian alkaline and neutral RNase superfamily in various reviews (Sierakowska & Shugar, 1977; Beintema, 1987; Beintema et al., 1988). The superfamily includes reptilian enzymes (Beintema et al., 1985; Zhao et al., 1994), amphibian enzymes (Ardelt et al., 1991; Titani et al., 1987), avian enzymes (Klenova et al., 1992) and over 40 mammalian members. There are three distinct pyrimidine-specific RNases in the various tissues and body fluids of all tested mammals: the secretory (eg. bovine RNase A) and non-secretory (eg. EDN) RNases defined by Sierakowska & Shugar (1977) and the angiogenins (Strydom et al., 1985). The distinct groups of mammalian enzymes have arisen from gene duplications after the divergence of the mammals and the non-mammals (Beintema et al., 1988) some 150 million years ago and are distant relatives of the reptilian, amphibian and avian enzymes (Beintema et al., 1988).

These enzymes range from 104 amino acids (ONC) to 140 amino acids (bovine brain RNase) in size, have a net positive charge at neutral pH and the mammalian members of the superfamily are glycoproteins. Table 1.1 lists several biochemical properties of representative mammalian members of this superfamily and ONC. Table 1.2 is an amino

Table 1.1

Pyrimidine-specific RNase superfamily of vertebrates

			<del></del>
124	13 753	9.2	digestion of RNA in gut
125	14 615	9.6	
134	15 463	10.4	host defense unknown (perhaps differentiation)
104	11 845	9.4	
	134	134 15 463	134 15 463 10.4

<sup>†</sup> RNase A and angiogenin are the bovine enzymes, EDN is isolated from human serum and ONC is from Rana pipiens oocytes.

<sup>‡</sup> Estimated from the amino acid composition.

acid sequence alignment of representative members of the pyrimidine-specific RNase superfamily of vertebrates. There are 19 residues conserved in the five amino acid sequences of Table 1.2. They include six cysteine residues that form three conserved disulfide bridges, three catalytic residues (RNase A: His12, Lys41, His119) and five residues from two \(\beta\)-strands that form the active site (RNase A: Asn44, Thr45, Phe46, Pro117, Val118), four hydrophobic residues (RNase A: Phe8, Tyr97, Val1C8), and one residue from a loop (RNase A: Ser77). Table 1.3 lists the percentage amino acid sequence identity between representative mammalian members of the superfamily and ONC. The pair-wise amino acid sequence identity between secretory RNases of different mammals, non-secretory RNases of different mammals and angiogenins of different mammals is typically 70% (Beintema et al., 1988). The relatively low sequence identity and the presence of several large deletions in the Onconase sequence alignment with RNase A originally led us to construct a comparative molecular model of ONC based upon RNase A (Chapter 2). The model addresses the following questions; Can ONC adopt an RNase Alike fold? Is the published amino acid sequence alignment correct? Can Lys9 of ONC replace the conserved Gln (RNase A: Gln14) at the active site of mammalian RNases?

# Cytotoxic Ribonucleases of the RNase Superfamily of Vertebrates

The amphibian members of the RNase superfamily include the lectins (Rana japonica, Rana catesbiana) and ONC (Rana pipiens). These enzymes are present in abundance in oocytes and the early blastomere of frog species and their biological functions are unknown. ONC was initially identified and purified based upon its therapeutic activity in several animal models of cancer (Mikulski et al., 1990). ONC has completed phase I and phase II human clinical trials in the United States as a prospective cancer treatment (Mikulski et al., 1993; unpublished results). It also possesses an anti-HIV activity (Youle et al., 1994) in vitro and displays a neurotoxic activity similar to that

Table 1.2

Amino acid sequence alignment of members of the RNase superfamily

	1	*		30	*
Seminal	kesaaak	FerqHmdsgn	spssssnyCn	lmMccrkmtq	gkCKpvNTFv
RNase A	ketaaak	FerqHmdsst	saasssnyCn	qm $M$ k $s$ r $n$ l $t$ k	drCKpvNTFv
EDN	kppqftwaqw	FetqHinmts	qqCt	naMqvinnyq	rrCKnqNTFl
Angiogenin	.aqddyryih	FltqHydakp	kgrnd.eyCf	hmMknrrlt.	rpCKpvNTFi
ONC	jdwlt	FqkkHitntr	dvdCd	niMstnlf	.hCKdkNTFi
	ααα	ααααα	αα	ααα	βββββ

		60			90
Seminal	hesladvkav	Csqkkvtckn	gqtncyqs	kStmritdCr	etgsskyp
RNase A	hesladvqav	Csqknvackn	gqtncyqs	yStmsitdCr	etgsskyp
EDN	lttfanvvnv	Cgnpnmtcps	nktrknchhs	gSqvplihCn	lttpspqnis
Angiogenin	hgnkndikai	Cedrngqpyr	gdlris	kSefqitiCk	hkggssrp
ONC	ysrpepvkai	Ckgiia.	sknvlt	tSefylsdCn	vtsrp
	β ααααααα	α	βββββ	<u> </u>	

				* 124	
Seminal	nCaYkttqve	khiiVaCggk	p	svPVHfdasv	
RNase A				yvPVHfdasv	
EDN	nCrYaqtpan	mfyiVaCdnr	dqrrdppqyp	vvPVHldrii	
Angiogenin				.lPVHfdesf	
ONC	.CkYklkkst	nkfcVtCenq		.aPVHfvgvg	sc
ββββββ βββββββ			βββββ		

A structure based amino acid sequence alignment (one letter code) of representative members of the RNase superfamily. Residues in capitals are conserved among the five sequences and catalytic residues are indicated with an asterix. Residues with  $\alpha$ -helical or  $\beta$ -strand conformations in all five structures are indicated by an  $\alpha$  (helices) or a  $\beta$  (strands), respectively. The sequence numbers are those of RNase A. The enzymes are bovine seminal RNase, bovine RNase A, human EDN, bovine angiogenin and Rana pipiens ONC. The symbol 'j' represents the N-terminal pyroglutamyl of ONC.

Table 1.3

Percent amino acid sequence identity between members of the RNase superfamily

Enzyme†	RNase A	EDN	angiogenin	ONC
RNase A	100	33	32	28
EDN		100	22	27
Angiogenin			100	27
ONC				100

<sup>†</sup> RNase A and angiogenin are the bovine enzymes, EDN is isolated from human serum and ONC is isolated from Rana pipiens oocytes.

of EDN (Newton et al., 1994). The RNases of Rana japonica and Rana catesbiana were identified as lectins that preferentially agglutinate cancer cells and specifically bind to sialylglycoproteins (Titani et al., 1987). The lectins share approximately 53% amino acid sequence identity with ONC and suggested an early model for the mechanism of ONC cytotoxicity. Perhaps, ONC binds to specific cell surface receptors and enters the cell cytosol where it degrades RNA and kills the cell. ONC exhibits saturable binding to the susceptible 9L glioma cells and its affinity is similar to its IC50, 2.5 x 10-7 M. In addition, each of the cytotoxic activities of ONC requires its ribonuclease activity (Wu et al., 1993; Youle et al., 1994; Newton et al., 1994). To further our understanding of the function and enzymatic activity of ONC, its three-dimensional structure has been determined and analyzed (Chapter 3). The structure determination also allows us to critically assess the There are several large previously constructed comparative molecular model (Chapter 2). conformational differences in a comparison of ONC, the comparative molecular model and the non-cytotoxic RNase A structures. Surprisingly, the N-terminal pyroglutamyl residue (Pyr1) folds back against the N-terminal α-helix in the crystal structure and hydrogen bonds to Lys9 at the active site of the enzyme. This suggests Pyr1 has a role in the enzymatic activity of ONC. This is significant as recombinant ONC (rONC) with the Nterminal sequence fMet\_1, Glu1, Asp2... lacks appreciable RNase and cytotexic activity (Lin et al., 1994). To determine the structural basis of the inactivity of rONC, its structure has been determined and refined at 2.2 Å resolution (Chapter 4).

An additional factor in the cytotoxicity of these enzymes is their sensitivity to the cytosolic ribonuclease inhibitor (RI). The inhibition of pyrimidine specific RNases by the cytosolic RI displays an interesting species specificity. Amphibian RNases are inhibited by the amphibian RI but not by RI of mammalian species and the mammalian RNases are only inhibited by mammalian RIs (Youle *et al.*, 1993). It has been suggested that this cytosolic inhibitor protects cells from RNases of this superfamily (Beintema, 1987).

EDN and ECP share 67% amino acid sequence identity (Gleich et al., 1986) and are

major constituents of the eosinophil granule. Eosinophils release the contents of their granules into the extracellular medium when appropriately stimulated and are a major line of defense against parasites (reviewed in Gleich & Adolphson, 1986). EDN and ECP are clinical indicators of activated eosinophils (Reimert et al., 1993) and are active against parasites in vitro (McLaren et al., 1981). They also play a role in a wide variety of human diseases characterized by excessive eosinophil levels (eosinophilia). These diseases are primarily associated with inflammatory processes and include bronchial asthma (Venge, 1993) and atopic dermatitis (Ott et al., 1994) among many others (Youle et al., 1993 and references therein). EDN and ECP are neurotoxins when injected intracerebrally (Gleich et al., 1986). In animal models the neurotoxicity is expressed as muscle rigidity, incoordination, ataxia and spasmodic paralysis that are consistent with cerebellar deficit. Neuropathological studies show Purkinje cells are selectively sensitive to these RNases (Gordon phenomenon; Durack et al., 1981). As with ONC, the neurotoxicity of EDN and ECP requires its RNase activity (Sorrentino et al., 1992). Precisely how EDN and ECP selectively kill Purkinje cells is still an open question. Unlike the amphibian enzymes, EDN and ECP are strongly inhibited by mammalian RIs (Beintema, 1987).

In parallel with our structural investigations of ONC, the three-dimensional structure of rEDN has been determined and analyzed to examine its function and enzymatic activity (Chapter 5). Of particular interest is a highly polar stretch of residues (Asp115-Tyr123) that are unique to and conserved among the non-secretory members of the superfamily.

Bovine seminal RNase is a unique member of the pyrimidine specific RNase superfamily of vertebrates. It is a covalent homodimer, has significant enzymatic activity towards single and double stranded RNA (ssRNA, dsRNA) and displays antispermatogenic and anti-tumor activities that require its RNase activity (for review, D'Alessio, 1993). It shares 81% amino acid sequence identity with the non-cytotoxic, monomeric RNase A. When seminal RNase is prepared as a monomer, it retains its

enzymatic activity towards ssRNA and dsRNA but loses its cytotoxic potential suggesting dimerization is important in its cytotoxic mechanism. Studies showing chemically cross linked dimers of RNase A posses anti-tumor activity (Bartholeyns & Zendergh, 1979) support this interpretation. How dimerization relates to the cytotoxicity of seminal RNase is unclear. Dimerization may facilitate entry into the cell or perhaps its reduced sensitivity to the RNase inhibitor allows it to function as a cytotoxin (Blackburn & Galivanes, 1980). Alternately, the RNase activity of seminal RNase in its dimeric form may lead to its cytotoxicity.

# Non-Cytotoxic Members of the RNase Superfamily

RNase A, its major glycosylated form (RNase B) and their mammalian counterparts are secretory RNases that have been intensively studied using biochemical and biophysical techniques (for reviews see Sierakowska & Shugar, 1977; Blackburn & Moore, 1982; Wlodawer, 1985; Beintema, 1987; Beintema et al., 1988). In these reviews and the referenced literature, pancreatic RNase, RNase C, RNase U1, RNase I, alkaline RNase and neutral RNase are alternate names for the secretory enzymes. The initial translation product of the secretory RNase gene includes a signal peptide and is responsible for the extracellular targeting and the naming of this subfamily of enzymes (Beintema et al., 1988). Secretory RNases of the pancreas are found in considerable quantity in the gut of ruminants and in species with ruminant-like digestion and cecal digestion. In these species the pancreatic RNase content is from 10 and 1000 fold greater than that observed in other herbivores and carnivores. The elevated levels of pancreatic RNase may be the response to the necessity of digesting large amounts of RNA derived from the microflora of the stomach of ruminants (Barnard, 1969). Notably, the extreme differences in the levels of pancreatic RNases among vertebrates are not reflected in the RNase levels of their serum. This implies the existence of an efficient mechanism(s) for regulating serum RNase levels.

Secretory RNases are also expressed at low levels in tissues other than the

pancreas. This would suggest secretory RNases have functions other than digesting RNA. In humans these RNases have a primary structure that is identical to the pancreatic enzyme. In contrast, at least three bovine secretory RNases are known to exist; the pancreatic enzyme, a dimeric form isolated from seminal fluid and the brain RNase that contains 16 additional residues at the C-terminus. These enzymes share 81% and 77% amino acid sequence identity with RNase A (Beintema et al., 1988). At present, only bovine species are known to have these separate secretory RNases.

The non-secretory RNases have been referred to as RNase II, RNase U<sub>s</sub>, acid RNase and EDN in the previously mentioned literature. They are somewhat misnamed as non-secretory RNases are found in the urine, serum, cerebrospinal fluid and semen of mammals. EDN is not unique to the eosinophil as its primary sequence is identical to those of non-secretory RNases isolated from human liver, urine, kidney and placenta. EDN and ECP are glycoproteins containing N-linked carbohydrates. Interestingly, the carbohydrate composition of non-secretory RNases differs depending upon the tissue of origin (Yamashita *et al.*, 1986). While the function of the non-secretory enzymes of the various tissues is unknown, EDN retains its neurotoxicity after the enzymatic removal of its N-linked carbohydrate moieties (Gleich *et al.*, 1986). Recently, EDN has been reported to contain a carbohydrate attached to the C2 atom of the Trp7 indole (Hofsteenge *et al.*, 1995). The significance of this novel carbohydrate attachment is not understood.

Angiogenin is not known to be cytotoxic. Instead, it is as an abundantly secreted protein from the human colon adenocarcinoma cell line HT-29 that stimulates the formation of blood vessels (angiogenesis; Strydom et al., 1985). While tumor growth and metastasis may be facilitated by the overexpression and secretion of angiogenin, this is not true in other carcinomas. The mRNA of angiogenin can be detected in a variety of tissues and is predominantly found in liver (Weiner et al., 1987). In vitro, angiogenin interacts with an angiogenin-binding protein from endothelial cells (Hu et al., 1991) and elicits a second-messenger response (Heath et al., 1989) that results in angiogenesis. The angiogenic

activity of angiogenin requires its RNase activity (Hallahan et al., 1991) just as ONC and EDN require their RNase activity for their cytotoxicity.

Other than the amphibian RNases, the non-mammalian RNases are not known to be cytotoxins. Instead their relationship to RNase A and the molecular evolution of the RNase superfamily have been examined. The amphibian and avian enzymes form distinct groups of related enzymes that do not conform to mammalian classifications. This is not surprising as the gene duplication events that gave rise to the distinct mammalian enzymes occurred long after the divergence of mammals and non-mammals. Interestingly, the amino acid sequence of turtle ribonuclease indicates it is an angiogenin-like enzyme (Beintema et al., 1988). It shares 43% amino acid sequence identity with angiogenin and less than 30% identity with RNase A and EDN (Table 1.3). This suggests an angiogenin-like gene has been conserved since the divergence of mammals and reptiles and suggests it is the common ancestor of all mammalian pyrimidine-specific RNases.

# Ribonucleolytic Mechanism of the Pyrimidine Specific RNases

By definition, these enzymes are those which, at or near neutral pH, cleave RNA endonucleolytically to yield 3'-monoribonucleotides and 3'-oligoribonucleotides ending in Cp or Up via a 2',3'-cyclic phosphate intermediate (Richards & Wyckoff, 1971). The reaction proceeds through a two-step general acid-general base mechanism involving (1) a transesterification in which the 2'-OH nucleophilically attacks the phosphodiester to form the 2',3'-cyclic phosphate intermediate and (2) a hydrolytic step that cleaves the intermediate and yields a product with a terminal 3'-phosphate. The nucleophilic attack by the 2'-OH (step 1) and the solvent molecule (step 2) have both been shown to proceed with an in-line geometry (Usher et al., 1972). Chemical evidence (Crestfield et al., 1963), site-directed mutagenesis (Trautwein et al., 1991), NMR spectroscopy (Rico et al., 1991) and X-ray crystallographic structures of RNase A-inhibitor complexes have agreed that His12, Lys41 and His119 have roles in the catalytic activity of RNase A. Using NMR and X-ray

abstracts the proton of the 2'-oxygen that becomes a nucleophile and attacks the phosphorus, forming a penta-coordinate transition state intermediate. The O(5') leaving group is protonated by His119, the general acid. A salt-bridge between Lys41 and an equatorial oxygen of the phosphate was postulated to stabilize the additional negative charge of the transition state. The refined X-ray crystallographic structures of RNase A in complex with uridine vanadate, a transition-state mimic (Wlodawer et al., 1983) and 3'-cytidine (Zegers et al., 1994) show Lys41 interacts directly with the 2' oxygen as opposed to a phosphate oxygen. This result agrees with charge distribution studies (Davis et al., 1988) and X-ray crystallographic studies on the non-homologous RNase T1 of bacteria that catalyzes an identical reaction. Figure 1.1 is a schematic representation of the reaction catalyzed by members of the pyrimidine specific RNase superfamily of vertebrates.

Recent evidence that imidazole-catalyzed hydrolysis of RNA displays a 'bell-shaped' pH versus rate profile has attracted widespread attention (Anslyn & Breslow, 1989). These results led to a proposed mechanism (triester-like) in which the formation and the breakdown of the pentacovalent phosphorane transition state are catalyzed by different mechanisms. This interpretation differs significantly from the mechanism described above and has been vigorously challenged (Haim, 1992) and defended (Breslow & Xu, 1993) in the literature. Subsequent studies have concluded catalysis by RNase A is not consistent with the triester-like mechanism (Herschlag, 1995) and others have attributed the 'bell-shaped' profile of imidazole-catalyzed hydrolysis of RNA to medium effects (Kirby & Marriott, 1995).

# Enzymatic Activity of the Pyrimidine Specific RNases

The enzymatic activity and specificity of the pyrimidine specific RNases have been characterized using diribonucleotides and natural oligoribonucleotide substrates. Unfortunately, the buffers, salts and ionic strengths used to assay the activity of RNase A

Figure 1.1 The two step ribonucleolytic mechanism shared by members of the pyrimidine (py) superfamily is schematically represented. In the transesterification reaction, the general base abstracts a proton from the O(2') making it nucleophilic and the general acid protonates the leaving group. In the slower hydrolysis step, a solvent molecule is the nucleophile and O(2') is the leaving group. In RNase A, the general acid (+A-H) and general base in these reactions are the invariant histidines, H119 and H12, respectively.

affect the experimental results. Despite these difficulties, RNase A and other secretory enzymes are known to be optimally active at approximately pH 8.0 and to generate products ending in Cp or Up. The base, ribose and phosphate moieties of substrates are referred to using the nomenclature of Richards and Wykoff (1971; Figure 1.2). In a recent study of the transesterification reaction, the measured  $K_m$  (Michaelis constant) of RNase A for UpX and CpX diribonucleotides (where X is a purine nucleotide) was between 0.3 and 2mM and the  $k_{cat}/K_m$  for these substrates was between 0.18 x  $10^5$  and 6.0 x  $10^6 M^{-1} sec^{-1}$ (Bond & Vallee, 1990). In UpX and CpX substrates, the U and C are the B<sub>1</sub> nucleoside and X corresponds to the B2 nucleotide. RNase A shows a slight preference for cytidine in the B<sub>1</sub> position and for adenylate in the B<sub>2</sub> position. Secretory RNases also cleave CpC, UpC, CpU and UpU at much reduced rates and under non-physiological conditions can cleave dsRNA and homopolymers of pA. Table 1.4 list the enzymatic activities of some representative members of the pyrimidine specific RNase superfamily towards various diribonucleotide substrates. The enzymatic activity of EDN is 10 to 100 fold less than of RNase A (Table 1.4) and is optimally active near pH 6.5 (Sierakowska & Shugar, 1977). EDN has a specificity similar to RNase A but with a greater preference for adenylate in the B<sub>2</sub> position. Non-secretory RNases are about 20 times more active against poly(U) substrates than their secretory counterparts, which makes their activity against poly(U) and poly(C) roughly equivalent (Youle et al., 1993). Angiogenin and ONC have pH optima near 6.0. The  $k_{cat}/K_m$  of angiogenin and ONC for these substrates is from  $10^2$  to  $10^5$ times lower than the comparable values for RNase A and EDN. ONC cleaves UpG and CpG with comparatively high specificity and its specificity is the opposite of the specificity of RNase A. It has a 30 fold preference for uridine in the B<sub>1</sub> position and a 500 fold preference for guanylate in B2. Angiogenin has a 10 fold preference for cytidine in the B1 position and has a slight guanylate preference in the B2 position. Recently, angiogenin and ONC have been shown to preferentially cleave tRNA species in reticulocyte lysates (Saxena et al., 1992, Lin et al., 1994).

Figure 1.2 A schematic representation of a short RNA substrate. The base, ribose and phosphate moieties are referred to using the nomenclature of Richards and Wykoff, 1971. The scissile phosphodiester bond is P<sub>1</sub>-O(5'). Note that the P<sub>1</sub>, R<sub>1</sub> and B<sub>1</sub> moieties together are equivalent to a 3'-monophosphate nucleotide. The base, ribose and phosphate numbering increases on the 5' side of the scissile phosphate (P<sub>1</sub>) of the RNA substrate and decreases on the 3' side.

Table 1.4

Enzymatic Activity and Specificity of members of the RNase superfamily.

 $k_{cat}/K_{m} (M^{-1} sec^{-1})$ 

	UpG	UpA	CpG	СрА	C>p <sup>†</sup>	U>p†
RNase Aa	1.8 x 10 <sup>5</sup>	4.0 x 10 <sup>6</sup>	5.1 x 10 <sup>5</sup>	6.0 x 10 <sup>6</sup>	7.7 x 10 <sup>3</sup>	$2.7 \times 10^3$
EDN <sup>b</sup>	1.4 x 10 <sup>3</sup>	1.2 x 10 <sup>5</sup>	1.6 x 10 <sup>3</sup>	2.6 x 10 <sup>5</sup>	1.1 x 10 <sup>2</sup>	ND
Angiogenin <sup>c</sup>	0.4	1.1	4.0	12	0.5	0.02
ONCd	4.7 x 10 <sup>3</sup>	9.4	1.6 x 10 <sup>2</sup>	<2.3	ND	ND

<sup>†</sup> C>p and U>p are cyclic 2',3' cytidine monophosphate and 2',3' uridine monophosphate, respectively.

ND Values have not been determined

<sup>&</sup>lt;sup>a</sup> Values from Harper & Vallee (1990).

b Values from Iwami et al. (1981).

c Values from Bond & Vallee (1990) and Russo et al. (1994).

d Values from W. Ardelt and E. Boix, unpublished results.

Diffraction quality crystals of ONC in complex with a substrate or transition state analogs have not been obtained. To investigate the enzymatic specificity of ONC, a model of ONC has been constructed in complex with UpG, an analog of its preferred substrate. The model allows us to infer the identity of residues that contribute to the B<sub>1</sub> and B<sub>2</sub> specificities of ONC (Chapter 4). In the rEDN structure (Chapter 5), a second sulfate anion binding site has been observed that most likely corresponds to P<sub>-1</sub> phosphate of extended RNA substrates.

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### **CHAPTER 2**

COMPARATIVE MOLECULAR MODELING AND CRYSTALLIZATION OF P-30 PROTEIN: A NOVEL ANTI-TUMOR PROTEIN OF RANA PIPIENS OCCYTES AND EARLY EMBRYOS<sup>†</sup>

#### INTRODUCTION

P-30 Protein (Onconase)‡ is an abundant, basic protein that is isolated from the oocytes or early embryos of *Rana pipiens* (1). It exhibits a specific anti-tumor activity against several human tumor cell lines *in vitro* including notoriously resistant solid tumors (1,2). The P-30 Protein (hereafter referred to as P-30) is also effective against the M109 Madison carcinoma in mice (3). Treatment of susceptible tissue cultures with P-30 results in the accumulation of cells arrested in G<sub>1</sub> phase of the cell cycle and cytotoxicity. Subsequent examination of those cells arrested in G<sub>1</sub> phase reveals heterogeneous total RNA contents, with many cells exhibiting very low levels of RNA (1). This suggests P-30 is able to preferentially enter susceptible cells by an undetermined mechanism. P-30 is currently undergoing phase II human clinical trials in the United States as a prospective solid tumor treatment.

The complete amino acid sequence of P-30 has recently been published (4). The protein consists of 104 amino acids including four disulfide bridges and a blocked N-terminus (pyroglutamyl). The calculated molecular weight is 11,837 Da when basic amino acid side chains are assumed to be fully protonated. More interestingly, it has been shown that P-30 is a member of the pancreatic RNase superfamily of enzymes (4). Currently, P-30's ribonucleolytic activity has been confirmed, however, its catalytic activity is low in

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<sup>‡</sup> Onconase is a trademark of the Alfacell Corporation.

comparison with other members of this family. It is intriguing that P-30's anti-tumor activity is completely inhibited by chemical modification of suspected catalytic residues (4). This suggests that P-30's ribonucleolytic activity is somehow required for its anti-tumor effect.

In light of P-30's exhibited anti-tumor activity, and in the absence of three dimensional structural information, a comparative molecular model (5-7) has been constructed with a view towards understanding the specificity of P-30. The pancreatic RNases have been extensively studied by physiochemical methods (8) and we are fortunate to have several refined, high resolution (≤ 2.0 Å) structures of bovine pancreatic RNase A (9-11). In addition, molecular structures of RNase S (an active form of RNase A resulting from subtilisin cleavage between Ala20 and Ser21) are available (12-13). In this work, RNase A's crystallographic coordinates serve as a template for the comparative molecular modeling of P-30.

### RESULTS

Alignment of P-30 and RNase A sequence — One of the most critical steps in the construction of a comparative molecular model is the alignment of the amino acid sequences, including the placement of both insertions and deletions. The current axiom for modeling homologous structures is attributed to J. Greer; if the alignment is wrong, the structure is guaranteed to be wrong (14,15).

In the present case, P-30 is considerably smaller (104 residues) than bovine pancreatic RNase A (124 residues). The sequence alignment we have deduced is shown in Table 2.1. It results from a two step process. First, the use of automated alignment programs (16-18), and subsequently, a revision of the alignment based on inspection of the P-30 sequence fitted into the RNase A structure. The present alignment makes use of three conserved disulfide bridges, the three catalytic residues and their flanking residues to align the bulk of the sequence. It should be pointed out that one of the disulfides of RNase A

Table 2.1

Complete amino acid sequence alignment of pancreatic RNase A and P-30 Protein

Amino acid homology between RNase A and P-30 Protein are indicated as boxed residues\*. The underscored region represents a revision of the original sequence alignment based on the inspection of the RNase A structure. Open circles above residues indicate constituents of the active site while closed circles above the sequence represent catalytic residues. Open circles below the sequence alignment indicate residues contributing to the hydrophobic core.

\*28% identity and 48% homology.

(Cys87 to Cys72) has no counterpart in P-30, whereas the fourth disulfide bridge of P-30 (Cys87 to Cys104) involves an inserted residue, the C terminal cysteine. An example of sequence alignment revision based on inspection of the RNase A structure is underlined in Table 2.1. In this region P-30 contains a five residue deletion relative to the RNase A sequence Cys65 to Gln74. Various automated alignment programs (16-18) position the deletion at RNase A residues Thr70 to Gln74, as a result of the strong homology at Cys65 to Asn67 (RNase A: Cys Lys Asn, P-30: Ser Lys Asn). However, in light of the structure of RNase A, it is clear that residues Cys65 to Gln69 form part of an exposed loop (bracketed by the Cys65-Cys72 disulfide bridge), whereas residues Thr70 to Gln74 contribute to one of the central strands of a four stranded anti-parallel β-sheet that constitutes part of RNase A's core. Figures 2.1a and 2.1b illustrate the consequences of positioning the deletion at Cys65 to Gln69 and Thr70 to Gln74 respectively. Since deletion of the Thr70 to Gln74 β-strand is likely to change RNase A's overall fold, we have chosen to position this deletion at residues Cys65 to Gln69.

The resulting alignment of P-30 with RNase A has 28% (29/104) sequence identity. When conservative amino acid replacements are included, the sequence alignment has 48% (50/104) similarity (Ala = Val = Leu = Ile = Met = Phe = Tyr, Glu = Gln = Asp = Asn, His = Arg = Lys, Ser = Thr). A complete accounting of the sequences deleted in the alignment of P-30 with RNase A is shown in Figure 2.2, where it can be seen that deleted sequences map mainly to exposed loops on the surface of RNase A. As a result, the alignment is expected to conserve the secondary structural elements of RNase A and many features of its hydrophobic core, in addition to its active site.

Model building and energy minimization — The program MUTATE (Randy Read, unpublished) was used to provide atomic coordinates for the P-30 sequence. In this process the amino-acid side chains of RNase A, were replaced by the side chains of P-30 according to the sequence alignment given in Table 2.1. In cases where small residues in RNase A were replaced by larger side chains in P-30, the extra atoms of the side chain were

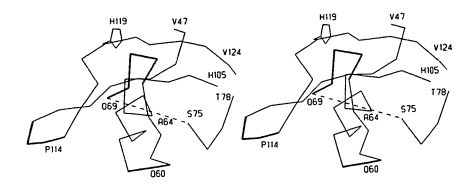


Figure 2.1a An example of sequence alignment revision based upon inspection of RNase A is shown. The dashed lines indicate regions that are deleted and the thick lines represents regions that are retained in two potential sequence alignments. Automated alignment programs (16-18) predict that RNase A residues Thr70-Gln74 are deleted relative to P-30 Protein's sequence.

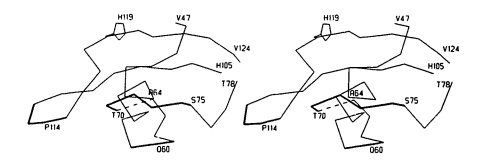


Figure 2.1b: Again, the dashed lines indicate regions that are deleted and the thick lines represents regions that are retained in two potential sequence alignments. In this alignment, Cys65-Gln69 are deleted. This has the advantage of preserving the antiparallel β-sheet (which forms part of RNase A's core) and leaving the ends of the deletion near to one another. The model of P-30 Protein has made use of the alignment depicted in b.

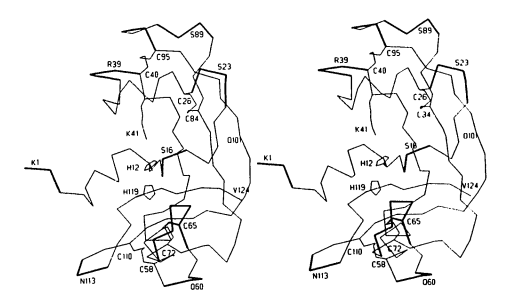


Figure 2.2: A  $C^{\alpha}$  representation of RNase A is presented with deletions (relative to P-30 Protein alignment in Table 2.1) indicated in thick lines and disulfide bridges shown as dashed lines. Note that the deletions occur primarily in exposed loops of the RNase A structure.

positioned according to expected stereochemistry (19,20). The inevitable too close van der Waal's contacts that arise from such amino acid replacements were relieved using the interactive graphics program TOM (21,22). As an example, the too close contacts that result from the replacement of Ala5 (RNase A) by Trp3 (P-30) are shown in Figure 2.3. Figure 2.3 also shows the identical region after graphical intervention. In this instance, rotations about the torsion angles  $\chi_1$  and  $\chi_2$  of Trp3, facilitate the location of a low energy conformation that is free of steric clash. A noteworthy consequence of this reorientation, is the 3.10 Å separation between N<sup>E1</sup> of Trp3 and the carbonyl oxygen of Glu91 suggesting a possible hydrogen bond interaction.

Once all non-hydrogen contacts less than an arbitrarily chosen distance of 2.6 Å were relieved in this manner, the residues at each end of a deletion and the residues adjacent to them in the primary sequence were subjected to 200 cycles of stereochemical regularization using J. Hermann's routine within TOM (21,22). At this point, the additional C terminal Ser103 and Cys104 residues were added to the structure, and the disulfide bridge involving the C-terminal cysteine (Cys87 to Cys104) was constructed, again using J. Hermann's regularization routine within TOM. The P-30 model was then subjected to energy minimization using the CHARMm program within QUANTA (23). The relatively robust steepest descents minimization (25 cycles) procedure was used to tidy up unacceptably close contacts and poor stereochemistry in the model. This was followed by conjugate gradient minimization until convergence, at which point, the N-terminal glutamate was cyclized and manipulation of the model was complete.

As all energy minimization was done in the absence of solvent and ions, some accounting of potential artifacts is required. To this end, the crystallographic coordinates of the RNase A structure were also subjected to energy minimization, under conditions identical to those used on the P-30 model coordinates. In each case, energy minimization was performed with model polar hydrogens, and parameters were invoked which assign electrostatic neutrality to potentially charged residues in an effort to account for the

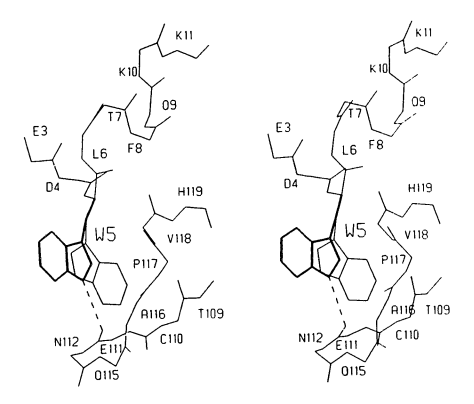


Figure 2.3: An extreme example of unacceptable close contacts introduced by the replacement of Ala5 (RNase A) with Trp (P-30 Protein) and its relief through graphical intervention. The RNase A numbering scheme is used in the figure and the thick lines correspond to the reoriented Trp. In this instance, the rotation of  $\chi_1$  and  $\chi_2$  torsion angles of Trp has relieved close contacts and indicates the potential formation of a H-hydrogen bond (shown as a dashed line) between Trp N<sup>E2</sup> and a mainchain carbonyl oxygen.

lack of solvent. Finally, it should be stated that the model can only be considered tentative.

RNase A and energy minimized RNase A — The root mean square (rms) deviation in atomic coordinates for the minimized structure of RNase A and the model of P-30 from the crystallographic structure of RNase A are listed in Table 2.2. The rms deviation of 0.59 Å between all atoms of the crystallographic RNase A structure and the energy minimized RNase A structure, indicates that gross structural perturbations have not been introduced by the energy minimization procedures adopted. As an indicator, each of the secondary structural elements of RNase A remain intact throughout energy minimization procedure. This is confirmed visually in Figure 2.4, where the RNase A structures ( $C^{\alpha}$ and selected residues), before and after the minimization process have been superposed. It is noteworthy, that in Table 2.2, the all-atom, active site only superposition of RNase A before and after energy minimization actually has an rms deviation greater than that of the remainder of the structure. In fact, the side chains of the catalytic residues His119 and Lys41 (shown in Figure 2.4) undergo some of the largest movements between structures. Under normal circumstances these results would be distressing. However, the crystallographic structure of RNase A contains a bound phosphate anion that interacts noncovalently with each of the three catalytic residues; His12, Lys41 and His119. Thus, it is presumed that the absence of the phosphate anion during energy minimization has led to the significant reorientation of His119 and Lys41 in the minimized RNase A structure. Overall, the series of small differences between the two structures is expected to reflect errors in both the crystallographic coordinates and probably to a greater extent, assumptions inherent in current energy minimization strategies.

When constructing a comparative molecular model, energy minimization procedures serve to relieve too close van der Waal's contacts and to ensure chemically reasonable structures. Given the agreement between the crystallographic RNase A structure and the energy minimized RNase A structure, it is expected that this energy minimization strategy will yield meaningful results when applied to the P-30 model coordinates.

Table 2.2

Rms deviation in atomic coordinates for minimized RNase A and the P-30 Protein model\*

	all atoms (Å)	mainchain (Å)∆	all active site (Å)°
RNase A (crystal) vs. RNase A (minimized)	0.59	0.42	0.63
P-30 Protein model vs. RNase A (minimized)	1.38	1.29	0.52

<sup>\*</sup> Comparisons between atoms in common in accordance with alignment in Table 2.1.

<sup>&</sup>lt;sup> $\Delta$ </sup> Mainchain atoms include N,  $C^{\alpha}$ , C and O.

o The active site includes all atoms of the catalytic triad (His12, Lys41, His119) and 6 other residues that contribute to the substrate binding cleft (Gln11, Asn44, Thr45, Phe46, Phe120 and Asp121).

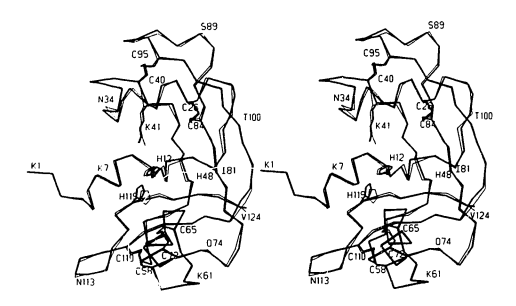


Figure 2.4: Superposition of RNase A crystallographic structure before and after energy minimization. Only  $C^{\alpha}$  atoms, catalytic residues and disulfide bridges are shown. The minimized RNase A structure is shown in heavy lines and disulfide bridges are shown as dashed lines.

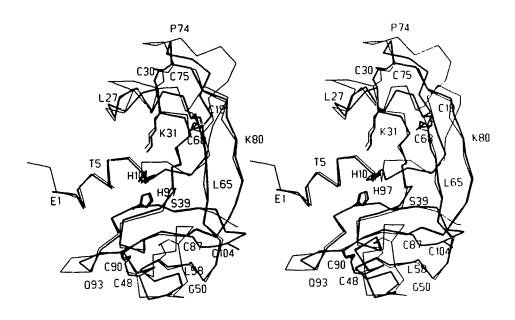


Figure 2.5: Superposition of minimized RNase A and the P-30 Protein model. Again, C<sup>α</sup> atoms, catalytic residues and disulfide bridges are presented. The P-30 Protein model is represented by the heavy lines in this figure and disulfide bridges are shown as dashed lines.

P-30 comparative molecular model — The rms deviations for the several superpositions of P-30 and minimized RNase A are presented in Table 2.2. Figure 2.5 shows a superposition of the catalytic residues, disulfide bridges and  $C^{\alpha}$  atoms of P-30 and the minimized RNase A.

Architecturally, RNase A consists of three regions; an N terminal helix (~ 15 residues) and two larger  $\beta$  sheet containing elements (~ 45 residues) that are separated by a hinge of 3 peptide strands. The two  $\beta$  sheet containing elements are related by approximate 2 fold symmetry and each of these regions contributes a catalytic residue to the active site. In more detail, the N-terminal helix of P-30 (S peptide of RNase S) provides Lys9 and His10 (Gln11 and His12 in RNase A) to the active site and makes contacts with both of the remaining regions. The active site Lys31 and Lys33-Thr35 of P-30 and the corresponding residues of RNase A (Lys41 and Val43-Lys45) are supplied by the large region that is composed of a relatively long three-stranded antiparallel  $\beta$ -sheet and a short helix that packs against the  $\beta$  sheet on the far side of the active site. The remaining element contributes His97-Val99 (His119-Asp121 in RNase A) and is formed by a shorter four-stranded antiparallel  $\beta$ -sheet. This domain also contains a short helix that packs against the  $\beta$  sheet on the side furthest from the active site.

As previously indicated, the sequences in RNase A that are deleted in the P-30 model, primarily involve exposed surface loops. Excluding the two residues deleted from the N terminus of P-30, there are five deletions involving more than one residue (see Table 2.1). In each of these cases, the secondary structural elements that bracket the deleted sequence are retained in the comparative molecular model of P-30. This conservation of secondary structural elements is related to the preservation of both identical and equivalent tertiary interactions that form the hydrophobic core of RNase A. In Table 2.1 it can be seen that the alignment of hydrophobic core residues is 50% (12/24) identical and 79% (19/24) homologous.

Of the five deletions referred to, three occur at a distance from the active site

(Ser22-Tyr25, Gly88-Ser90, Asn113-Pro114) of RNase A. The two remaining deletions (Lys37-Arg39, Cys65-Gln69) map to the periphery of the RNase A active site. Of these the Lys37-Arg39 deletion is only one residue removed from the catalytic Lys41. Despite this, the Lys31 coordinates (P-30) can remain essentially unchanged when compared to Lys41 of the minimized RNase A structures. It is notable that the intervening residue (Cys40 of RNase A; Cys30 in P-30) is involved in a conserved disulfide bridge that presumably contributes to the correct positioning of the catalytic Lys41 in RNase A (Lys31 in P-30). With reference to the deduced mechanism for pancreatic RNases, the catalytic Lys interacts with the pentacovalent transition state. The other deletion, Cys65-Gln69, has been previously discussed and the chosen alignment is somewhat justified by the involvement of Ser54-Leu58 in the four-stranded β-sheet of the P-30 model (Thr70-Gln74 of RNase A).

The sole insertion in the alignment of P-30 and RNase A sequences involves the two C-terminal residues, Ser103 and Cys104. The C-terminal Cys104 participates in a disulfide bridge with Cys87, that is unique to P-30. These additional residues are peripheral to the active site and form a turn on the surface of the P-30 model, immediately following the active site residues His97-Val99.

With reference to the active site residues mentioned previously and defined again in Table 2.2, P-30 only differs from RNase A in three of nine positions. The increased conservation of active site residues is reflected in the all atom, active site rms deviation, which is only 0.52 Å. When comparing the active site differences between P-30 and RNase A, the respective replacements are Lys9, Lys33 and Val99 (P-30) for Gln11, Val43 and Asp121 (RNase A). In an effort to evaluate these amino acid replacements in terms of the specificity of P-30, various crystallographic structures of RNase S- (8,24-26) and RNase A-inhibitor complexes (27,28) have been examined. In general, the results of the inhibitor structures are internally consistent and supportive of earlier chemical studies (29-30). Unfortunately, these studies have been hampered by the lack of available

coordinates for RNase-inhibitor complexes.

The specificity of RNase A is such that the phosphate at the active site has a 3' linkage to the pyrimidine ribonucleotide and a scissile 5' linkage to the remaining ribonucleotide. In native and I hibitor structures of RNase A, N<sup>2</sup> of Gln11 donates a hydrogen bond to the active site phosphate. In the P-30 model, Lys9 has the potential to form a similar hydrogen bond. Despite the increased size (1 C-C bond) of Lys9, there are few steric constraints imposed by neighboring residues, and thus it is possible that this contact is retained in both P-30 and RNase A.

The contribution of Val43 to the RNase A active site is primarily through its mainchain atoms. As such, the non-conservative Val43 to Lys33 replacement present in P-30 probably will not alter substrate binding significantly. Alternatively, the additional positive charge introduced by this substitution could affect the substrate specificity of P-30.

The greatest active site differences between RNase A and P-30 involve the Asp121 to Val99 change in conjunction with the deletion of the Cys65-Gln69 loop. In the several RNase structures, Asp121 makes side chain contacts with the Cys65-Gln69 containing loop, and the active site His97 side chain. These contacts remain unchanged in structures of RNase A-inhibitor complexes. However, one or both of Gln69 and Asn71 contribute side chain hydrogen bonds to the 3' base (purine or pyrimidine) in diribonucleotide inhibitor-RNase A complexes (24-26). In the P-30 model, the Cys65-Gln69 region of RNase has been deleted and thus P-30 most likely will not form the analogous interactions. It is noteworthy that this active site difference involves the 3' ribonucleotide binding site, since RNase A does not show a rigid specificity at this site. To expand on this point, it has been shown biochemically, that RNase A activity towards diribonucleotides, varies over two orders of magnitude in response to various 3' ribonucleotides (31). This tolerance of RNase A for 3' ribonucleotides would suggest that other interactions are of greater importance in substrate binding and catalysis.

In summary the comparative molecular model of P-30 has been constructed with the

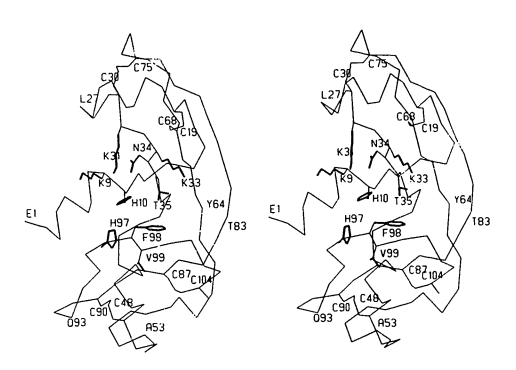


Figure 2.6: The tentative P-30 Protein model. Active site residues are included as well as catalytic residues and disulfide bonds. The active site and catalytic residues are shown in thick lines and the disulfide bridges are indicated by dashed lines.

aim of understanding some of the underlying features of its specificity. In comparison with RNase A, the P-30 model has three significant active site differences which are shown in Figure 2.6. First, Lys9 of P-30 replaces Gln11 of RNase A which donates an important hydrogen bond to the active site phosphate. Second, there are topological changes at the periphery of the active site that result from an insertion and deletions. Since these regions of RNase enzymes may function in the binding of oligoribonucleotides, they may contribute to the specificity of P-30. Finally, the absence of the Cys65-Gln69 (RNase A) loop in P-30, indicates that P-30 probably cannot bind the 3' ribonucleotide in a fashion similar to RNase A. As P-30's specificity with respect to various substrates is currently unknown, it will be interesting to note the similarities and differences between the specificities of P-30 and RNase A as more data become available.

Crystallization — Recently, our lab has grown two separate forms of P-30 crystals, orthorhombic and hexagonal. Of these, the orthorhombic form has been characterized by X-ray diffraction techniques.

Crystals of P-30 were grown by vapor diffusion using a hanging drop. Typically,  $2 \mu L$  of 15 mg/mL P-30 (in H<sub>2</sub>O) are mixed with  $2 \mu L$  of reservoir solution (50% saturated (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>), 10 mM acetate pH 4.5) and allowed to equilibrate with 1 mL of reservoir solution. Crystals grow as rosettes in 3-7 days and achieve maximum size after approximately 1 month. Single crystals can be broken off the rosettes for subsequent diffraction experiments. The crystals have unit cell dimensions of a = 40.76, b = 69.77, and c = 32.54 Å and belong to the space group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>. We are currently optimizing the crystallization conditions in order to obtain large single crystals suitable for high resolution diffraction experiments. It is hoped that this comparative molecular model will contribute to the solution of the three dimensional structure of P-30, when satisfactory crystals are obtained.

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## CHAPTER 3

# THE REFINED 1.7 Å X-RAY CRYSTALLOGRAPHIC STRUCTURE OF P-30 PROTEIN, AN AMPHIBIAN RIBONUCLEASE WITH ANTI-TUMOR ACTIVITY<sup>†</sup>

## INTRODUCTION

P-30 Protein (Onconase)‡ is an abundant amphibian ribonuclease (RNase) that is isolated from the oocytes and early blastomeres of Rana pipiens. It is a member of the pancreatic RNase superfamily (Ardelt et al., 1991) and it is one of several RNases that display a cytotoxic activity in vivo (Mikulski et al., 1990a)‡. P-30 is active against a number of the tumor cell lines tested in vitro. At concentrations of 1.5 µg/mL, P-30 can reduce the number of cells that undergo mitosis by up to 90% (Darzynkiewicz et al., 1988; Mikulski et al., 1990b). It is a distant relative of the extensively characterized RNase A and seminal RNase enzymes (Beintema et al., 1988). The seminal RNase homodimer shows a similar growth inhibition (up to 95%) at 50 µg/mL and RNase A is still weaker as a cytotoxin (Tarnowski et al., 1976). The cytotoxic mechanism of P-30 and related enzymes is under investigation. While it is known that RNase A is a potent cytotoxin when injected (Saxena et al., 1991) or when targeted for entry (Ryback et al., 1991) into Xenopus oocytes; it is not known if intact, active RNase A enters susceptible cells in vivo. The seminal RNase homodimer is internalized by both susceptible and resistant cell lines (Vescia et al., 1980). P-30 binds to 9L glioma cells with two separate affinities  $(K_d=6.2\times10^{-8}M)$  and  $K_d=2.5\times10^{-7}M$ ) and its IC50 is similar to the low affinity  $K_d$  (Wu et al., 1993). Wu et al. (1993) have also shown that P-30's RNase activity and cytotoxicity are inhibited by the alkylation of histidine residues. These results suggest P-30 has an

<sup>†</sup> A version of this chapter has been published. Mosimann SC, Ardelt W. & James MNG. 1994. The refined 1.7 Å x-ray crystallographic structure of p-30 protein, an amphibian ribonuclease with anti-tumor activity. J. Mol. Biol. 236, 1141-1153.

<sup>‡</sup>P-30 is currently undergoing phase II human clinical trials (U.S.A.).

intracellular site of action that depends upon its ribonucleolytic activity. If this is true, it may be significant that amphibian RNases are not inhibited by mammalian placental ribonuclease inhibitor (PRI) and that the mammalian RNases are not inhibited by the amphibian counterpart to PRI (Beintema et al., 1988). This may explain P-30's enhanced cytotoxicity when compared to the mammalian enzymes. Alternatively, it may be P-30's distinctive RNase activity or its specific binding to the surface of susceptible cells that contribute to its increased cytotoxicity.

In this work, the X-ray crystallographic structure solution and refinement of P-30 is reported at 1.7 Å resolution. The molecular conformation is described with an emphasis on the stereochemistry of the active site. The discussion assesses the accuracy of the previously published comparative molecular model (Mosimann et al., 1992).

## **METHODS**

P-30 was isolated and purified from Rana Pipiens oocytes and early embryos using previously described protocols (Ardelt et al., 1991). Orthorhombic crystals (space group, P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>) of P-30 were grown according to Mosimann et al. (1992). The unit cell parameters for the high-resolution native data set used in the refinement are  $a = 40.53 \pm 0.08$  Å,  $b = 69.64 \pm 0.09$  Å,  $c = 32.52 \pm 0.08$  Å and  $\alpha = \beta = \gamma = 90^{\circ}$  The calculated solvent content for this crystal is 36.3% and the specific volume,  $V_{\rm m}$ , is 1.94 Å<sup>3</sup>/Da (Matthews, 1968).

## (a) Structure determination

The structure of P-30 was determined using the multiple isomorphous replacement (MIR) method. The X-ray intensity data were collected locally on San Diego Multiwire Systems (SDMS) multiwire proportional counters. Graphite monochromated  $CuK_{\bar{\alpha}}$  X-rays were generated with a Rigaku Rotoflex RU-200BH rotating anode source operated at 40 kV and 150 mA. The data were processed to 1.7 Å for the native protein and to

maximum resolutions of 2.4 Å, 2.7 Å and 3.0 Å for the three heavy atom derivatives using the SDMS software (Hamlin, 1985; Howard et al., 1985). Statistics for the processing of native and derivative data sets are presented in Table 3.1. The heavy atom binding sites were located from difference Patterson maps and subjected to both positional and occupancy parameter refinement with the program MLPHARE (Otwinowski, 1991). The MIR phasing statistics for each of the heavy atom derivatives are presented in Table 3.2. The absolute hand was determined using the anomalous signal of the trimethyl lead acetate derivative and confirmed by the presence of right-handed helical density in the resulting electron density map. The 2.7 Å MIR electron density map was calculated using figure of merit (FOM) weighted structure factor amplitudes and phases having a mean FOM of 0.68. This map was of sufficient quality to enable the tracing of the complete polypeptide chain. The comparative molecular model of P-30 (Mosimann et al., 1992) was fitted to the electron density on an SGI 4D-25 workstation using a locally modified interactive graphics program TOM Alberta/CalTech 2.8.0 (based on FRODO: Jones, 1985). The three refined heavy atom positions are located on the surface of the polypeptide chain tracing in the MIR map and each is within 5 Å of P-30 and a symmetry related molecule.

During the MIR structure determination of P-30, a number of attempts were made to solve the structure by molecular replacement. Both the comparative molecular model of P-30 (Mosimann et al., 1992) and the crystallographic structure of the phosphate containing bovine RNase A (Wlodawer et al., 1983) were used as search models for rotation function calculations with the following programs: YORROT (Dodson, E.J., 1985), MOLECULAR REPLACEMENT PROGRAM PACKAGE (Alzari & Navaza, 1990) and X-PLOR (Brünger et al., 1987). Additional search models representing truncated versions of the original molecular models were also tested with YORROT. For each search model a series of rotation function maps were calculated as several parameters were varied: (1) the resolution range (the maximum resolution was varied between 5.0Å and 2.0Å; the minimum resolution was varied between 15.0Å and 5.0Å) and

Table 3.1 Summary of data collection statistics for P-30 native and derivative data sets+

	Res.	No.	Unique	%			unit cell lengths (Å)		
	(Å)	Ref.	_	Complete	I/σIII	R <sub>m</sub> ‡	a	b	С
Native	1.7	43858	9495	89.4	11.0	0.050	40.53	69.64	32.52
Native	2.1	22458	5104	88.7	20.0	0.029	40.51	69.63	32.53
Pt Deriv.	2.4	13032	3833	98.3	20.7	0.051	40.59	69.61	32.51
Pb Deriv.	2.7	12549	2670	96.9	10.4	0.065	40.48	69.65	32.51
V Deriv.	3.0	8248	1833	89.4	22.9	0.036	40.65	69.73	32.48

<sup>+</sup> Space Group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>; wavelength (Cu $K_{\bar{\alpha}}$ ) = 1.5418Å

<sup>‡</sup>  $R_m = \sum_{l:kl} \sum_{l:kl} |I_{ave} - I_{obs}|/\sum_{l:kl} |I_{ave}|$  Pt = platinum ethylenediamine dichloride; Pb = trimethyl lead acetate; V = oxo-vanadium diflouride.

Table 3.2

Multiple Isomorphous Replacement phasing statistics for P-30 at 2.7Å resolution

	sites	Rdiff <sup>†</sup>	fractional coordinates			В	Occ.	Phasing
			x	у	z	(Å <sup>2</sup> )	(e-)	Power <sup>‡</sup>
Pt	1	0.049	0.0480	0.2975	0.4189	44.16	64.87	1.21
Pb	1	0.158	0.2235	0.2141	0.0431	19 <b>77</b>	33.66	1.77
v	1	0.091	0.1208	0.3555	0.21	< 	28.50	1.61

Mean Figure of Merit = 0.678

Pt = platinum diethylamine dichloride; Pb = trimethyl lead acetate; V = oxo-vanadium diflouride.

 $<sup>\</sup>tau \; R_{diff} = \Sigma ||F_{PH}| \text{-} |F_{nat}||/\Sigma |F_{nat}|$ 

<sup>‡</sup> Phasing Power =  $\sqrt{\Sigma F_H^2} / \sqrt{\Sigma (|F_{PH obs}| - |F_{PH calc}|)^2}$ 

(2) the threshold intensity of the measured data and (3) the integration radii of the rotation function map. A consistent solution to the rotation function was not obtained using any of these search models. In general, there was a lack of discrimination between the highest peak in the rotation function map and other peaks within the same map.

## (b) Structure refinement

The initial model of P-30 was subjected to molecular dynamics refinement using the program X-PLOR (Brünger, et al., 1987). Data between 20.0-2.0 Å resolution with intensities greater than 3oIII were selected for the refinement using the simulated annealing protocol described in the X-PLOR manual. The protocol includes: (1) energy minimization using restraints to the X-ray data, (2) a molecular dynamics simulation at 2000 K, (3) a slow cooling to 300 K, (4) positional parameter refinement and (5) an individual B-factor  $(B = 8\pi \overline{u^2})$ , where  $\overline{u^2}$  is the mean-square vibrational amplitude) refinement. During this procedure the crystallographic R-factor  $(R = \Sigma ||F_0| - |F_c|| / \Sigma ||F_0||)$ , where  $|F_0|$  and  $|F_c|$  are the observed and calculated structure factor amplitudes) was reduced from 0.543 to 0.275. The resulting model coordinates were then subjected to five cycles of restrained leastsquares refinement using PROLSQ (Hendrickson & Konnert, 1981) prior to manual refitting. For this and each subsequent refitting session the model was displayed using TOM Alberta/Caltech 2.8.0 on the interactive graphics system with two types of electron density maps: (1) a map calculated with FOM weighted coefficients (2|F<sub>0</sub>|-|F<sub>c</sub>|) and calculated phases and (2) a map computed with FOM weighted difference coefficients (IF<sub>0</sub>I-IF<sub>c</sub>I). In all cases the FOM weighted coefficients were generated with the program SIGMAA (Read, 1986). At several points in the refinement standard omit maps were calculated. In these maps selected atoms were omitted from a short (5-9 cycle) refinement and the subsequent map calculation. The resulting electron density in the region of the omitted atoms represents a relatively unbiased view of the true electron density at these positions. During the initial manual refitting session the amino acids Pyr1 (pyroglutamyl), Asp2, Thr25, Asn26 and Leu27 occupied regions of ambiguous electron density and were removed from the model. These residues were then rebuilt into the relatively unbiased omit density during the following refitting sessions. After a total of 23 cycles of PROLSQ refinement and three manual refits the R = 0.233 for data between 10.0-2.0 Å having  $I \ge 3$ olll. During subsequent refitting sessions, a sulfate anion was added to the active site and 21 ordered solvent molecules were included. The active site anion was modeled as a sulfate due to its presence in the crystallization medium (and the absence of phosphate). The sulfate oxygens were fit to the electron density and refined throughout (final B-factors range from  $24\text{\AA}^2$  to  $26\text{\AA}^2$ ). This reduced the R-factor to 0.209 after a total of 47 cycles of restrained least-squares refinement using 10.0-2.0 Å data with  $I \ge 3$ olll.

The model was further refined against 10.0-1.7 Å data having  $I \ge 1.5\sigma III$  and additional ordered water molecules were included in the model. Throughout the refinement no effort was made to refine water molecules with alternate positions and/or variable occupancies. The final model contains all 104 amino acids, a sulfate anion and 96 ordered water molecules and has an R-factor of 0.178. The stereochemical parameters associated with the final cycle of least-square refinement are given in Table 3.3. The P-30 coordinates have been deposited in the Protein Data Bank (pdb1onc.ent) at Brookhaven National Laboratory.

#### RESULTS

## (a) Quality of the refined structure

In the final 2|F<sub>0</sub>|-|F<sub>c</sub>| map (1s contour level) there is continuous electron density for all main-chain atoms. Two or more side-chain atoms of the residues Lys8, Arg40, Glu42, Lys44, and Lys78 do not have associated electron density. Lys8 and Lys44 have no density beyond C<sup>7</sup> and Lys78 has none beyond C<sup>8</sup> at this contour level. Arg40 and Glu42

Table 3.3

Stereochemical statistics for the final cycle of restrained least-square refinement

R-factor	0.178 for 8653 reflections > 1.5olII between 10.0 - 1.7 Å resoluti				
	0.231 for all 9495 unique reflections between ∞ - 1.7 Å resolution				
Model	B-factor (mean)				
	826 protein atoms	13.4 Å <sup>2</sup>			
	96 waters	$30.2  \mathrm{\AA}^2$			
	1 sulfate anion	23.9 $^2$			
Restraints		rmsd(Å)	Target(Å)		
	1 - 2 bond distance	0.016	(0.016)		
	1 - 3 bond distance	0.030	(0.022)		
	1 - 4 planar distance	0.034	(0.030)		
	planar groups	0.009	(0.010)		
	chiral volumes	0.216	(0.200)		
	non-bonded contacts				
	single torsion contacts	0.218	(0.200)		
	multiple torsion contacts	0.167	(0.200)		

rmsd =  $\sqrt{\sum (X_x - \bar{x})^2 / m}$ ; Target is the stereochemical weight assigned during the last cycle of least squares refinement.

have electron density have electron density beyond  $C^{\beta}$  but it is weak and discontinuous. The Arg40-Glu42 residues form intermolecular contacts with Thr25'-Leu27' and the electron density associated with these residues is shown in Figure 3.1.

A Ramachandran plot of the P-30 dihedral angles is presented in Fig. 3.2. P-30 has three glycine residues and 101 non-glycine residues. In the case of the glycine residues, two (Gly50 and Gly102) participate in reverse turns and adopt conformations that are most commonly associated with glycine residues while the third (Gly106) is part of the C-terminal strand  $\beta$ 5. Gly50 and Gly102 each occupy the i+2 position of their respective reverse turns (Table 3.4). Of the remaining 101 residues five (His29, Cys30, Cys75, Asn92, Gln93) have left-handed helical conformations. All of these residues are in reverse turns occupying the second or third positions (Table 3.4). Residues 91 to 94 form a type I' (Sibanda & Thornton, 1985) reverse turn (Fig. 3.3.). The short contact (3.01 Å) between the C $\beta$  of Gln93 (i+2) and the carbonyl oxygen of Asn92 (i+1) that is common to this conformation is stabilized by several hydrogen bonding interactions: Glu91 O to Ala94 N (3.21 Å), Gln93 N to Ile51 O (2.69 Å) and Gln93 O<sup>E1</sup> and Wat119 (2.76 Å). Thr71 has a conformation ( $\phi = -95^{\circ}$ ,  $\psi = -162^{\circ}$ ) that lies slightly outside the allowed  $\beta$ -conformation. This residue participates in an Asx type turn in which the side-chain Oel forms a hydrogen bond to the main-chain nitrogen (3.11 Å) of the residue i+2 in the polypeptide (Arg73 N). The Thr71 N also donates a hydrogen bond to the Lys76 O (2.97 Å).

The B-factor versus sequence number profile of P-30 is shown in Fig. 3.4. In general, the lowest thermal parameters in the structure are associated with the β-strands while the residues with the highest thermal parameters comprise loop regions. Eleven residues of P-30 have average B's greater than 15 Å<sup>2</sup>. Of these, the only ones that belong to a helix or a β-strand are Arg40 and Glu42 (helix α3). Six of the eleven residues occur in two loops (Ser24-Leu27, Ser72-Arg73) that are covalently linked by the nearby Cys30 to Cys75 disulfide bridge. This results in a cluster of relatively mobile residues found at

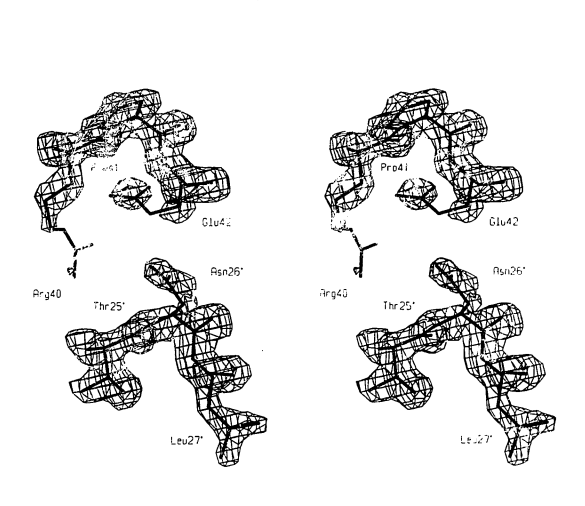


Figure 3.1 The 2|Fo|-|Fc| electron density (1 $\sigma$  contour) associated with Arg40-Glu42 and Thr25'-Leu27' of a symmetry related molecule. This represents one of the least well-defined regions of electron density in the P-30 structure.

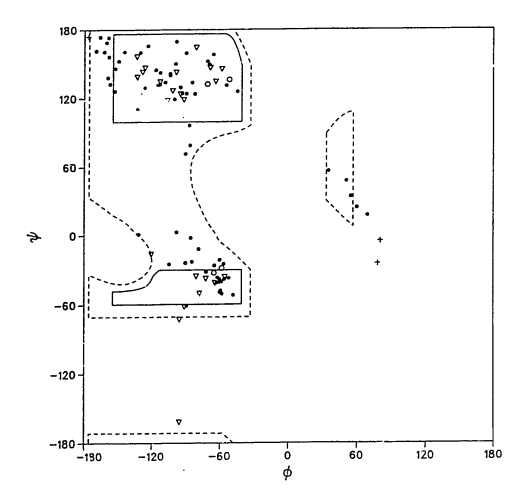


Figure 3.2 The Ramachandran plot of the completed P-30 model is presented with the various allowed regions of conformational space indicated in boxes. The 71 ( $\phi = -95^{\circ}, \psi = -162^{\circ}$ ) and Gln93 ( $\phi = 69^{\circ}, \psi = 17^{\circ}$ ) are the only non-glycine residues that occurs in mildly disallowed regions of the plot. The crosses are glycine residues, the inverted triangles represent amino acids with branched side chains, the open circles correspond to proline residues and closed circles are used for the remaining residues.

Table 3.4

Reverse turns in P-30

Residues	Туре	φ2 ψ2 φ3 ψ3	H bond distance (Å) O <sub>i</sub> ····N <sub>i+3</sub>
Phe28-Lys31	I'	35, 56, 55, 30	2.97
Cys48-Ile51	II	-54, 131, 81, -5	3.01
Arg73-Lys76	П	-52, 136, 60, 24	3.07
Glu91-Ala94	I'	51, 48, 69, 17	3.21

The reverse turns have been classified according to Sibanda & Thornton, 1985.

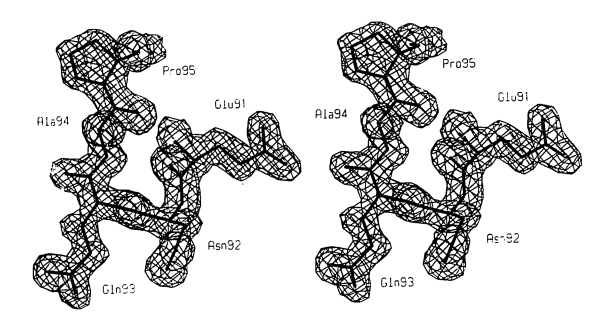


Figure 3.3 The  $2|F_0|-|F_c|$  electron density (1 $\sigma$  contour) belonging to Gln93 ( $\phi = 69^\circ$ ,  $\psi = 17^\circ$ ). Gln93 occupies the i+2 position of this type I' turn, a position typically occupied by glycine. As a result, Asn92 O forms a short intramolecular contact (3.01 Å) with the Gln93 C $\beta$ .

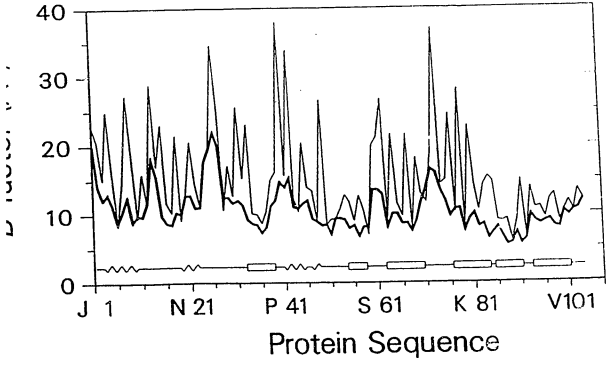


Figure 3.4 A plot of the average thermal parameter (B-factor) against the amino acid sequence number of P-30. The thick line represents the average main-chain B-factor while the thin line represents the average side-chain B-factor: P-30's secondary structures are indicated above the sequence numbers; rectangles represent β-strands, waves represent helices and straight lines indicate turns and loops. The highest average B-factors are associated with several loops (Thr25-Leu27, Arg40-Glu42 and Ser72-Arg73) that are involved in intermolecular contacts.

one end of the molecule (Fig. 3.1). Several of the other residues that have relatively high average main-chain B-factors make intermolecular van der Waal's contacts with these mobile loops. Asn13' contacts Thr71 (3.92 Å) and Glu42' contacts Asn26 (3.76Å). The final residue with an average main-chain B-factor greater than 15 Å<sup>2</sup> is the N-terminal residue, Pyr1. The largest B-factor for a protein atom is 44.7 Å<sup>2</sup> (Arg40 N<sup>η1</sup>) and the highest average side-chain B-factors are associated with Asn26, Arg40, Glu42 and Arg73. Other side chains with high a erage B-factors are Lys8, Lys49, Glu62 and Lys78. In each of these cases, the side chains are relatively long and extend into a disordered solvent channel.

There are 96 full occupancy solvent molecules in the P-30 model. They have B-factors ranging from 12 to 51 Å<sup>2</sup> (the highest in the structure). The water molecules are approximately equally distributed within this range. There are 16 waters with B's less than 20 Å<sup>2</sup>, 28 waters with B's between 20-30 Å<sup>2</sup>, 35 waters with B's between 30-40 Å<sup>2</sup> and 17 waters with B's greater than 40 Å<sup>2</sup>. The solvent molecules were refined with full occupancy and variable B-factors in all cases.

### (b) Molecular conformation of P-30 Protein

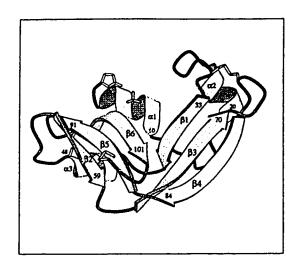
P-30 is a single domain protein that is presently the smallest known member of the pancreatic RNase superfamily. It contains 104 amino acids and has a calculated  $M_T$  of 11,834 Da (pH 7.0, His protonated). The fold of its polypeptide chain is shown in Fig. 3.5a and as stick model in Fig. 3.5b. P-30's fold is closely related to that of RNase A. It is a predominantly antiparallel  $\beta$ -structure with a prominent N-terminal  $\alpha$ -helix that makes contacts to each of its two lobes.

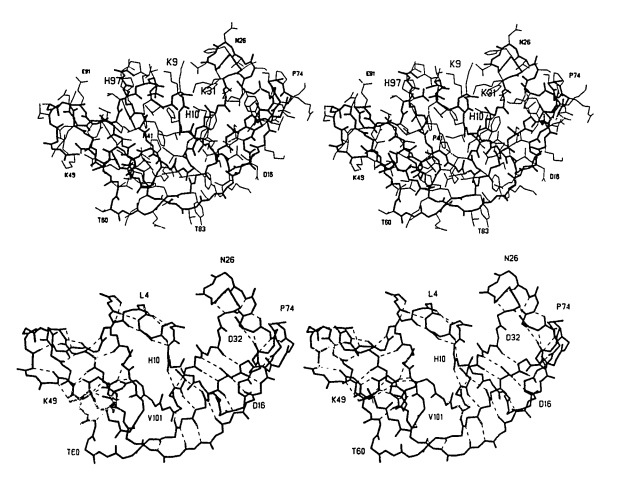
The secondary structure of P-30 has been analyzed with the DSSP program (Kabsch & Saunders, 1983), giving the following proportions: 40.4% antiparallel β-strand, 20.2% helix, 39.4% turns and aperiodic segments (loops). The intramolecular

Figure 3.5a A Molscript (Kraulis, 1991) ribbons diagram of the molecular conformation of the P-30 protein. The helices and strands are labeled sequentially as they occur in the primary sequence. Three of the four disulfide bonds are visible while the fourth (Cys19-Cys68) is hidden behind strand β3. The N-terminal pyroglutamyl residue and the active site sulfate anion are shown with all their bonds.

Figure 3.5b An all atom representation of the P-30 molecule. The main-chain atoms are connected by thick lines while the thin lines represent the bonds between side-chain atoms. The sulfate anion at the active site is also shown in thin lines.

Figure 3.5c The main-chain tracing of the P-30 molecule. The solid thick lines connect all main-chain atoms and the thin dashed lines represent hydrogen bonds between pairs of main-chain atoms (  $2.50 \text{ Å} \le d \le 3.30 \text{ Å}$  ).





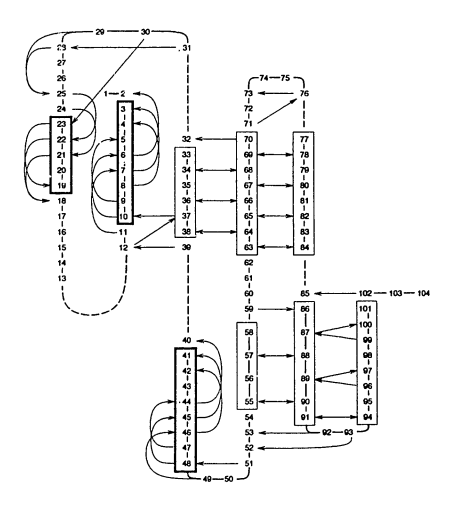


Figure 3.6 The intramolecular main-chain hydrogen bonding network for P-30. The  $\beta$ -strands are indicated by thin boxes and the helices by thick boxes. The arrows represent hydrogen bonds ( 2.50 Å  $\leq$  d  $\leq$  3.30 Å ) and the arrowheads are directed at the hydrogen bond acceptors. Residues Cys87 and Thr89 of strand  $\beta$ 5 each form main-chain hydrogen bonds with two residues from strand  $\beta$ 6.

Those residues that do not adopt helical or  $\beta$ -strand conformations account for the remainder of the P-30 structure. These segments include four reverse turns that have  $i \rightarrow i+3$  hydrogen bonds and several aperiodic segments. The reverse turns are listed in Table 3.4. Of the segments that do not fit the  $\alpha$ -helix,  $\beta$ -strand or reverse turn classification only Ile11-Asp18 is longer than four residues. These residues form an irregular loop connecting helices  $\alpha 1$  and  $\alpha 2$ . The first three residues (Ile11-Asp13) of this loop have an extended  $\beta$ -strand like conformation but only form a single hydrogen bond with the neighboring strand  $\beta 1$  (Thr12N $\rightarrow$ Ile37 O, 3.10 Å). The other five residues of this loop are involved in reversing the direction of the polypeptide chain. The equivalent loop

in RNase A contains a subtilisin cleavage site that is not present in P-30.

There are three short connections between the lobes of P-30 that may act as a hinge between the two lobes. The residues at these connections include Ser39-Arg40, Thr59-Glu62 and Lys85. The final segments of the P-30 structure with non-helix and non-strand  $\phi$ ,  $\psi$  torsion angles are the two N-terminal and the three C-terminal residues. The N-terminal Pyr1 and Asp2 are folded back against the helix  $\alpha$ 1 and Pyr1 is directed into the active site of the enzyme. The C-terminal residues Gly100-Ser103 form a structure closely related to a reverse turn but lacking the  $i\rightarrow i+3$  hydrogen bond. The Gly100 C=O bond vector lies in the plane of the  $\beta$ -sheet ( $\beta$ 2,  $\beta$ 5,  $\beta$ 6) whereas the Ser103 N-H bond vector is perpendicular to the plane of the  $\beta$ -sheet. As a result, this turn-like structure does not have an  $i\rightarrow i+3$  hydrogen bond. The C-terminal amino acid, Cys104, forms a disulfide bridge with Cys87.

P-30 is a disulfide-rich protein with eight cysteine residues forming four disulfide bonds (Fig. 3.5a,b). Three of the four disulfide bonds have counterparts in RNase A and these involve the following residues: (1) Cys19 and Cys68, (2) Cys30 and Cys75 and (3) Cys48 and Cys90. The fourth disulfide connects Cys87 and Cys104 and is unique to the amphibian members of the RNase superfamily. The respective -S-S- torsion angles for these disulfide bonds are -79°, -77°, -72° and +99°. The Cys19 to Cys68 bond connects the N-terminus of helix  $\alpha$ 2 to the central strand ( $\beta$ 3) of the antiparallel  $\beta$ -sheet that it packs against. Similarly, the Cys48 to Cys90 disulfide tethers the  $\alpha$ 3 helix to the central strand ( $\beta$ 5) of its associated  $\beta$ -sheet. Whereas these two disulfides fulfill structurally similar roles, the other two disulfides are structurally unrelated as indicated by their  $\phi$ ,  $\psi$  angles and especially their  $\alpha$ 3 torsion angles. In the Cys30 to Cys75 bridge, the cysteine residues are both involved in reverse turns. Cys30 and Cys75 each occupy the  $\alpha$ 4 position of their respective reverse turns and each adopts a left-handed helical conformation as a consequence. The Cys87 to Cys104 disulfide links the C terminus to the central strand ( $\beta$ 5) of the  $\beta$ 2,  $\beta$ 5,  $\beta$ 6 antiparallel sheet.

### (c) Intermolecular contacts

Table 3.5 contains a compilation of the direct intermolecular contacts within the P-30 crystal and Fig. 3.7 is an intermolecular packing diagram. The contacts listed in Table 3.5 have been separated into nine van der Waal's contacts (3.20 < d < 3.90 Å) and 14 hydrogen-bonding interactions. All together, these intermolecular contacts involve 29 of the 104 residues in P-30. All of the nine van der Waal's contacts involve one or more residues that are on loops. The contacts involve or are adjacent to residues in previously discussed sequences possessing relatively high main-chain B-factors. The Pro74 side-chain is involved in four separate close contacts. The shortest intermolecular contact occurs between Tyr64 OH and Arg73  $C^{\beta}$  (3.22 Å). When examining the intermolecular hydrogen bonds in Table 3.5, two interactions are particularly noteworthy. The first involves the Lys55 N and the sulfate oxygen, O4. In Fig. 3.8, the long Lys55 side-chain adopts an almost fully extended conformation ( $\chi_1 = -175^{\circ}$ ,  $\chi_2 = -171^{\circ}$ ,  $\chi_3 = -154^{\circ}$ ,  $\chi_4 = 168^{\circ}$ ) that allows it to bridge the solvent channel. The side-chain atoms have welldefined electron density and individual B-factors that are below 15 Å2. Lys55 is the only residue that makes an intermolecular contact with the active site (Fig. 3.8). Its  $N^{\zeta}$  atom is 2.79 Å from the sulfate O4 and 2.73 Å from the Phe98 O. As a result of this position, Lys55 and the sulfate anion sterically block His97 from adopting X1 torsion values near 180°. The second intermolecular interaction of note involves the Glu91 side chain. Glu91 OE1 makes two separate hydrogen bonds to the side chains of Asp16 and Lys81. In turn, the Asp16 side chain accepts an intramolecular hydrogen bond from Lys81 N\u00e4. Given the pH of the crystallization buffer (pH 4.5) and the pKa's of the carboxyl side-chains (-4.5); the most likely effect is that the Asp16 and Glu91 carboxyls ingle proton in a short hydrogen bond (2.49 Å) and have a single net negative charge. Similar carboxyl-carboxylate interactions have been reported for a variety of other systems

Table 3.5

Summary of the intermolecular close contacts and hydrogen bonds for the refined P-30 model

Symmetry operation	Van der Waal's contact d < 3.90 Å	s 	Hyd: d	rogen bonds < 3.30 Å	
x, y, z+1		3.22 3.36 3.77 3.56 3.84 3.20	Ser24 O Ser61 OY	- Arg40 Nm1 - Asp32 Oδ1	2.71 2.52
x+1/2, -y+1/2, -z	Asp $20  C^{\beta}$ - Val $101  O$ Asp $20  O^{\delta 1}$ - Ser $103  C^{\beta}$ Asn $21  C^{\gamma}$ - Val $101  C^{\gamma 1}$ Thr $25  C^{\gamma 2}$ - Lys $80  C^{\epsilon}$	3.38 3.30 3.87 3.39	Asn21 Oδ1 Asn21 Nδ2 Arg73 N <sup>n1</sup> Arg73 N <sup>n2</sup> Lys76 N <sup>ζ</sup>	<ul> <li>Lys80 N<sup>ζ</sup></li> <li>Asp67 O<sup>δ1</sup></li> <li>Ser103 O<sup>γ</sup></li> <li>Ser103 O<sup>γ</sup></li> <li>Gly102 O</li> </ul>	2.58 2.88 3.05 3.05 2.58
x+1/2, -y+1/2, -z+1			Тут38 ОН	- Lys85 N <sup>ζ</sup>	2.97
-x+1/2, -y, z+1/2			Lys55 N <sup>Ç</sup> Lys55 N <sup>Ç</sup>	<ul><li>Phe98 O</li><li>Sulfate O4</li></ul>	2.79 2.75
-x+1, y+1/2, -z+1/2			Asp16 O <sup>δ1</sup> Leu79 O Lys81 N <sup>ζ</sup>	<ul> <li>Glu91 O<sup>ε1</sup></li> <li>Asn92 N<sup>δ2</sup></li> <li>Glu91 O<sup>ε1</sup></li> </ul>	2.49 2.94 3.29

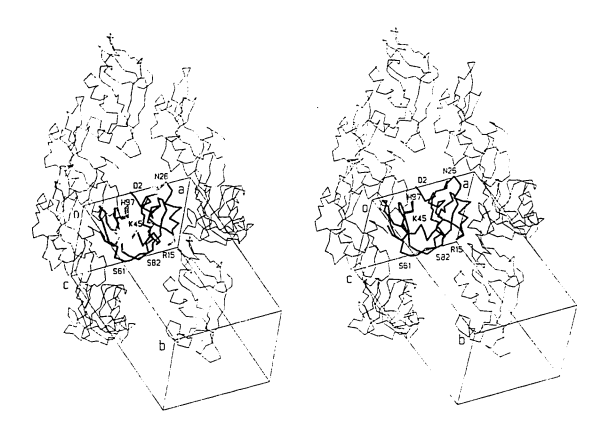


Figure 3.7 The intermolecular packing diagram of orthorhombic P-30. The unit cell is shown as a rectangular box and the thick C<sup>α</sup> tracing represents a single P-30 molecule. Seven other neighboring P-30 molecules are included in thin lines.

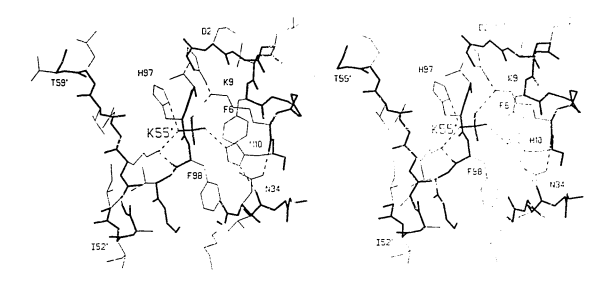


Figure 3.8 The intermolecular contact at the active site is shown as a stick model. The symmetry related Lys55' residue is observed to extend across a solvent channel and into the active site where it makes contacts with the sulfate anion and Phe98. This intermolecular contact prevents His97 from adopting  $\chi_1$  torsion angle near 180°. It should be noted that residues adjacent to Lys55' are not involved in intermolecular contacts.

(Sawyer & James, 1982) and very short hydrogen bonds are also found in protein structures (Fraser et al., 1992).

## (d) Active site of F-30 Protein

The active site is a long cleft that runs approximately parallel to strand β6. It is formed at the junction of helix α1, strand β1 and strand β6. The active site residues are Pyr1, Lys9, His10, Lys31, Thr35, His97 and Phe98. The residues and their associated electron density are shown in Fig. 3.9. The aromatic Phe98, the catalytic His10 and the polar Thr35 side chains line the interior of the cleft. His10 packs against Phe6 (not shown) and the Cβ of Phe98 and Thr35 packs against the relatively exposed aromatic ring of Phe98. The hydrogen bonding requirements of the His10 and Thr35 side chains are satisfied by a well-ordered sulfate anion and several ordered water molecules and for non-covalently bound to the active site. The sulfate anion is presumed to be a mimic of the phosphodiester moiety of RNA bestrates

Lys9, His10, His97 and the 8 form salt bridges and hydrogen bonds with the oxygen atoms of the sulfate anion. Lys9 N\z is 3.15 \hat{A} from the sulfate oxygen, O2 and forms a 3.00 \hat{A} hydrogen bond with Pyr1 O\xi1. His10 N\xi2 is directed between the O1 (2.64 \hat{A}) and O3 (3.32 \hat{A}) atoms of the anion. His97 N\xi01 (2.95 \hat{A}) forms a salt bridge and Phe98 N (3.20 \hat{A}) form hydrogen-bonded contacts with the sulfate oxygen O\xi4. This network of salt bridges and hydrogen bonds at the P-30 active site is shown in Fig. 3.10. Active site residues that do not contact the the anion directly include Pyr1, Lys31 and Thr35. The pyroglutamyl residue is folded back against the N-terminal helix and its maintain nitrogen atom forms a hydrogen bond with Val96 O (2.69 \hat{A}). This directs the O\xi1 atom of Pyr1 into the active site where it accepts hydrogen bonds from Lys9 N\z and Wat306 (2.64 \hat{A}). In a model of an enzyme-substrate complex, the Pyr1 side chain is expected to pack against the ribose moiety 5' to the scissile phosphodiester bond. The inclusion of Lys31 as an active site residue is based upon its conserved character within the

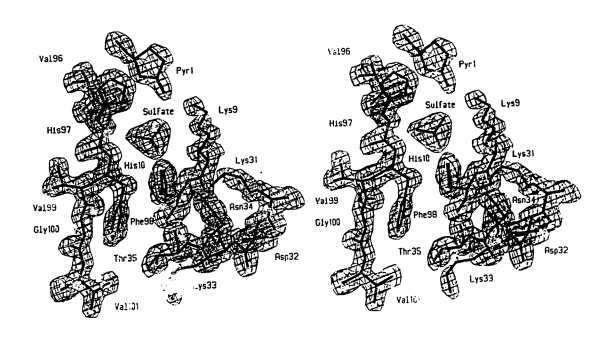


Figure 3.9 The electron density associated with the active site is shown. The contour level is 1σ. A sulfate anion occupies the active site in this orthorhombic crystal form. The refined model is represented by the thick times. Residues Pyr1, Lys9-His10, Lys31-Thr35 and Val96-Val101 are shown

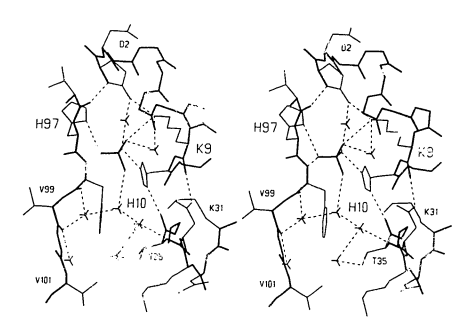


Figure 3.10 An all atom representation of the active site of P-30 is shown with the detailed hydrogen bonding network indicated as dashed lines. The thin lines correspond to side chains while the thick lines represent the bonds between the main-chain atoms of the molecule. The sulfate anion and seven ordered solvent molecules are also included.

pancreatic RNase superfamily and upon experiments in the related RNase A system. Chemical modification studies (Hirs et al., 1965; Raetz & Auld, 1972) and X-ray crystallographic structures of RNase A and RNase S-inhibitor complexes (Richards & Wyckoff, 1973; Wodak et al., 1977; Borkakoti, 1983; Wlodawer et al., 1983) demonstrate that the equivalent residue in RNase A (Lys 41) has a role in catalysis or in substrate binding. In our structure of P-30, Lys31 N\(^{\zeta}\) is 5.49 \(^{\zeta}\) from the nearest sulfate oxygen (O3). If a similar role is expected for this residue, a conformational change is necessary. The Thr35 N and CY1 atoms form hydrogen bonds with the ordered water molecules Wail17 (3.07 Å) and Wat217 (2.81 Å) respectively within the active site. In the previously mentioned RNase A-inhibitor complexes, the equivalent Thr atoms give rise to the pyrimidine specificity of RNase A by forming hydrogen bonds to O2 and N3 of the base, respectively. P-30 has a similar pyrimidine specificity and it is active against vidine-3',5'-guanylate (UpG) and weakly active against cytidine-3',5'-guanylate (CpG) (Ardelt, unpublished). In contrast, RNase A cleaves all diribonucleotide combinations that begin with a pyrimidine (UpX, CpX where X=A,C,G,U). Three water molecules are involved in the remaining hydrogen bonds to the sulfate anion. Wat306 is 2. / Å from the sulfate oxygen O1, Wat405 is 3.02 Å from the sulfate oxygen O3 and Wat426 is 2.77 Å from sulfate oxygen O2.

P-30 and KNase A share a two-step, pyrimidine specific endoribonucleolytic activity that is catalyzed by two invariant histidines and an invariant tysine (P-30: His10, Lys31, His97). In RNase A, the first step of the reaction is an intramolecular transesterification that produces a cyclic-2',3'-phosphate intermediate and releases the 3' end of the RNA substrate. The second step is the hydrolytic cleavage of the intermediate to a polyribonucleotide that ends in a 3'-phosphoester of cytidine (Cp) or uridine (Up) (for review see Richards and Wyckoff, 1973). Each of the reaction steps proceeds with an in line geometry and involves a pentacovalent phosphorus transition state. In P-30, His10 and His97 are the equivalent general acid/general base catalysts while Lys9 and Phe98

stabilize the active site anion. In the RNase A-transition state analog X-ray structure (Wlodawer et al., 1983) the invariant lysine interacts with the O2' of the phosphate mimic. The equivalent lysine in P-30 is Lys31 and it is 5.49 Å from the nearest sulfate oxygen. As mentioned above, in order for Lys31 to interact with the P-30 transition state during catalysis, it will have to move.

### DISCUSSION

P-30 Protein is a distant relative of bovine RNase A. It is twenty residues smaller and one can align the sequences and achieve 28% identity. Prior to the X-ray structure solution, a comparative molecular model of P-30 was constructed using bovine RNase A coordinates and the aligned sequences (Mosimann et al., 1992). Since comparative molecular model building is dependent upon accurate sequence alignments (Greer, 1985), regions with high dentity are expected to show the closest agreement in a comparison of the X-ray structure (Chothia & Lesk, 1986). Fig. 3.11 shows a superposition of the Ca atoms of the P-30 model as derived above with the X-ray structure of P-30. The root mean square (RMSD) deviation for the 81 Ca atoms that can be superposed within 3 Å is 1.26 Å.

The comparative model of P-30 accurately predicts  $\alpha$ -helical and  $\beta$ -strand conformations with two exceptions. The exceptions involve Pyr1 and helix  $\alpha$ 2. In the model, Pyr1 is part of the N-terminal  $\alpha$ -helix. In the X-ray structure Pyr1 folds back against the N-terminal helix becoming part of the active site. This is a significant difference in the modeling of P-30's active site based on RNase A. Helix  $\alpha$ 2 is located far from the active site. In the X-ray structure, helix  $\alpha$ 2 is four residues shorter than predicted in the model and it is found systematically shifted by more than 2.0 Å from its predicted position. This difference is related to the conformational differences in the adjacent loop structures and possibly to P-30's intermolecular packing.

The P-30 model does not accurately predict the conformation of loops that possess

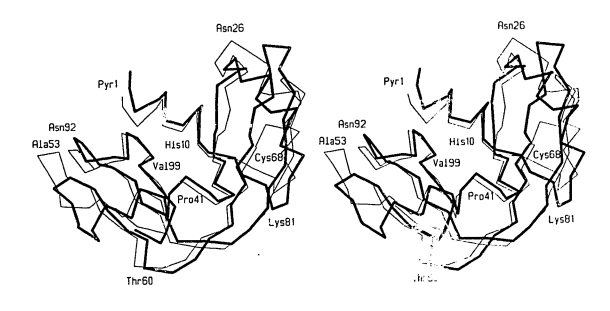


Figure 3.11 The C<sup>α</sup> superposition of the comparative molecular model (this, ames) and P-30 X-ray crystallographic structure (thick lines). The RMS deviation is 1.26 Å for the 81 C<sup>α</sup> atoms pairs that superpose within 3 Å.

low sequence identity and are the sites of deletions relative to RNase A. Not surprisingly, these represent the areas of greatest divergence in the X-ray structures of P-30 and RNase A (Fig. 3.11). The loops following helix  $\alpha$ 3 (P-30: Lys49-Ser54; RNase A: Ser59-Thr70) and strand  $\beta$ 3 (P-30: Thr71-Cys75; RNase A: Thr87-Cys95) deserve comment.

In the case of the loop following helix  $\alpha 3$ , several sequence alignment protocols (Dayhoff, 1979; McLachlan, 1972) position a five residue deletion in P-30 at RNase A residues Thr70-Gln74 (Mosimann et al., 1992). These residues form part of strand  $\beta 2$  in the structure of RNase A. Since the deletion of strand  $\beta 2$  seemed unreasonable, the automated alignment was revised in the comparative molecular model and the deletion was positioned in a loop region of RNase A (Cys65-Asn69). The loop's predicted conformation in P-30 is similar to the RNase A residues Ser59-Ala64. While the alignment revision is justified by the presence of strand  $\beta 2$  in the X-ray structure of P-30, the loop's conformation is radically different in the predicted and real structures. The pairwise positional differences reach 11 Å when comparing the predicted and real structures. Fig. 3.12 is a superposition of the loop region between helix  $\alpha 3$  and strand  $\beta 2$  in the real and predicted structures. Such large differences in predicted and real loop conformations have been observed for eucaryotic and procaryotic serine proteinase (Delbaere et al., 1979; Read et al., 1984).

The loop following strand β3 is the site of a three residue deletion in P-30. In the comparative molecular model this deletion was placed between Thr71-Ser72 (Fig. 3.11). As a result, Ser72, Arg73 and Pro74 were aligned with the RNase A residues Lys91, Tyr92 and Pro93. In light of the P-30 X-ray structure, this deletion is now placed between Pro74-Cys75. In the modified alignment Ser72, Arg73 and Pro74 are equivalent to Gly88, Ser89 and Ser90 of RNase A. In this case, the original alignment was chosen over the alternative after examining several related sequences. In the closely related amphibian lectin (53% aminc acid identity with P-30), there is no deletion in this loop. The additional residues occur between the P-30 residues Thr71-Ser72; the site of the original deletion. In

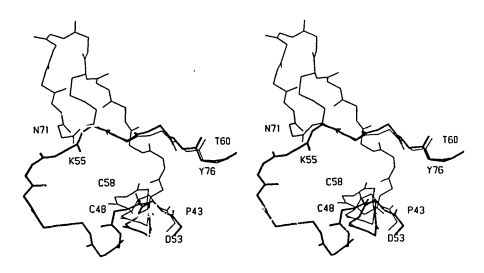


Figure 3.12 The predicted and real conformations of the P-30 loop containing residues Lys49-Asn54. The predicted conformation of the loop is shown in thin lines and the conformation observed in the X-ray structure is shown in thick lines. The region between Cys48 and Lys55 exhibits differences up to 11 Å between equivalent C<sup>α</sup> atoms.

bovine angiogenin, the sequence equivalent to Lys91, Tyr92 and Pro93 (RNase A) is Ser, Arg, Pro; the same sequence found in 2-35. In spite of this

evidence for the correct alignment of the deletion, the X-ray structure clearly positions the deletion between residues Pro74-Cys75. The differences in the predicted and real structures of these two loops point to two 'facts of life' in comparative molecular modeling. First, the correct sequence alignment does not guarantee an accurate structure prediction and second, the correct structural alignment is not necessarily the one with the greatest sequence identity.

Finally, it should be pointed out that the Ser72-Pro74 sequence of ONC is involved in several close Van der Waal's contacts with a symmetry related molecule (Table 3.5). Interestingly, these contacts would be relieved by the alignment used in the comparative molecular modeling.

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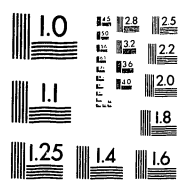
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### **CHAPTER 4**

# REFINED 2.2 Å X-RAY CRYSTALLOGRAPHIC STRUCTURE OF RECOMBINANT ONCONASE AND COMPARISON WITH ONCONASE AND RNASE A<sup>†</sup>

### INTRODUCTION

Onconase<sup>TM</sup>(ONC) is a small (104 amino acids), basic ribonuclease (RNase) isolated from the eggs of rana pipiens (Ardelt et al., 1991). Its amino-acid sequence is distantly related to members of the pyrimidine specific RNase superfamily of vertebrates (Ardelt et al., 1991). A comparative molecular model constructed on the basis of this sequence similarity confirmed that the sequence of ONC makes it a member of the RNase superfamily (Mosimann et al., 1992). ONC displays a specific cytotoxic activity against several tumor cell lines in vitro (Darzynkiewicz, Z. et al., 1988) and in vivo (Mikulski et al., 1990; Mikulski et al., 1993). Its RNase activity is required for its cytotoxic action (Wu et al., 1993). When compared to RNase A, ONC has a greatly reduced RNase activity (0.001-1.9%) towards various diribonucleotide substrates and exhibits a different substrate specificity (W. Ardelt, unpublished results).

The X-ray crystallographic structure of ONC has been determined at 1.7Å resolution (Mosimann et al., 1994). Its overall fold is similar to RNase A (Zegers et al., 1994), RNase S (Richards & Wykoff, 1971) and angiogenin (Acharya et al., 1994). One novel feature of the ONC structure is the hydrogen bond between the N-terminal pyroglutamyl 1 (Pyr1; 5-oxo-2-pyrollidine carboxylate)  $O^{\epsilon 1}$  atom and Lys9N $^{\zeta}$  at the enzyme's active site.

Recently, recombinant Onconase (rONC) having the N-terminal sequence of <sup>f</sup>Met. 1(formyl methionine)Glu<sub>1</sub>Asp<sub>2</sub>... has been produced in a bacterial expression system and the expressed enzyme was shown to have no significant ribonucleolytic activity or cytotoxic activity (less than 2% in high-salt conditions, Y. Wu & R.J. Youle, unpublished

results). The rONC X-ray crystallographic structure has been determined and refined at 2.2Å resolution to further investigate the structural basis for its lack of activity. The resulting structure has been compared to that of wild-type ONC. To further our understanding of the ONC system's ribonucleolytic activity and specificity, ONC-deoxyuridine-3',5'-deoxyguanylate (d(UpG)) and rONC-d(UpG) models have been constructed based upon the RNase A-deoxycytidine-3',5'-deoxyadenylate (d(CpA)) crystallographic structure (Zegers et al., 1994).

### RESULTS

### Quality of the refined structure

The refined rONC structure includes 828 non-hydrogen protein atoms, a single sulfate anion bound in the active site and 72 ordered solvent molecules. The N-terminal fMet<sub>1</sub> residue is not ordered in the electron density maps and its position not included in the refined model. Table 4.1 presents the conventional R-factor, the root mean-squared deviation of various stereochemical parameters and the average B-factors for several classes of atoms for the rONC structure. The rONC coordinates will be deposited in the Protein Data Bank at the Brookhaven National Laboratory (Bernstein *et al.*, 1977).

In the final 3|F<sub>0</sub>|-2|F<sub>c</sub>| (|F<sub>0</sub>| and |F<sub>c</sub>| are the measured and calculated structure factors respectively) electron density map contoured at 1 $\sigma$ , the electron density is continuous and well defined for all main chain atoms excepting Glu1N. At the same contour level, there is no electron density for the Glu1 side-chain and several other solvent exposed side-chain groups have discontinuous electron density. At the 0.8 $\sigma$  contour level the Glu1 side-chain electron density is weak and discontinuous. The electron density associated with three of the twelve lysine side chains (33, 49 and 80) is absent beyond CY. These residues represent the most-poorly defined regions of electron density in the structure. Figure 4.1 shows a stereo diagram of the electron density associated with residues Glu1 to His10

Table 4.1

Stereochemical statistics for the final cycle of restrained parameter least-square refinement

R-factor	$0.165$ for 4968 reflections $ F_0  > 1.0\sigma  F_0 $ between 20.0 - 2.2 Å resolution			
Model			B-factor (mean)	
	417 mainchain atoms <sup>1</sup>		$15.0~\mathrm{\AA}^2$	
	410 sidechain atoms		19.0 Å <sup>2</sup>	
	72 waters	32.7 Å <sup>2</sup>		
	1 sulfate anion		26.4 Å <sup>2</sup>	
Stereochemistry		rmsd <sup>2</sup>	Restraint <sup>2</sup>	
	Bond length	0.013Å	0.700	
	Bond Angles	1.62°	0.700	
	Pseudorotation Angles	13.95 <sup>6</sup>	1.00	
	Trigonal planar groups	0.009Å	1.50	
	General planar groups	0.008Å	3.00	
	non-bonded contacts	0.035Å	5.00	
	Temperature factor	$2.17$ Å $^2$	0.200	

<sup>&</sup>lt;sup>1</sup>The Met -1 residue was not observed in this work and has not been included in the model.

 $<sup>2</sup>_{rmsd} = ((\Sigma_i(X_i-X_j)^2/N)^{0.5})$ , where  $X_i$  and  $X_j$  are the observed and ideal values and N is the number of values being considered.

<sup>&</sup>lt;sup>3</sup>Restraint is the relative weight assigned to the stereochemical parameter during the final cycle of refinement.

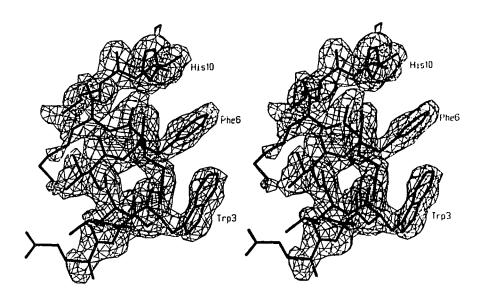


Figure 4.1 The 3lF<sub>0</sub>l-2lF<sub>c</sub>l α<sub>calc</sub> electron density distribution associated with the N-terminal α-helix of rONC is shown at the 1σ contour level. Residues 1-10 are drawn with thick lines and the electron density in represented by thin lines. The Glu1 and Asp2 residues adopt an α-helical conformation. Electron density corresponding to the fMet<sub>-1</sub> residue is not observed; this residue has not been included in the rONC structure. The density associated with the Glu1 sidechain is the weakest in the structure. Lys9 and His10 make hydrogen bonded contacts to the active site sulfate anion through their respective side-chain atoms.

for the rONC structure. The bound sulfate anion does not have distinct lobes of electron density corresponding to the four oxygen atoms in maps calculated with either  $3|F_0|-2|F_c|$  or  $2|F_0|-|F_c|$  coefficients at this resolution. In omit maps (Bhat, 1988) calculated after refinement without the anion, the sulfate anion oxygen positions are observed in the electron density and correspond to the refined sulfate oxygen coordinates. The  $|F_0|-|F_c|$   $\alpha_{calc}$  omit map electron density contoured at  $3\sigma$  is shown in Figure 4.2.

The Ramachandran plots and average B-factor (B =  $8\pi u^2$ , where  $u^2$  is the mean-square vibrational amplitude) versus sequence number plots for rONC and ONC (Mosimann et al., 1994) are virtually identical and have not been provided. Only Asp2 has appreciably different  $\phi$ ,  $\psi$  dihedral angles in the Ramachandran plots of the two structures. Asp2 has  $\beta$ -strand like  $\phi$  (- $160^\circ$ ) and  $\psi$  ( $169^\circ$ ) angles in ONC but adopts an  $\alpha$ -helical conformation ( $\phi$ =- $77^\circ$ , y=  $44^\circ$ ) in rONC. While the average main chain and side chain B-factor versus sequence number plots are similar for the two structures, the rONC average B-factors are roughly 25% larger than the equivalent average B-factor in the ONC structure. The one large divergence in the average B-factor ( $\langle B \rangle$ ) plots occurs at the N-terminus: ONC:Pyr1 main-chain  $\langle B \rangle$ =19.9Ų, side-chain  $\langle B \rangle$ =22.7Ų, rONC:Glu1 main-chain  $\langle B \rangle$ =49.7Ų, side-chain  $\langle B \rangle$ =64.8Ų. The absence of electron density corresponding to  $\langle Met_{-1} \rangle$  and the high B-factors associated with Glu1 indicate the N-terminus of rONC: more mobile than its counterpart in ONC.

### Molecular conformation

The rONC structure can be superimposed on the ONC structure with an rmsd of 0.69Å for all 104  $C^{\alpha}$  atoms and 0.18Å for the 103  $C^{\alpha}$  atoms of residues Asp2-Cys104 (Figure 4.3). The secondary structural elements (helix  $\alpha$ 1 to  $\alpha$ 3, strand  $\beta$ 1 to  $\beta$ 6) described in the ONC structure are present here with one significant difference. In rONC, helix  $\alpha$ 1 is two residues longer and extends from Glu1 to His10. The conformational difference in Asp2 adds this residue and Glu1 to helix  $\alpha$ 1 in rONC. As a result,

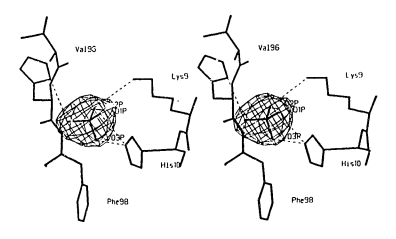


Figure 4.2 The  $|F_0|$ - $|F_c|$   $\alpha_{calc}$  electron density associated with the sulfate anion site after refinement leaving out the atoms of the anion. The electron density is contoured at the  $3\sigma$  level. The density is roughly spherical and does clearly indicate four distinct lobes of electron density corresponding to the sulfate oxygens.

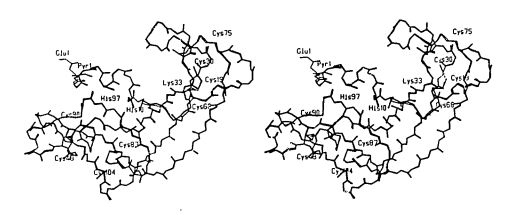


Figure 4.3 A representation of the main chain atoms of the least-squares superposed structures of ONC (thin lines) and rONC (thick lines). The rmsd for  $103~\text{C}^{\alpha}$  atoms is 0.18Å. Significant conformational differences in the two structures are confined to the N-terminus. In the ONC structure the Pyr1 and Asp2 residues fold back against the N-terminal  $\alpha$ -helix. In rONC the Glu1 and Asp2 residues continue in an  $\alpha$ -helical conformation. The  $\text{C}^{\alpha}$  atoms of Pyr1 and Glu1 are separated by 7Å.

the  $C^{\alpha}$  and  $C^{\delta}$  atoms of Pyr1 (ONC) and Glu1 (rONC) are separated by 7.0 and 8.7 Å, respectively. This leads to the large rmsd for the 104  $C^{\alpha}$  atoms of these two structures.

Figure 4.4 is a stereo diagram of the superposed active sites of rONC and ONC and Table 4.2 lists the hydrogen bonds involving the sulfate anion oxygens. Other than the conformational differences of the two N-terminal residues, the active sites of the two structures are virtually identical. Lys9N<sup>\zeta</sup>, His10N<sup>\zeta</sup>2, His97N<sup>\delta1</sup> and Phe98N occupy equivalent positions in the least-squares superposition of these structures and the sulfate anions are bound in nearly identical orientations. The sulfur atom positions of the superposed anions are only slightly different (0.29Å). In both structures, His10N<sup>£2</sup> forms a bifurcated hydrogen bond with the sulfate oxygens O1 (SulO1) and O3 (SulO3) whereas His 97N $^{\delta 1}$  and Phe 98N donate hydrogen bonds to SulO4. In ONC, Lys 9N $^{\zeta}$  donates hydrogen bonds to SulO2 (3.09Å) and Pyr1O<sup>E1</sup> (2.91Å). In rONC there is no structural equivalent to Pyr1 and the Lys9Nζ to SulO2 hydrogen bond distance is increased by 0.4Å. Wat254 and Wat 258 of rONC hydrogen bond to SulO2 and SulO1, respectively and have counterparts in ONC. There is an additional intermolecular hydrogen bond from K55'N\$\xi\$ of a symmetry related molecule at -x+1/2, -y, z-1/2 to SulO4 in both structures. The K55' side chain is not expected to perturb the conformation of the active site very much (Mosimann et al., 1994).

There are 72 and 96 ordered solvent molecules in the rONC and ONC structures, respectively. A total of 54 of the 72 ordered solvent molecules are common (within 1.0Å) in the two structures (rmsd=0.48Å). The difference in the number of ordered solvent molecules likely results from two sources, differences in the resolution (2.2Å versus 1.7Å for ONC) and in the refinement technique. The rONC structure has been refined with TNT release 5D (Tronrud, 1992) using a solvent continuum whereas ONC was refined using PROLSQ (Hendrickson & Konnert, 1981) and no solvent continuum.

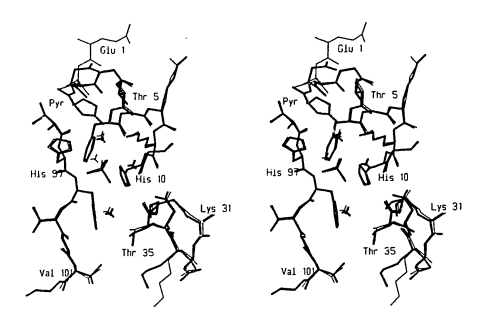


Figure 4.4 An all atom representation of the active sites of the least-squares superposed structures of ONC (thick lines) and rONC (thin lines). Ordered solvent molecules (small tetrahedra) and the bound sulfate anion observed in each structure are also included. The active site residues Lys9, His10, His97 and Phe98 have nearly identical conformations in the two structures. The sulfate anion is bound in the same orientation with a slightly shifted position. The major conformational differences in the two structures are confined to the N-terminus (Glu1 and Asp2).

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Table 4.2

Conserved hydrogen bonds involving the sulfate anion in the superposed ONC and rONC structures

Protein Atom name+	Sulfate Atom name	distance(Å) ONC	distance(Å) rONC	
Lys9N <sup>ζ</sup>	O2	3.09	3.51	
His10NE2	<b>O</b> 1	2.69	2.88	
His10N <sup>€2</sup>	O3	3.32	3.05	
His97Nδ1	04	2.93	3.08	
Phe98N	<b>O</b> 4	3.20	3.03	
Wat254	O2	3.02	2.99	
Wat258	<b>O</b> 1	2.51	2.50	
Lys55'N <sup>ζ‡</sup>	04	2.79	2.82	

<sup>+</sup> Wat numbers are taken from rONC. In ONC, Wat405 and Wat306 are equivalent to Wat 254 and Wat258.

<sup>‡</sup> Lys55' is from a symmetry related molecule and forms an intermolecular contact.

# Superpositions of rONC and ONC with RNase A

The rONC and ONC structures have been superposed by least-squares methods with RNase A coordinates (1RPF, 1RPG, 1RPH; Zegers *et al.*, 1994) obtained from the Protein Data Bank (Bernstein *et al.*, 1977). The rmsds for the overlaps of rONC and ONC with RNase A (1RPG) are 3.16Å and 3.22Å, respectively for the 102 equivalent  $C^{\alpha}$  atoms. Alternatively, the rmsds for the  $C^{\alpha}$  atoms that can be superposed within 3.8Å are 1.37Å for rONC (89  $C^{\alpha}$ ) and 1.31Å for ONC (87  $C^{\alpha}$ ). The N-terminal residue and 14 residues from loop structures in ONC (15-18, 24-27, 49-54) account for the 15 residues whose  $C^{\alpha}$  atoms do not superpose within 3.8Å of equivalent RNase A atoms.

The ONC and rONC residues that comprise the anion binding site come from helixα1 (rONC:Glu1-His10, ONC:Trp3-His10) and strand-β6 (Ala94-Val101). In the ONC superposition with RNase A based upon the 87  $C^{\alpha}$  atoms that superpose within 3.8Å, the secondary structures contributing residues to the active site are shifted relative to their counterparts in RNase A. In this superposition the N,  $C^{\alpha}$ , C, O and  $C^{\beta}$  atoms of the ONC active site residues (Lys9, His10, His97 and Phe98) and their counterparts in RNase A superpose with an rmsd of 0.62Å. The difference in position of these residues is a result of a shift observed in the position of the two lobes of ONC (lobe 1: helices- $\alpha$ 1,  $\alpha$ 3, strands-β2, β5, β6; lobe 2: helix-α2, strands-β1, β3, β4) when compared to RNase A. The 52  $C^{\alpha}$  atoms of lobe 1 of ONC (residues 1-12, 41-62, 85-102) have been leastsquares superposed with the equivalent  $C^{\alpha}$  atoms of RNase A to improve the overlap of the active site residues. The rmsd is 0.80Å for the 45 out of 52  $C^{\alpha}$  atoms of lobe 1 that can be superposed within 3.8Å and the main chain and  $C^{\beta}$  atoms the ONC and RNase A active site residues have an rmsd of 0.32Å in this improved superposition. Figure 4.5 shows the least-squares superposition of ONC and RNase A using the 45 of 52  $C^{\alpha}$  atoms of lobe 1 that can be superposed within 3.8Å. In Figure 4.5, the ONC and RNase A residues of lobe 2 are rotated slightly relative to one another as a result of the improved

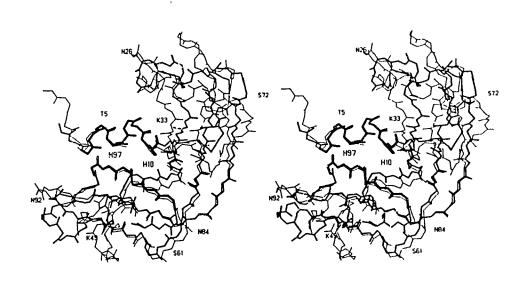


Figure 4.5 The least-squares superposition using 52 C<sup>α</sup> atoms of lobe 1 of rONC(1-12, 41-62 and 85-102; thick lines) and the equivalent atoms in RNase A (1RPG; thin lines). The rmsd is 0.80Å for 45 out of 52 C<sup>α</sup> coordinates that can be superposed within 3.8Å. The helix-α1 and strand-β6 of lobe 1 contain the active site residues in these structures and are shown in heavy lines. The rmsd between the superposed main-chain and C<sup>β</sup> atoms of the active site residues of ONC (Lys9, His10, His97 and Phe98) and RNase A is 0.32Å in the superposition based on lobe 1. The rmsd is 0.62Å for same active site atoms when the superposition based on the entire molecule.

agreement in the overlap of the active site residues of lobe 1. The angles between strands- $\beta$ 1,  $\beta$ 3,  $\beta$ 4 of lobe 2 of ONC and their counterparts in RNase A are 7.9°, 7.4° and 7.5° respectively.

The position and orientation of the sulfate anion bound to ONC has been compared to the sulfate anion bound to RNase A. The superposed ONC and sulfate containing RNase A (1RPH; Zegers et al., 1994) active sites are shown in Figure 4.6 using the improved superposition. The sulfate anions bind in closely similar positions and orientations in the two structures and suggest that ONC and RNase A will bind substrate in an equivalent manner. The conserved His10 and Phe98 residues contact equivalent anion oxygens in the two structures as does Lys9N $^{\zeta}$  (ONC) and its counterpart Gln11 (RNase A). His97 has a side chain conformation ( $\chi_1=-73^\circ$ ,  $\chi_2=-53^\circ$ ) that is different from that of the equivalent RNase A residue ( $\chi_1=153^{\circ}$ ,  $\chi_2=80^{\circ}$ ) in the complex. The ONC His97 side chain conformation is similar to one of two His119 conformations observed in several uncomplexed RNase A structures (Borkakoti et al., 1982; Martin et al., 1987; deMel et al., 1992). In both the ONC and the rONC structures, the Lys55N $^{\zeta}$  intermolecular contact with the sulfate anion sterically prevents His97 from adopting the side chain conformation observed in the RNase A structure. In solution, His97 will be able to adopt a conformation equivalent to that seen in the RNase A. The invariant Lys31 (ONC) and Lys41 (RNase A)  $C^{\alpha}$  atoms are 3.04Å apart in the superposition.

In order to deduce a possible substrate binding mode, ONC-d(UpG) and rONC-d(UpG) model structures have been constructed based upon the RNase A-d(CpA) complex structure. To generate the models, the individual lobes of ONC were superposed with the RNase A-d(CpA) complex and the d(CpA) inhibitor was replaced by d(UpG), an analog of the preferred substrate for ONC. In the RNase A reaction mechanism, the ribose 0(2') is the nucleophile in the initial transesterification that produces a cyclic-2',3'-monophosphate intermediate and 0(2') is the leaving group in the subsequent hydrolysis to

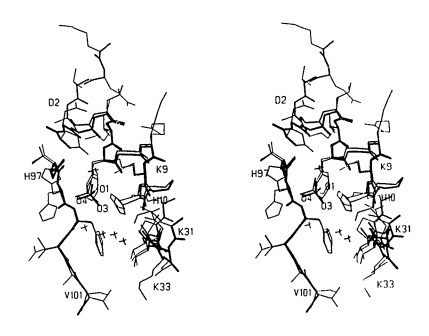


Figure 4.6 The superposed ONC and sulfate containing RNase A (1RPH) active sites using the superposition from Figure 4.5. The ONC structure is shown as bold lines and RNase A as light lines. Main chain atoms are represented by thicker lines than side chain atoms. The position and orientation of the bound sulfate anion of ONC and RNase A are similar. The active site histidines residues contact equivalent anion oxygens in the two structures as do Lys9 (ONC) and its counterpart Gln11 (RNase A). The His97 conformation in ONC corresponds to an alternate conformation observed in other RNase A structures.

a 3'-monophosphate product. d(CpA) is a non-hydrolysable deoxy form of a diribonucleotide substrate that lacks the ribose 0(2'). The ONC-d(UpG) and rONC-d(UpG) model structures are seen in Figures 4.7a-b as well as suggested hydrogen bonds to the inhibitor. The sulfur atoms of the ONC and rONC anions are 0.57Å and 0.62Å from the superposed position of the phosphate atom of the d(CpA), respectively. An equivalent superposition of sulfate containing RNase A and the RNase A-d(CpA) complex, the comparable distance is 0.31Å.

The proposed hydrogen bonds in the ONC-d(UpG) modeled structure have been compared to the hydrogen bonds in the RNase A-d(CpA) structure. The base, ribose and phosphate subsites are referred to as  $B_1$ ,  $R_1$ ,  $P_1$ ,  $B_2$  and  $R_2$ , respectively (Richards & Wykoff, 1971). In the ONC-inhibitor model, the hypothetical hydrogen bonds from Lys9N $^{\zeta}$  and His10N $^{\varepsilon}$ 2 to the phosphate moiety of the inhibitor ( $P_1$ ) are of reasonable length (2.81Å and 3.20Å). The His10N $^{\varepsilon}$ 2 hydrogen bond is similar to that seen in the RNase A complex structure and the Lys9N $^{\zeta}$  hydrogen bond is analogous to RNase A's Gln11N $^{\varepsilon}$ 2 hydrogen bond. Phe98N makes a 2.80Å hydrogen bond with the phosphate oxygen equivalent to SulO1. By adopting the conformation of the equivalent RNase A residue ( $\chi_1$ =153°,  $\chi_2$ =80°), His97 (ONC) can be brought into hydrogen bond distance of the phosphate (2.62Å). Likewise, the invariant Lys31 (ONC) side chain is modeled with an all-trans conformation as observed for Lys41 (RNase A) and is within 3.2Å of the O(3') ribose of uridine.

In Figure 4.7, the uracil base O(2) and N(3) atoms are paired with the main chain N and side chain  $O^{\gamma 1}$  atoms of the invariant Thr35 (Thr45 in RNase A) in the B<sub>1</sub> subsite. These hydrogen bonds are 3.06Å and 2.43Å long in the ONC-d(UpG) model. The O(4) atom of the uracil base is directed towards Ser123 in RNase A. The equivalent residue in ONC is Val101. The nearest potential protein hydrogen bond acceptor or donor is Lys33N $^{\zeta}$  and it is 3.06Å from O(4) when Lys33 adopts an all-trans side chain

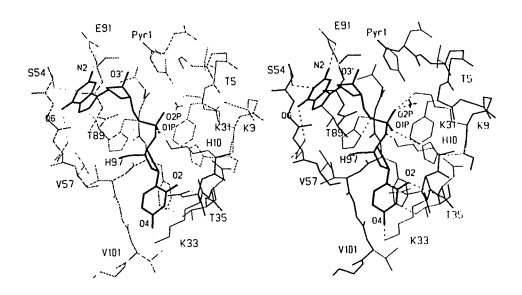


Figure 4.7a The active site of the ONC-d(UpG) model based on RNase A-d(CpA). The ONC side chains are shown as thin lines, the main chain as thicker lines and d(UpG) is shown as thick lines. Potential hydrogen bonds between ONC and d(UpG) are indicated as dashed lines. The ONC active site residues Lys9, His10, His97 and Phe98 make contacts equivalent to those observed in the RNase A-complex structure. Adjustment of the Lys33 side chain torsion angles brings Lys33N $^{\zeta}$  within hydrogen bonding distance of the uridine O4. The guanine base has three nearby hydrogen bonding partners; Ser54O $^{\gamma 1}$  (N1), Asn56N $^{\epsilon 2}$  (O6) and Glu91O $^{\epsilon 1}$  (N2).

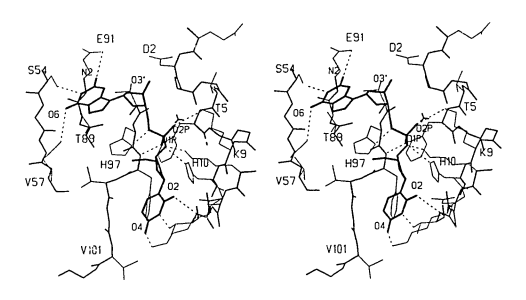


Figure 4.7b The rONC-d(UpG) model is shown in a manner equivalent to 7a. The guanosine deoxyribose packing interaction with Pyr1 (Figure 7a) is partially replaced by the Asp2 carboxylate in the rONC model. This change in the shape and charge of the ribose binding pocket may be responsible for rONC's lack of significant ribonucleolytic activity.

conformation. The guanine binding site (B<sub>2</sub>) in ONC is considerably different from that in RNase A as a result of a five residue insertion in RNase A (65-69) relative to ONC. In the RNase A-d(CpA) complex, adenine N(1) and N(6) atoms are paired with Gln69 and Asn71 in the B<sub>2</sub> binding site. In the ONC-d(UpG) model the N(1) and O(6) atoms of guanine are within hydrogen bond distance of Ser54OY<sup>1</sup> (2.7Å) and Asn56N<sup>6</sup> 2 (3.5Å). Structure based sequence alignments indicate Ser54 and Asn56 are equivalent to Thr70 and Cys72 (RNase A) (Mosimann *et al.*, 1994). ONC Glu91O<sup>£1</sup> is 2.6Å from the N(2) position of the base and provides a third hydrogen bond in the model. The conserved Glu111O<sup>£1</sup> is the equivalent residue in RNase A.

#### DISCUSSION

Significant main chain differences in the rONC and ONC protein structures are confined to the N-terminus. In rONC, the N-terminal helix (Glu1-Trp10) extends into a solvent channel. The fMet-1 residue is disordered and not seen in the electron density maps. The high B-factors and relatively weak density associated with Glu1 sidechain suggests a high degree of mobility and are consistent with fMet-1 being disordered. In ONC, the N-terminal helix ends at Trp3 and Pyr1 is folded back against this helix and the cyclized pyroglutamyl residue is directed into the active site.

The positional and conformational differences in the anion binding residues in rONC and ONC are small and the bound sulfate anion is only slightly shifted (0.29Å between sulfur atoms) in the two structures. In ONC, Pyr10<sup>e</sup> forms hydrogen bonds with Lys9N<sup>c</sup> (2.91Å) and Wat306 (2.66Å), respectively. While rONC does not have a structural counterpart to Pyr1, the Lys9N<sup>c</sup> and Wat258 positions differ by less than 0.5Å in the superposition. This suggests the anion binding residues of rONC are correctly positioned for catalysis. In the absence of additional structural information, there is no obvious structural reason for the rONC enzymatic inactivity. The ONC-d(UpG) and rONC-d(UpG)

models may explain rONC's inactivity. In the ONC-d(UpG) model, the guanine ribose O(3') is 3.9Å from Pyr1C\beta as the ribose packs against the Pyr1 ring in the R2 subsite. The C(2')-exo conformation of the ribose and the anti conformation of the glycosyl torsion angle allow the purine base to interact with up to three hydrogen bonding partners in the B2 subsite. In the rONC-d(UpG) model the wan der Waals contacts to the ribose are lost and are only partially replaced by the Asp2 carboxylate. Maximizing the non-bonded contacts in the R<sub>2</sub> subsite of rONC brings O(3') nearer to the Asp2 negative charge. Since the O(3') of the ribose of guanosine forms an ester bond with the anionic phosphate group in oligoribonucleotide substrates, an unfavorable electrostatic interaction may result. In addition, Asp $20^{82}$  is already 5.31Å from the Glu91  $\gamma$ -carboxylate that hydrogen bonds to N(2) of guanine in the model. Large conformational changes involving the ribose to improve van der Waals interactions or relieve electrostatic interactions may move the purine base from its binding pocket (B2). Together, the change in the shape and electrostatic nature of the ribose binding pocket of guanosine (R2) may give rise to rONC's lack of activity towards extended substrates. It is also possible the mobility of the N-terminus of rONC contributes to its lack of activity.

ONC has a  $k_{cal}/k_m$  of 4.73 x 10<sup>3</sup> M<sup>-1</sup>sec<sup>-1</sup> for UpG diribonucleotide substrates at its optimal pH of 6 (Boix et al., unpublished results). Under the same conditions, RNase A has a  $k_{cal}/k_m$  of 2.93 x 10<sup>5</sup> M<sup>-1</sup>sec<sup>-1</sup> for the same substrate. ONC has a marked preference for UpG but it will slowly hydrolyze CpG as well (W. Ardelt, unpublished results). In contrast, RNase A cleaves UpX and CpX (X = A,G,C,T) diribonucleotides and is least-active against UpG (Harper & Vallee, 1989). The superposition of ONC with the sulfate containing RNase A structure shows that ONC and RNase A bind the active site anion in equivalent positions. In the RNase A-d(CpA) structure and in the ONC-d(UpG) model, the N and OY¹ atoms of the invariant Thr (ONC Thr35; RNase A Thr45) form hydrogen bonds with the O(2) and N(3) atoms of the pyrimidine base (B<sub>1</sub>). This suggests that differences in the pyrimidine-base specificity of these two enzymes arise from differences in the binding of

the N(4) (cytosine)/O(4) (uracil) atom. In the RNase A-inhibitor structures the N(4) (cytosine) or O(4) (uracil) atom is directed at Ser123O $^{\gamma 1}$ . Direct hydrogen bonds between N(4) and Ser123O $^{\gamma 1}$  (Richards & Wykoff, 1971) and water-mediated interactions between O(4) and Ser123O $^{\gamma 1}$  (Wlodawer *et al.*, 1983; Borah *et al.*, 1985) have been reported. Since Ser123O $^{\gamma 1}$  and water molecules can either accept or donate hydrogen bonds, they account for the pyrimidine base preference of RNase A.

In the ONC-d(UpG) model (Figure 4.7) the uracil O(4) hydrogen-bonding potential is unsatisfied. There are three ONC atoms within 6Å of the uracil O(4) that are capable of participating in hydrogen bonds: Lys33N<sup>\(\zeta\)</sup>, Thr35O<sup>\(\gamma\)</sup> and Val101N. From previous crystallographic work, the invariant Thr35071 atom is known to interact with the N(3) position of pyrimidine bases (B<sub>1</sub>). Val101N is 4.11Å away and the conformation of this residue is fixed by the neighboring Cys87 to Cys104 disulfide bridge. In ONC, Lys33 and Val101 are equivalent to Val43 and Ser123 of RNase A. Adjustment of the Lys33 side chain torsion angles can bring the N<sup>\zeta</sup> atom to within 3.1Å of the uridine O(4) atom while retaining good stereochemistry. After adjusting the Lys33 torsion angles, the angle between C(4), O(4) (uridine) and Lys33N\(^{\zeta}\) is 127° which is close to the expected 120° hydrogen bonding angle. An interaction between Lys33N5 and N4 (cytosine)/O4 (uracil) provides an attractive explanation of the observed  $B_1$  site specificity of ONC. As Lys33N $^{\zeta}$ is only capable of donating hydrogen bonds at physiologically relevant pH, it would strongly favor the uridine O(4) atom (hydrogen bond acceptor) over the cytosine N(4) atom (hydrogen bond donor). The guanine specificity of ONC in the B2 site can also be explained by the nature of the residues in this subsite. Ser54 is 2.5Å from the guanine N(2) and can form hydrogen bonds with the N(2) atom of either purine base. In both ONC and rONC, Asn56N<sup>82</sup> hydrogen bonds with Thr89O<sup>71</sup> (2.66Å, 2.56Å). In this conformation, Asn $56N^{\delta 2}$  is available to interact with O(6) of guanine but not the N(6) of adenine. Similarly, the Glu91 carboxylate can productively interact with guanine N(2) (2.5Å) but not with adenine. The Asn56N $^{82}$  and Glu91O $^{\epsilon1}$  interactions are proposed to account for the guanine specificity of ONC.

The difference in the relative positions of the RNase A and ONC lobes (see Results) suggests the need for a conformational change in ONC during substrate binding and catalysis. The superposition of ONC and RNase A based on 45 out of 52 equivalent  $C^{\alpha}$ atoms of lobe 1 that can be overlapped within 3.8Å is justified by the improved agreement in the superposed positions of the active site residues of ONC and RNase A. The rmsd for the main chain and C<sup>β</sup> atoms of the active site residues (Lys9, His10, His97, Phe98) drops from 0.62Å to 0.32Å in superpositions based on the entire structure and the atoms of lobe 1, respectively. In Figure 4.5, lobe 2 of ONC is rotated relative to lobe 2 of RNase A and the active site cleft of ONC is more exposed. As a consequence, the Lys31 of ONC is 2.0Å ( $C^{\alpha}$ - $C^{\alpha}$  distance) from the required Lys41 of RNase A. In the RNase-d(CpA) complex, the Lys41 side chain adopts an all-trans conformation and hydrogen bonds to O(3') in the R<sub>1</sub> subsite. Lys31 of ONC is over 4Å from O(3') and must undergo a relatively large conformational change in order to make an equivalent contact. The difference in the lobe 2 positions of ONC and RNase A in Figure 4.5 is a hinge-like closing of approximately 8<sup>d</sup> (see Methods). In Figure 4.7, lobe 1 and lobe 2 of ONC were separately superposed with RNase A. In the resulting ONC-d(UpG) model, Lys31 has a position equivalent to Lys41 of RNase A and Lys33 is brought closer to the B<sub>1</sub> subsite. The closer approach of Lys33N<sup>\(\zeta\)</sup> to the uridine O(4) position supports the interpretation that Lys33N<sup>\(\zeta\)</sup> contributes to the pyrimidine specificity of ONC. Finally, this type of conformational change has not been observed in complexed and uncomplexed RNase A structures. In the recent crystallographic structure of angiogenin, the pyrimidine binding site (B<sub>1</sub>) is sterically blocked by the Glu117 side chain (Acharya, et al., 1994). As a consequence, substrate binding requires a conformational change that moves Glu117 and adjacent residues. Within the RNase superfamily, angiogenin also has a relatively small RNase activity (2%) in comparison with RNase A (Shapiro et al., 1986).

The authors realize that the ONC-d(UpG) and rONC-d(UpG) models developed

above cannot replace direct structural studies of inhibitor complexes. These models suggest differences in the shape and electrostatic nature of the R<sub>2</sub> result in the lack of rONC catalytic activity. In addition, the modeling has identified residues of the novel B<sub>2</sub> subsite and provided chemically reasonable explanations for the observed substrate specificity.

### MATERIALS AND METHODS

# Protein Purification and Crystallization

A synthetic rONC gene having an N-terminal sequence fMet\_1Glu<sub>1</sub>Asp<sub>2</sub>... has been expressed in an *Escherichia coli* expression system and the resulting protein has been purified from cell lysates (Wu, Y.N. & Youle, R.J., unpublished results). The identity of the amino-terminal residues has been confirmed by determining the amino acid composition and by N-terminal amino acid sequencing (W. Ardelt, unpublished data).

The purified rONC was dissolved in water (10 mg/mL; pH 6.0) and screened against a set of crystallization conditions, based upon a sparse matrix sampling method (Jancarik & Kim, 1991), using hanging drops. rONC was also screened against the crystallization conditions that produced native ONC crystals (Mosimann *et al.*, 1992) and small rosettes were obtained. Ultimately, the largest single crystals (up to 0.15 mm x 0.12 mm x 0.5 mm) were grown over a 20% w/v polyethylene glycol (PEG) 1000, 0.2 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.010 M acetate solution at pH 4.5. The initial hanging drop conditions contained 4 μL of 10mg/mL rONC and 6 μL of reservoir solution at pH 4.5.

#### Data collection

The unit cell and space group of the rONC crystals were determined from X-ray diffraction patterns and found to be isomorphous with the native ONC crystals. Data were collected by exposing the crystal to  $CuK_{\bar{\alpha}}$  radiation ( $\lambda$ =1.5418Å) produced by a Rigaku RU-200 rotating anode X-ray source operated at 40 kV and 150 mA. The X-ray diffraction

intensity data were measured on a San Diego Multiwire Systems (SDMS) multiwire proportional counter (Hamlin et al., 1985) while the crystal was moved through a series of ω-motor scans and φ, χ settings determined using the program LATTICEPATCH (Klinger & Kretsinger, 1989). The measured intensity data were corrected with Lorentz and polarization factors, scaled and multiple measurements of individual reflections were merged and averaged using the SDMS software. The data evere subsequently placed on an absolute scale. Table 4.3 lists the unit cell parameters, the completeness of the collected data and the agreement among symmetry equivalent structure factor amplitudes for the 2.2Å resolution data set.

#### Refinement

The initial phases for the rONC structure were derived from the native ONC coordinates (10NC; Mosimann et al., 1994) using residues 2-104. The initial rONC model was refined against the observed structure-factor amplitudes ( $|F_0| \ge 1 \sigma |F_0|$ ) between 10Å and 2.2Å. resolution using the simulated annealing slow-cooling protocol described in the X-PLOR manual (A. Brünger et al., 1987). This individual B-factor ( $B = 8\pi^2\overline{u^2}$ , where  $\overline{u^2}$  is the mean-square vibrational amplitude) refinement and positional parameter refinement resulted in a conventional crystallographic R-factor ( $R = \Sigma ||F_0| - ||F_0|| / \Sigma ||F_0||$ , where  $F_0$  and  $F_0$  are the observed and calculated structure factor amplitudes) of 0.213. Subsequent refinement was performed with TNT version 5C and 5D (Tronrud, 1992) using data between 20.0Å and 2.2Å resolution with  $|F_0| \ge 1 \sigma ||F_0||$ . Electron density maps were calculated using SIGMAA  $2mF_0-DF_0$  and  $mF_0-DF_0$  coefficients (Read, 1986) and using TNT  $3|F_0|-2|F_0|$ ,  $2|F_0|-|F_0|$  and  $|F_0|-|F_0|$  coefficients. Interactive graphics refitting sessions were performed using TOM 3.0 Alberta/CalTech (based upon Jones, 1985). Throughout the refinement procedure the stereochemical quality of the rONC model was monitored using PROCHECK (Laskowski, 1993).

Table 4.3

Summary of data collection statistics for Onconase(Met-1,Glu1) native data set+

	Resolution	No.	Unique	%			ι	ınit cell (Å	<b>(</b> )
	(Å)	Ref.	Ref.	Complete	I/ơIII	R <sub>m</sub> ‡	a	b	c
Native	2.2	36332	5066	98.0	12.0	0.053	41.14	69.37	32.53

<sup>+</sup> Space Group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>; wavelength (Cu $K_{\tilde{\alpha}}$ ) = 1.5418Å

 $<sup>^{\</sup>ddagger}R_{m} = \Sigma_{hkl}\Sigma |I_{obs} - I_{ave}|/\Sigma |I_{ave}|$ , where  $I_{obs}$  and  $I_{ave}$  are the observed intensity of a single reflection and the average intensity of all measurements of a reflection and its Freidel mate, respectively.

A sulfate ion bound in the active site region and the Glu1 residue were fit to the electron density maps calculated after the initial TNT refinement and Asp2 was refit. There was no electron density corresponding to fMet\_1 and this residue has not been included in the model. Ordered solvent molecules were added during interactive graphics refitting sessions. Solvent molecules were required to meet three conditions before being selected: (1) the |F<sub>0</sub>|-|F<sub>c</sub>| electron density map had a peak greater than 2.5 $\sigma$  at the proposed solvent site, (2) the electron density was roughly spherical in shape and (3) a productive hydrogen bond was formed with a protein atom (or with an already established ordered solvent molecule). All solvent molecules were subsequently refined at unit occupancy and were required to have refined B-factors less than 80Å2 (approximately 1.2 times the value of the largest B-factor of a protein atom). The electron density of the Glu1 side-chain remained discontinuous throughout the course of the refinement and electron density corresponding to Met was never observed. Prior to the final refinement cycles a model omitting Glu1 was subjected to a short refinement and the resulting model was used to calculate electron density maps (Bhat, 1988). The electron density in the region of Glu1 was not significantly different from that observed in electron density maps calculated with the complete model.

The final refined model of rONC contains 828 protein atoms, a sulfate anion and 72 ordered solvent molecules. As in the refinement of ONC, the anion was presumed to be  $SO_4^{2-}$  based upon its inclusion in the crystallization media (0.2 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>) and the absence of added phosphate to those solutions. The refined coordinates of several water molecules with high B-factors were fixed during the latter stages of refinement. These solvent molecules have refined hydrogen bond distances of slightly less than 2.5Å. Given the relatively weak electron density associated with these solvent molecules, they have been fixed at a hydrogen bonding distances of 2.5Å or greater.

Three amino acid side-chains have been modeled with alternate conformations. Ser 54 is modeled in two conformations while Thr60 and Thr83 are modeled in three

conformations. The occupancies of the various alternate conformations were fixed at 0.33 or 0.5 and each alternate conformation had to improve the fit between the model and the resulting electron density. These residues are all located on the surface of the rONC structure and extend into solvent channels.

All least-squares superpositions were performed using O (Jones *et al.*, 1991). The superposition of rONC and ONC was performed using the  $C^{\alpha}$  coordinates for residues 2-104. Superpositions involving RNase A (1RPG) made use of the published amino acid sequence alignment (Mosimann *et al.*, 1992) with the modifications described in Mosimann *et al.*, 1994. The superposition in Figure 4.5 was performed based on the 52 equivalent  $C^{\alpha}$  atoms of lobe 1 of ONC (residues 1-12, 41-62 and 85-102) and RNase A. In this superposition, lobe 2 of ONC is rotated relative to lobe 2 of RNase A. The angle between the antiparallel  $\beta$ -sheet (strands  $\beta$ 1,  $\beta$ 3 and  $\beta$ 4) of lobe 2 of ONC and RNase A is approximately 8°. For strands- $\beta$ 1 and  $\beta$ 4, the angle has been measured from using the N-terminal residue of the rONC strand, the C-terminal residue of the ONC strand and the N-terminal residue of the RNase A strand. For strand- $\beta$ 2, the angle has been measured from using the C-terminal residue of the rONC strand, the N-terminal residue of the ONC strand and the C-terminal residue of the RNase A strand. In ONC-d(UpG) model in Figure 4.7, lobe 1 and lobe 2 of ONC (residues 13-40 and 63-84) have been separately superposed with their counterparts from RNase A.

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### CHAPTER 5

# X-RAY CRYSTALLOGRAPHIC STRUCTURE OF RECOMBINANT EOSINOPHIL-DERIVED NEUROTOXIN AT 1.83Å RESOLUTION<sup>†</sup>

#### INTRODUCTION

Eosinophil-derived neurotoxin (EDN) is a small, basic ribonuclease (RNase) of the eosinophil granule that is a member of the pyrimidine-specific RNase superfamily of vertebrates (Gleich et al., 1986). It has been variably referred to as RNase Us, RNase II and non-secretory RNase (Sierakowska & Shugar et al., 1977) in the existing literature. Recombinant EDN (rEDN) is a neurotoxin that selectively kills cerebellar Purkinje cells (Gordon phenomenon; Durack et al., 1981), is a potent cytotoxin when injected into Xenopus oocytes (Saxena et al., 1991) and plays a prominent role in several human pathologies (Youle et al., 1993 and references therein). When recombinant EDN (rEDN) is conjugated to monoclonal antibodies raised against a cell surface receptor, it is a potent and specific cytotoxin with favorable immunogenic properties (Ryback et al., 1991). The homologous Onconase (Mikulski et al., 1989), eosinophil cationic protein (ECP; Fredens et al., 1982) and seminal RNase (D'Alessio et al., 1991) also display cytotoxic activity in vivo. Both ECP and Onconase (Newton et al., 1994) selectively kill Purkinje cells of the cerebellum. In animal models the neurotoxicity is expressed as muscle rigidity, incoordination, ataxia and spasmodic paralysis that are consistent with cerebellar deficit. Neuropathological studies show Purkinje cells are selectively sensitive to these RNases (Gordon phenomenon; Durack et al., 1981). Other homologous enzymes, ribonuclease A (RNase A) and angiogenin, are not cytotoxic. In order to extend our understanding of the

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structure and function of cytotoxic, vertebrate RNases, rEDN has been crystallized and its structure has been determined using molecular replacement techniques. The rEDN structure has been refined at 1.83Å resolution to a conventional R-factor (= $\Sigma ||F_0| - ||F_c|| / \Sigma ||F_0|$ , where  $||F_0||$  and  $||F_c||$  are the observed and calculated structure factor amplitudes, respectively) of 0.152 for all data with 1.83Å  $\leq d_{min} \leq 20$ Å and intensities  $||E_0||$ . The final molecular model of the crystallographic asymmetric unit contains an N-terminal formyl-methionine residue ( $||M_0||$ ), all 134 amino acids of EDN, two non-covalently bound sulfate anions and 121 ordered solvent molecules. The molecular conformation of rEDN and its interactions with the sulfate anion ligands are described. We also compare the conformation of rEDN to those of the cytotoxic Onconase (Mosimann et al., 1994) and the non-cytotoxic RNase A (Zegers et al., 1994).

## **METHODS**

The rEDN gene has been synthesized and the rEDN protein expressed as inclusion bodies in *Escherichia coli* BL21. The inclusion bodies have been denatured, renatured and dialyzed before a two step chromatographic purification protocol (D. Newton *et al.*, 1994). The recombinant enzyme is enzymatically active and does not have the post-translational modifications that are present in EDN isolated from human body fluids. Wild-type EDN is glycosylated at C2 of the indole of Trp7 (Hofsteenge *et al.*, 1995) and at five of the six Asn-X-Ser/Thr sequences (where X represents any amino acid except proline); Asn17, Asn59, Asn65, Asn74 and Asn85 are N-glycosylated whereas Asn92 is not glycosylated (Beintema *et al.*, 1988). The recombinant enzyme retains the cytotoxic activities of wild-type EDN.

Crystallization trials were performed using the vapor diffusion technique and hanging drops. A sparse-matrix screening for appropriate crystallization conditions (Jancarik & Kim, 1991) yielded crystals in (NH<sub>4</sub>)<sub>2</sub> SO<sub>4</sub> solutions near neutral pH. The crystal used for the high resolution data collection was grown over a 1 mL reservoir

solution containing 40% (NH<sub>4</sub>)<sub>2</sub>,SO<sub>4</sub> 0.1M sodium cacodylate at pH 6.5 and 5% ethanol. The hanging drop, containing initially 3  $\mu$ L of 21 mg/mL rEDN and 3  $\mu$ L of the reservoir solution, produced a rod-shaped crystal with dimensions of 0.60 x 0.15 x 0.15 mm. The rEDN crystals are orthorhombic (space group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>) and the unit cell dimensions determined from the high-resolution data set are: a=53.42(0.07)Å, b=57.35(0.08)Å, c=42.26(0.06)Å and  $\alpha=\beta=\gamma=90^{\circ}$ . The calculated solvent content is 40% and the specific volume (V<sub>m</sub>) is 2.04Å<sup>3</sup>/Da (Matthews. 1968).

The X-ray intensity data were collected locally on twin San Diego Multiwire Systems (SDMS) multiwire proportional counters. Graphite monochromated  $CuK_{\bar{\alpha}}$  X-rays were generated with a Rigaku Rotoflex RU-200BH rotating anode source operated at 40 kV and 150 mA. The data were collected and processed to 1.83Å using the SDMS software (Hamlin, 1985). Statistics for the processing of the native data set is presented in Table 5.1. In the 1.97Å to 1.83Å resolution range, 69% of the unique reflections have  $I \ge \sigma(I)$ .

The structure of rEDN was determined by the molecular replacement method using the AMORE suite of programs (Navaza et al., 1994). The rotation and translation function solutions were obtained using a truncated model of RNase A. In the search model, residues 17 to 26 were deleted, residues 2, 7, 10, 29, 31, 76, 79 and 98 were truncated to alanine and residues 13, 33, 37, 61, 66, 85 and 90 were modeled as serine. The deleted residues of the RNase search model correspond to an insertion in the RNase A sequence relative to the EDN primary sequence. The residues that were modeled as alanine or serine have large, solvent exposed side chains that are not conserved in the alignment of the sequences of the two enzymes. With this search model, the correct rotation function solution was the fourth largest peak when using data extending from 12 to 2.5Å resolution and an integration radius of 24Å. The correct translation function solution has the highest correlation coefficient (0.386) and the lowest R-factor (52.4). Its correctness was evaluated using packing functions and by calculating and interpreting electron density

Table 5.1

Summary of data collection statistics for the rEDN native data set+

	ur a	nit cell (Å	) c		No. Refln.		% Complete	I/σ(I)	R <sub>m</sub> ‡
Native	53.42	57.35	42.26	1.83	58062	10840	90.8	12.3	0.053

<sup>+</sup> Space Group P2<sub>1</sub>2<sub>1</sub>2<sub>1</sub>; wavelength (Cu $K_{\tilde{\alpha}}$ ) = 1.5418Å

 $<sup>\</sup>ddagger R_m = \Sigma_{hkl}(\Sigma | I_{obs} - \langle I \rangle / \Sigma \langle I \rangle)$ , where  $I_{obs}$  and  $\langle I \rangle$  are the observed intensity of a single reflection and the average intensity of all measurements of a reflection and its Freidel mate, respectively.

maps.

The TOM Alberta/Caltech 2.8.0 interactive graphics program (based on Jones, 1985) was utilized to fit the molecular replacement search model to the electron density. The fMet and Gln116 to Pro122 amino acids were the only residues that could not be fit unambiguously into the initial electron density maps. Data between 8.0 and 1.83Å resolution with intensities greater than  $2\sigma$  were selected for the initial stages of refinement using the simulated annealing protocol described in the X-PLOR manual (Brünger et al., 1987). During this procedure the crystallographic R-factor ( $R = \Sigma I |F_0| - |F_c| / \Sigma |F_0|$ , where  $|F_0|$  and  $|F_c|$  are the observed and calculated structure factor amplitudes) was reduced from 0.519 to 0.263. Two sulfate anions and the segment of residues Gln116 to Pro122 were fit into a  $3|F_0| - 2|F_c|$  electron density map calculated after the molecular dynamics refinement. All subsequent steps in the refinement of rEDN made use of the TNT program suite (Tronrud, 1992) and data in the range  $1.83\text{Å} \leq d_{\text{min}} \leq 20\text{Å}$  with intensities greater than  $1\sigma$ . Interactive graphics refitting sessions utilized electron density maps calculated with  $3|F_0| - 2|F_c|$ ,  $2|F_0| - |F_c|$  and  $|F_0| - |F_c|$  coefficients obtained from TNT.

At several points during the refinement, standard omit maps were calculated (Bhat et al., 1988). In these maps selected atoms were omitted from a short refinement (5 cycles) and the subsequent map calculation. The resulting electron density in the region of the omitted atoms represents a relatively unbiased view of the true electron density at these positions. The residues Ser64 to Lys66, Pro88 to Ile93 and the sulfate anions were all checked against their density in these maps at several stages of the refinement. The side chains of Asn25, Asn39 and Asn104 were also examined in similar omit maps and subsequently modeled as having alternate conformations in the rEDN structure. At occupancies of 0.3 and 0.7 for the two alternate conformations, the average side chain B-factors of each of the conformations are between 17Å<sup>2</sup> and 26Å<sup>2</sup>. The N-terminal <sup>f</sup>Met residue was built into |F<sub>0</sub>|-|F<sub>c</sub>| difference electron density towards the end of the refinement. Throughout the refinement no effort was made to refine water molecules with alternate

positions and/or variable occupancies. The final model contains 121 ordered, full-occupancy water molecules and has an R-factor of 0.152 for all data with  $I/\sigma(I) \ge 1.0$  from 20 to 1.83Å resolution. The stereochemical parameters associated with the final cycle of least-squares refinement are given in Table 5.2. The rEDN coordinates will be deposited in the Protein Data Bank at the Brookhaven National Laboratory (Bernstein *et al.*, 1977).

The least-squares superpositions of RNase A, Onconase and rEDN were computed (Kabsch, 1978) using the O program suite (Jones *et al.*, 1991). The active sites of rEDN and RNase A were superposed using the main-chain and  $C^{\beta}$  atoms of the Thr6 to His15, Arg36 to Thr42 and Val126 to Ile133 residues of rEDN and the equivalent atoms of RNase A (Zegers *et al.*, 1994).

#### RESULTS

# Quality of the refined structure

The final (3|F<sub>0</sub>|-2|F<sub>c</sub>|)  $\alpha_{calc}$  electron density is continuous and well-defined for all main-chain atoms of residues 1 to 89 and 92 to 134 at the 1 $\sigma$  contour level. The N-terminal formyl-methionine (fMet) and residues Pro90 and Gln91 have discontinuous and relatively weak electron density. All side-chain atoms of residues in the hydrophobic core and nearly all side-chain atoms of residues on the surface of rEDN have continuous electron density at the same contour level. However, the solvent-exposed side chains of Gln40, Lys66, Gln91 and Arg117 are disordered beyond C7 and have no associated electron density at the 0.9 $\sigma$  contour level. The two non-covalently bound sulfate anions (SO<sub>4</sub>(A) and SO<sub>4</sub>(B)) have distinct lobes of electron density for their respective oxygen atoms. Figure 5.1a shows the (|F<sub>0</sub>|-|F<sub>c</sub>|)  $\alpha_{calc}$  difference electron density contoured at 2.5 $\sigma$  and 8 $\sigma$  for both sulfate anions after refinement with the atoms of the sulfate anions omitted from the structure factor calculations. The refined sulfate anion positions are also displayed in

Table 5.2

Stereochemical statistics for the final cycle of restrained least-squares refinement

R-factor resolution	0.152 for 10627 reflection	ons > 1.0oiII	between 20.0 - 1.83Å
Model			B-factor (mean)
	541 main-chain atoms		12.7 Å <sup>2</sup>
	548 side-chain atoms		17.4 Å <sup>2</sup>
	121 waters		$38.2  \text{\AA}^2$
	2 sulfate anion		42.7 Å <sup>2</sup>
Stereochemistry		rmsd‡	Restraint§
	1 - 2 bond distance	0.014Å	0.70
	Bond Angles	1.87°	0.55
	Pseudorotation Angles	17.11°	0.50
	Trigonal planar groups	0.013Å	1.50
	General planar groups	0.009Å	2.50
	non-bonded contacts	0.019Å	6.00
	Temperature factor	$3.04$ Å $^2$	0.15

<sup>‡</sup>rmsd =  $\sqrt{\sum (X_x - \overline{x})^2 / m}$ ; the root mean square deviation from expected values in small molecule structures

<sup>§</sup>Restraint is the relative weight assigned to each stereochemical parameter during the final cycle of refinement.

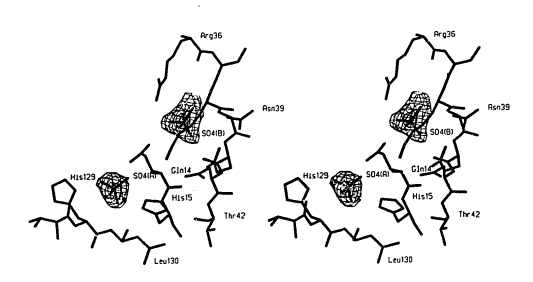


Figure 5.1a The (IF<sub>0</sub>I-IF<sub>c</sub>I)α<sub>calc</sub> electron density associated with the sulfate anions after a short refinement and map calculation with the contribution of the sulfate anions omitted. The density is contoured at 2.5σ and 8σ. The figure includes a stick model of the refined rEDN active site and sulfate anions. The are distinct lobes of electron density corresponding to the oxygen atom positions for each sulfate anion. The side chain conformation of Asn39 is one of the two alternate conformations for this side chain.

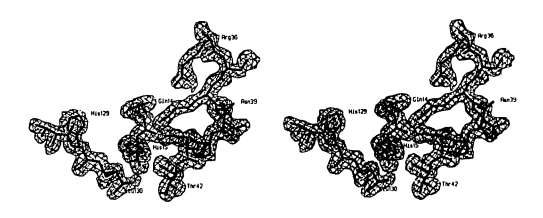


Figure 5.1b The (2|F<sub>0</sub>|-|F<sub>c</sub>|)α<sub>calc</sub> electron density associated with the active site residues of rEDN. The active site residues are shown as thick lines and the density is contoured at 1σ. The two alternate conformations of the Asn39 side chain are shown. The Gln40 side chain is partially disordered and does not have associated electron density beyond Cγ.

Figure 5.1a superposed on the electron density. Fig5.1b shows the ( $2|F_o|-|F_c|$ )  $\alpha_{calc}$  electron density associated with the active site residues contoured at  $1\sigma$ .

Figure 5.2 is a plot of the average B-factor of main-chain atoms ( $\langle B_{mc} \rangle$ ; (B =  $8\pi$  ,u<sup>2</sup>), where  $\overline{u^2}$  is the mean-square vibrational amplitude) and average side-chain B-factor ( $\langle B_{sc} \rangle$ ) versus amino acid sequence number. In both plots, the residues Ser64 to Lys66 and Thr87 to Ile93 possess the largest average B-factors in the structure. These residues are part of two large, solvent-exposed omega loops (Gly56-Lys69, Thr88-Cys96) and include two of the residues with partially disordered side chains. Lys66 and Glu91. The remaining residues with  $\langle B_{mc} \rangle \geq 20\,\text{Å}^2$  are Gly75, Ser76, Glu116 to Arg118, the N-terminal fMet and the C-terminal Ile133, Ile134. Residues Gly75 and Ser76 are part of a loop structure and residues Glu116 to Arg118 are part of a helical turn.

Figure 5.3 shows the Ramachandran plot for the 118 non-glycine, non-proline residues of rEDN. The main-chain  $\phi,\psi$  dihedral angles populate the core (103 of 118 residues) and allowed (15 of 118 residues) regions of the Ramachandran plot as defined in PROCHECK (Laskowski *et al.*, 1993). The EDN primary sequence contains 2 glycine and 12 proline residues. The NH of Gly56 is 2.78Å from Asn53O and Gly56 adopts a 3<sub>10</sub>-helical conformation ( $\phi = -84^{\circ}$ ,  $\psi = -9^{\circ}$ ) at the end of helix α2 (residues Thr47-Cys55). Gly75 has a conformation ( $\phi = 84^{\circ}$ ,  $\psi = -49^{\circ}$ ) not normally allowed to nonglycine residues and Gly75O forms a single intramolecular hydrogen bond with His73Nε2 (3.06Å). The 12 proline residues each adopt *trans*-peptide conformations and have  $\phi$ ,  $\psi$  dihedral angles typical of proline residues. Eight of the proline residues are found in loop structures and four in segments of β-strand. Pro102, Pro124 and Pro127 are parts of solvent accessible β-strands on the edge of their β-sheet. Pro79 is the second residue of a central β-strand and is also solvent exposed.

# Molecular conformation of rEDN

The polypeptide fold of rEDN is shown in Figures 5.4 and an all atom stereo

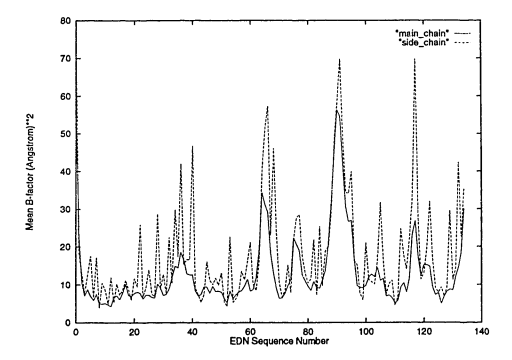


Figure 5.2 The average main-chain B-factor (solid line) and average side-chain B-factor (dashed line) versus sequence number plots for the rEDN structure. The segments Ser64 to Lys66 and Thr87 to Ile93 have the largest average main-chain B-factors and are parts of two long loop structures. The side chains of Lys66, Gln92 and Arg117 are disordered beyond C7 and represent the largest average side-chain B-factors.

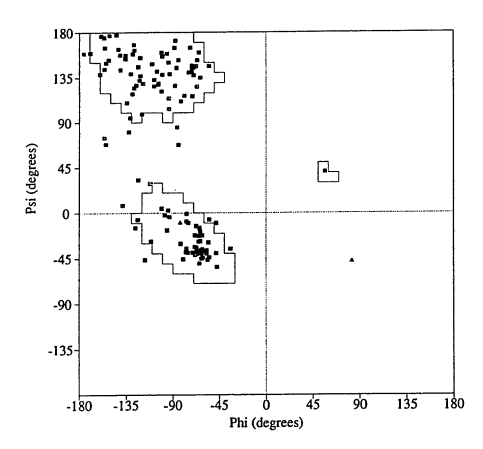


Figure 5.3 A Ramachandran plot of the φ, ψ dihedral angles of rEDN. The two glycine residues of rEDN are indicated by filled triangles and the remaining residues of rEDN are indicated by filled squares. The contours represent the most favorable regions of the plot as defined in PROCHECK (Laskowski et al., 1993) and include 103 of the 118 non-glycine, non-proline residues in rEDN. The remaining 15 residues are in the "allowed" regions of the plot (contour not shown).

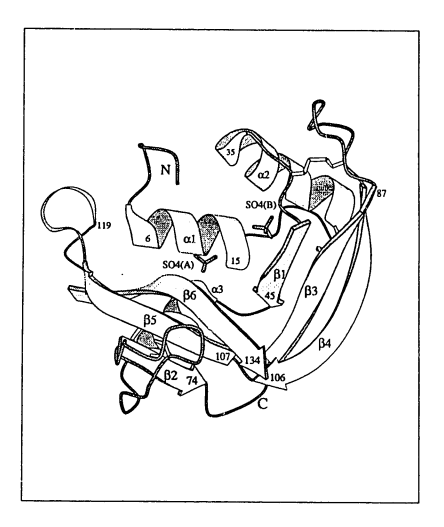


Figure 5.4 A molscript (Kraulis, 1991) diagram of the rEDN polypeptide fold. The helices, strands and loop structures of rEDN are shown as helical ribbons, arrows and filled lines, respectively. The four disulfide bridges and the two non-covalently bound sulfate anions are represented as rods. The helical turn Asp115 to Asp119 (shown as a one turn helix) is part of a nine residue insertion that is unique to EDN.

representation is presented in Figure 5.5. Table 5.3 lists the secondary structural elements present in the rEDN structure as defined by PROCHECK and Table 5.4 lists the  $\beta$ -turns and identifies omega loops (Sibanda & Thronton, 1985; Leszcz; 1ski & Rose, 1986). rEDN is comprised of an N-terminal  $\alpha$ -

helix and two related lobes. Each lobe contains a three-stranded antiparallel  $\beta$ -sheet and an  $\alpha$ -helix with a conserved disulfide bridge between the central strand of the  $\beta$ -sheet and the  $\alpha$ -helix. The strands  $\beta 1$ ,  $\beta 3$ ,  $\beta 4$  and helix  $\alpha 2$  form lobe 1 and strands  $\beta 2$ ,  $\beta 5$ ,  $\beta 6$  and helix  $\alpha 3$  comprise lobe 2. The conserved disulfide bridges of the two lobes are Cys23-Cys83 (lobe 1) and Cys55-Cys111 (lobe 2) and they share left-handed  $\chi 3$  torsion angles of -81° and -78°, respectively. Helix  $\alpha 1$  packs predominantly against lobe 2 where it contributes residues to the hydrophobic core. Residues at the C-terminus of helix  $\alpha 1$  (Gln14, His15, Ile16) pack against amino acids of lobe 1. The active site of rEDN is located at the junction of lobe 1, lobe 2 and the N-terminal  $\alpha$ -helix where a cleft is formed running parallel to strand  $\beta 6$ .

The remaining cysteine residues of rEDN form two additional disulfide bonds that help define the tertiary structure. The disulfide Cys37-Cys96 is between the loop following helix  $\alpha 2$  and the central strand ( $\beta 3$ ) of the  $\beta$ -sheet of lobe 1. The disulfide Cys62-Cys71 is part of the large loop (Gly56-Lys69) between helix  $\alpha 3$  and strand  $\beta 2$ , a strand at the edge of the  $\beta$ -sheet of lobe 2. These two disulfide bridges have right-handed  $\alpha 3$  torsion angles of 88° and 104°, respectively.

The stretch of polar residues from Asp115 to Tyr123 (Asp-Gln-Arg-Arg-Asp-Pro-Pro-Gln-Tyr) forms a single helical-turn followed by a type III  $\beta$ -turn like structure (Figure 5.6). This compact folding unit corresponds to an omega loop. Asp115O forms a hydrogen bond with Asp119N (2.89 Å) and Asp115O $^{\delta 1}$  hydrogen bonds to Arg118N (2.54 Å) initiating helical turn. Five additional polar interactions stabilize the conformation of this loop: Arg114N $^{\eta 1}$  donates a hydrogen bond to Gln116O (2.82Å); Arg118 $^{\eta 1}$  is 2.97 Å from Pro3 O; Asp119 O $^{\delta 2}$  accepts a hydrogen bond from Asp115N (2.84Å) and

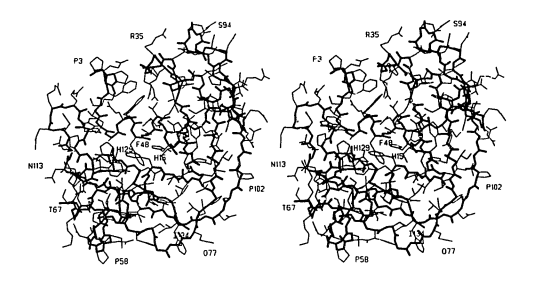


Figure 5.5 A stereo diagram of the all-atom structure of rEDN. The bonds between the main-chain atoms are shown as thick lines; those connecting the side-chain atoms are thin lines. The view is the same as that in Figure 5.4. The active site His15 and His129 and other selected residues are indicated.

Table 5.3
Secondary structures of rEDN

	Helices		Strands		loop structures		
α1	Thr6-Ile16	β1	Asn39-Leu45	11	Lys1-Phe5		
α2	Gln22-Arg35	β2	Asn70-Asn74	12	Asn17-Gln21		
α3	Thr47-Cys55	β3	Val78-Thr87	13	Arg36-Lys38		
		β4	Tyr97-Met104	14	Gly56-Lys69 <sup>‡†</sup>		
		β5	Phe105-Arg114	15	Gly75-Gln77		
		β6	Pro124-Ile134	16	Pro88-Cys96‡†		
				17	Asp115-Tyr123		

<sup>‡</sup> Residues Pro2 to Phe5, Asn62 to Asn65, Asn65 to Arg68 and Gln92 to Asn95 are type I β-turns.

<sup>§</sup> Residues Asp115 to Asp119 form a helical turn and Pro120 to Tyr123 form a type III  $\beta$ -turn.

<sup>†</sup> Loops 1 4, 1 6, and 1 7 are omega loops (Leszczynski & Rose, 1986).

Table 5.4
β-turn structures of rEDN

Туре	Residues i, i+1, i+2, i+3	φ <i>i+!</i> , Ψ <i>i+1</i>	φ <sub>i+2</sub> , ψ <sub>i+2</sub>
I	Pro2 Pro3 Gln4 Phe5	-66°, -21°	-79°, -7°
I	Cys62 Pro63 Ser64 Asn65	-54°, -35°	-85°, -10°
I	Asn65 Lys66 Thr67 Arg68	-46°, -11°	-98°, -1°
I	Asn92 Ile93 Ser94 Asn95	-42°, -33°	-66°, -12°
IΠ	Pro120 Pro121 Gln122 Tyr123	-59°, -26°	-44°, -43°

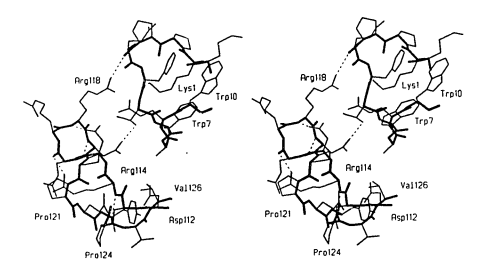


Figure 5.6 A stereo diagram of the residues of Asp115 to Tyr123 of rEDN and their packing interaction with the N-terminus. The main-chain and side-chain bonds are represented by thick and thin lines, respectively and hydrogen bonds are indicated as dashed lines. The residues of Asp115 to Asp119 form a single helical turn and Pro120 to Tyr123 forms a type III β-turn. The side chain of Asp119 forms a hydrogen bond to the side chain of Thr6 and the negatively charged carboxylate balances the positively charged dipole moment of the helix α1.

Asp $1190^{\delta 1}$  is positioned opposite the dipole of helix  $\alpha 1$  where it also forms a hydrogen bond with Thr6 OY (2.68 Å). The segment Pro120 to Tyr123 breaks the helix and adopts a type III turn like conformation that changes the direction of the polypeptide chain and leads into strand  $\beta 6$ .

The loop 17, and in particular the helical turn, packs against and influences the conformation of N-terminus of EDN. The side chains of Asp115, Arg118 and Asp119 pack against the N-terminus of helix  $\alpha$ 1. Helix  $\alpha$ 1 is initiated by helix-capping hydrogen bonds from the Thr6 OY1 and O atoms to Gln9N and Trp10N, respectively. The conformation of Thr6 is further fixed by the Thr6N to Gln9 OE1 hydrogen bond. Residues Pro2 to Phe5 adopt a type I  $\beta$ -turn conformation and the Lys1 N $\zeta$  forms a hydrogen bond with Phe5O. The conformation of the N-terminus is also stabilized by aromatic stacking interactions between the Phe5 and Trp10 rings (Burley & Petsko, 1986). The side chain of fMet does not interact with residues of the N-terminus or the residues Asp115 to Tyr123.

The omega loop, 14 of rEDN contains a disulfide bridge (Cys62-Cys71) and consecutive type I  $\beta$ -turns. In loop 14, Asn65N and Asn65O are 3.1Å and 2.8Å from Cys62O and Arg68N, respectively. Another omega loop, 16 contains Cys96 of the conserved Cys37-Cys96 disulfide and a single type I  $\beta$ -turn. The O atom of Asn92 of loop 16 is 2.5Å from Asn95N. The only other non-helical, mainchain i to i+3 hydrogen bond in the rEDN structure is between Ile93O and Cys96N (3.0Å).

### Sulfate anion binding sites

Figure 5.7 shows the hydrogen bonding network between rEDN and the two, non-covalently bound sulfate anions (SO<sub>4</sub>(A) and SO<sub>4</sub>(B)). Table 5.5 presents a list of the hydrogen bonds to the anions. The anions are presumed to be sulfate based upon the presence of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (and the absence of phosphate) in the crystallization media. The SO<sub>4</sub>(A) anion binds at the active site of rEDN where Gln14N<sup>E2</sup> and His15N<sup>E2</sup> of helix  $\alpha$ 1 are 3.37 Å and 3.08 Å from O4 and O1 of the SO<sub>4</sub>(A) anion,

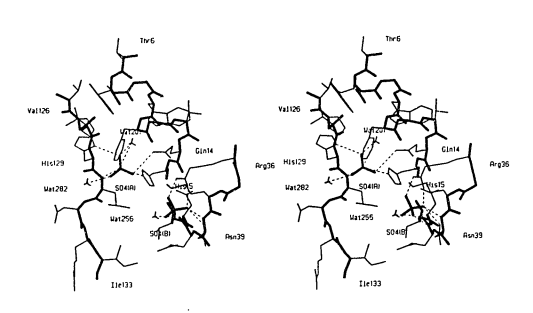


Figure 5.7 The interactions between rEDN and the non-covalently bound SO<sub>4</sub>(A) and SO<sub>4</sub>(B) anions are shown in a stereo diagram. The thick lines represent the main chain of rEDN the thin lines represent the side chains of rEDN and its anions and hydrogen bonds are represented as dashed lines. The SO<sub>4</sub>(A) anion at the active site contacts the side chains of Gln14, His15 and His129 and the main chain of Leu130. The anion SO<sub>4</sub>(B) is bound by the main chain of Asn39 and Gln40 and the side-chain atoms of Arg36 and Asn39. Ordered solvent molecules that make contacts with either anion are shown as small tetrahedra.

Table 5.5

The sulfate anion contacts distances (Å) in EDN

SO <sub>4</sub> (A)		SO <sub>4</sub> (B)	
Ol		O1	
His15N <sup>E2</sup>	3.08	Arg36N <sup>n2</sup>	2.50
Leu130N	2.61		
Wat201	3.13		
O2		O2	
Wat282	2.73	Gln40N	3.12
His $129N^{\delta 1}$	3.33	Wat256	3.06
O3		О3	
His129N <sup>δ1</sup>	2.63	†Asn39Nδ2	3.08
Wat201	2.96		
04		O4	
Gln14N <sup>€2</sup>	3.23	Arg36N <sup>n1</sup>	2.97
His15N <sup>E2</sup>	3.47	Asn39N	2.60
		$^{\dagger}$ Asn39N $^{\delta2}$	2.69

<sup>&</sup>lt;sup>†</sup> The Asn39 side chain is modeled as two separate conformations in the rEDN structure. The Asn39 side chain conformation that contacts the SO<sub>4</sub>(B) is the minor conformer (occupancy=0.3).

respectively. The remaining two hydrogen bonds between rEDN and SO<sub>4</sub>(A) are from His129 N<sup> $\delta$ 1</sup> (2.63Å) and Leu130 N (2.61Å) of strand  $\beta$ 6 to O3 and O1 of the anion, respectively. Two ordered solvent molecules, Wat201 and Wat282 also contact the bound sulfate anion. Wat201 accepts a 3.05Å hydrogen bond from Gln14N<sup> $\epsilon$ 2</sup> and is  $\leq$ 3.30Å from O1 and O3 of the anion. This water has counterparts in both the RNase A and the Onconase structures. Wat282 forms a 2.73Å hydrogen bond with O2. The side chain of the catalytically important Lys38 is well-defined in the electron density maps and the N<sup> $\zeta$ </sup> atom forms a hydrogen bond with the Gln14O<sup> $\epsilon$ 1</sup>. In this conformation, Lys38N<sup> $\zeta$ </sup> is 3.9Å from O4 of the anion.

The anion SO<sub>4</sub>(B) is bound to the surface of a single rEDN molecule by three main-chain hydrogen bonds and the side-chain groups of Arg36 and Asn39. Gln40N is 3.12Å from O2 and Asn39N is 2.60Å and 3.28Å from O4 and O2, respectively. Arg36 Nm1 and Arg36Nm2 are within hydrogen bond distance of O4 and O1, respectively. The alternate side-chain conformation (occupancy = 0.30) of Asn39 provides two additional contact with O3 and O4. Wat256 is the only ordered solvent molecule hydrogen bonding with SO<sub>4</sub>(B). The distance between the two sulfur atoms of SO<sub>4</sub>(A) and SO<sub>4</sub>(B) is 9.4Å. The second anion binding site has not been observed in structures of other members of the RNase superfamily.

## Intermolecular packing in rEDN

Figure 5.8 is a stereo diagram representing the intermolecular packing within the rEDN crystal. Table 5.6 lists the polar hydrogen-bonding interactions and van der Waals contacts between symmetry-related molecules. Two hydrogen bonds are accepted by Gln12O<sup>ε1</sup> (Arg97N<sup>ε</sup>, Arg97N<sup>η2</sup>) and by Arg35O (Arg114N<sup>ε</sup>, Gln116N<sup>e2</sup>), respectively and Gln21 N<sup>ε2</sup> donates hydrogen bonds to Cys55O Asn57O. Thr61 also forms a pair of hydrogen bonds and is involved in the sole polar intermolecular contact between mainchain atoms (Thr61N to Thr86O). Gln12, Gln21, Arg35 and Thr61 account for 8 of the

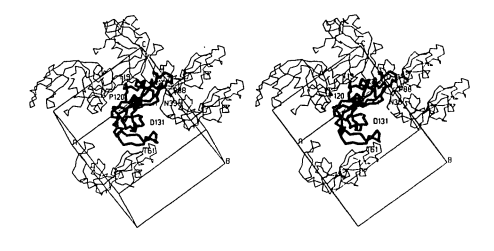


Figure 5.8 A C<sup>α</sup> backbone trace representing the intermolecular packing in the rEDN crystal is shown as a stereo diagram. The rEDN molecule at x, y, z is shown in thick lines and the five symmetry related molecules that make contacts with it are represented as thin lines. Residues in the rEDN molecule in thick lines are labeled as are the unit cell axes.

Table 5.6

The intermolecular contacts in EDN

Polar <3.30 Å		van der Waals < 3.90 Å	
$\frac{-x+1/2, -y, z+1/2}{-x+1/2}$			
-x+1/2, -y, $z+1/2Thr13 OY1 - Gln100 NE2$	3.29	Phe5 $C^{\delta 2}$ - Gln22 $N^{\epsilon 2}$	3.11
	3.13	Asn 17 N $^{82}$ - Ala 99 C $^{\alpha}$	3.74
Gln12 O <sup>E1</sup> - Arg97 N <sup>n2</sup>	2.87	Val29 C <sup>yl</sup> - Gln100 O	3.40
Gln12 O <sup>E1</sup> - Arg97 N <sup>E</sup>	2.01		3.84
		Tyr33 C <sup>ζ</sup> - Pro102 C <sup>γ</sup>	3.41
		Tyr33 OH - Gln100 CY	
		Phe48 $C^{\delta 2}$ - Lys66 O	3.52
-x+1, y-1/2, -z+1/2			2.24
Ile16 O - Lys69 N <sup>ζ</sup>	3.29	Ser19 O - Pro58 CY	3.31
Gln21 $N^{\epsilon 2}$ - Asn57 O	2.69	Thr47 CY2 -Lys66 C	3.65
Gln21 N <sup>2</sup> - Cys55 O	3.15		
Thr46 O - Lys69 N <sup>G</sup>	3.27		
Asn50 $O^{\delta 1}$ - Thr67 O	2.96		
x-1/2, $-y+1/2$ , $-z+1$			
Arg35 O - Gln116 Nε2	3.19	Pro3 C - Gly56 Cα	3.80
Arg35 O - Arg114 N <sup>n</sup> <sup>2</sup>	3.06	Pro3 C <sup>β</sup> - Val52 O	3.32
Mg55 O - /Mg11411		Gln4 C <sup>y</sup> - Gly56 O	3.70
		Tyr33 O - Pro124 C <sup>β</sup>	3.45
		Arg35 C <sup>y</sup> - Pro124 O	3.61
. 1 /2 . 1 /2 -	•	A1833 C1 - F10124 O	
x+1/2, -y+1/2, -z	2.02	Met60 C <sup>β</sup> - Thr86 O	3.44
Thr61 N - Thr86 O	2.92	Metou CP - 1 nr86 U	J. TT
Thr61 Oyl - Thr86 Oyl	3.16		

12 unique intermolecular hydrogen bonds and each residue is in contact with a different symmetry related molecule. The Pro3 and Gln4 residues at the N-terminus and Tyr33 account for 6 of the 14 van der Waals contacts <3.90 Å. Four additional proline residues Pro58, Pro88, Pro102 and Pro124 also participate in packing interactions. The shortest intermolecular van der Waals contact is  $3.11\text{\AA}$ , Phe5C $^{\delta2}$  to Gln22N $^{\epsilon2}$ .

### Superposition of rEDN with RNase A and Onconase

Figure 5.9 is a stereo diagram of the least-squares superposed X-ray crystallographic structures of rEDN, RNase A and Onconase. The superpositions of rEDN with RNase A and Onconase have rmsds of 1.27 Å and 1.42 Å for the 106 (of 118) and 86 (of 101) common  $C^{\alpha}$  atoms that can be superposed within 3.8 Å, respectively. The residues in helices  $\alpha 1$  to  $\alpha 3$  and those in strands  $\beta 1$  to  $\beta 6$  account for nearly all the equivalent residues in the three structures. The residues Ile30 to Arg35 of rEDN extend helix  $\alpha 2$  relative to RNase A and are the only residues of these secondary structures that do not superpose with their counterparts in RNase A and Onconase. The loops 11, 12, 14, 16 and 17 (Table 5.3) are the sites of insertions and deletions in the three structures and include those remaining residues that do not superpose within 3.8 Å in the overlaps. The conformation and intramolecular packing between the N-terminus (loop 11) and the 9 residue insertion of (helix  $\alpha 4$  and loop 17) rEDN have been described. The loop 12 of rEDN is 6 residues shorter and 1 residue shorter than the equivalent loops in RNasc A and Onconase, respectively; the conformation of 12 is completely different in all three structures (Figure 5.9). The relatively large loops, 14 and 16 of rEDN are 2 residues longer than the equivalent loops in RNase A. In each case, the additional residues of rEDN result in conformational differences along the entire length of the loop. The loops 14 and 16 of Onconase are 8 residues shorter and 6 residues shorter than the equivalent loops of rEDN and have very different conformations in RNase A and Onconase (Mosimann et al., 1995).

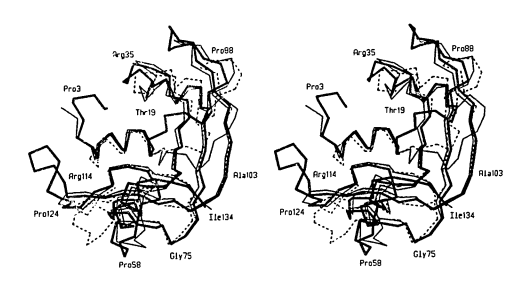


Figure 5.9 The least-squares superposition of rEDN, RNase A (Zegers et al., 1994) and Onconase (Mosimann et al., 1994) based on their  $C^{\alpha}$  coordinates rEDN, RNase A and Onconase are represented by thick, thin and dashed lines, respectively. The  $\alpha$ -helices and  $\beta$ -strands of the structures superpose relatively closely whereas the main conformational differences are in the connecting loop structures.

Figure 5.10 shows the superposed active sites of RNase A (Zegers et al., 1994) and rEDN. The mainchain and  $C^{\beta}$  atoms of residues Thr6-His15, Arg36-Thr42 and Val126-Ile133 of rEDN were superposed with the equivalent residues of RNase A by least-squares. The amino acid exquence identity for these 25 residues is 64% and the rmsd is 0.60Å for 125 common atoms. The sulfate anion of the RNase A structure and SO<sub>4</sub>(A) of rEDN have nearly identical conformations and their sulfur atoms differ in their positions by 0.6Å. This difference is less than the 0.9Å that separates Asp121C<sup> $\alpha$ </sup> (RNase A) and Asp131C<sup> $\alpha$ </sup> (EDN). The side chains of Gln11, His12, His119 and the main chain of Phe120 of RNase A form hydrogen bonds with the anion. Gln14, His15, His129 and Leu130 of rEDN make equivalent contacts with SO<sub>4</sub>(A). His119 of RNase A adopts two alternate conformations in the RNase A structure (Zegers et al., 1994). The His129 conformation of rEDN corresponds to the catalytically inactive conformation of His119 of RNase A. His129 can adopt the active conformation observed in RNase A by rotation about its  $\chi_1$  and  $\chi_2$  torsion angles. In order for His129 to adopt the active conformation, Wat282 would have to be displaced (Figure 5.7).

In the superposition of RNase A and EDN, the SO<sub>4</sub>(B) of rEDN is less than 1Å from the side chain of Pro42 of RNase A. Pro42 of RNase A is the residue equivalent to Asn39 of rEDN. In the crystal structure of rEDN, the NH and N<sup> $\delta$ 2</sup> atoms of Asn39 donate hydrogen bonds to SO<sub>4</sub>(B) (Figure 5.7). Obviously, Pro42 of RNase A cannot act as a hydrogen bond donor and as a result, RNase A cannot bind an anion in a manner equivalent to the SO<sub>4</sub>(B) binding observed in rEDN.

rEDN has been superposed with the RNase A-thymidylic acid tetramer (d(Tp)4) complex structure (Birdsall et al., 1992) by least squares. Figure 11 is a stereo diagram of rEDN and three and of the four superposed thymidilic acids. The base, ribose and phosphate moieties shown in Figure 11 are B<sub>2</sub>, B<sub>1</sub>, B<sub>0</sub>, R<sub>2</sub>, R<sub>1</sub>, R<sub>0</sub> and P<sub>1</sub>, P<sub>0</sub>, P<sub>-1</sub> using the nomenclature of R<sup>2</sup> hards & Wykoff, 1971 (Figure 12). The conformation of d(Tp)4 has not been altered though one nucleotide has been omitted for clarity.

In the rEDN-d(Tp)3 model, the B<sub>2</sub>, B<sub>1</sub>, R<sub>2</sub>, R<sub>1</sub>, P<sub>1</sub> and P<sub>-1</sub> moieties can make specific interactions with rEDN. The bases both adopt *anti* conformation. The side cnains of Asn70 and Asp112 are within the B<sub>2</sub> subsite and the N and Oγ1 atoms of the invariant Thr42 of rEDN complement the hydrogen bonding potential of the O2 and N3 atoms of B<sub>1</sub>. The hydrogen bonding potential of the O4 atom of B<sub>1</sub> is unsatisfied in Figure 10. The R<sub>2</sub> and R<sub>1</sub> moieties are involved in van der Waals contacts with the Trp7 ring and Leu130, respectively. The P<sub>1</sub> and P<sub>-1</sub> phosphates in the rEDN-d(Tp)3 model have positions similar to the SO<sub>4</sub>(A) and SO<sub>4</sub>(B) anion positions observed in the rEDN. The P<sub>1</sub> phosphate of the model is located in the rEDN active site and can hydrogen bond with Gln14Nε2, His15Nε2, His129Nδ1 and Leu130N. The P<sub>-1</sub> phosphorus atom of the model is more than 3.0Å from the sulfur atom position of SO<sub>4</sub>(B). In the RNase A-d(Tp)<sub>4</sub> structure, the P<sub>-1</sub> phosphate makes van der Waals contacts with Pro42.

### DISCUSSION

The polypeptide fold of rEDN is related to those observed in the homologous RNase A (Zegers et al., 1994), Onconase (Mosimann et al., 1994) and angiogenin (Acharya et al., 1994) X-ray crystallographic structures. These enzymes share less than 35% amino-acid sequence identity and they range in size from 104 residues (Onconase) to 134 residues (EDN). Given the relatively low sequence identity and the size variation it is not surprising that the molecular surface of EDN is different from that of other pyrimidine specific RNases of known structure. The loops 12, 13 and 16 of EDN have significantly different conformations from those observed in structures of other vertebrate RNases. This is significant since enzymes belonging to the vertebrate RNase superfamily have a variety of biological functions and properties in addition to that of digesting RNA. The non-ribonucleolytic properties of several of the vertebrate enzymes are known to be associated with their specific binding to targets on the cell surface (Wu et al., 1993; Hallahan et al.,

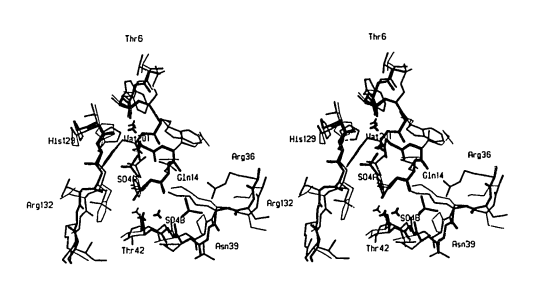


Figure 5.10 A stereo diagram of the least-squares superposed active site residues and bound anions of rEDN and RNase A are shown from the same vantage point as that in Figure 5.7. The rmsd for the 125 main-chain and Cβ atoms is 0.60Å. The thick lines represent rEDN and its anions and the thin lines represent RNase A and its anion. The conformations of the catalytic residue and the positions of the active site anions and are closely similar in the two structures. The conformation of His129 of rEDN is equivalent to one of the two alternate conformations (inactive) of the His119 (RNase A). The SO<sub>4</sub>(B) anion of rEDN makes a number of too-close contacts with Pro42 of RNase A. This proline residue probably prevents RNase A from binding an anion in a manner analogous to the binding of SO<sub>4</sub>(B) to rEDN.

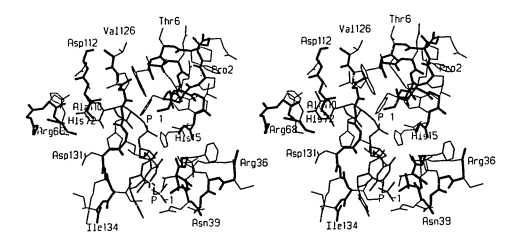


Figure 5.11 A stereo diagram of the rEDN-d(Tp)3 model based upon the RNase A-(Tp)4 structure (Birdsall et al., 1992). The rEDN backbone is shown as thick lines and the thin lines represent the side chain atoms. The d(Tp)3 inhibitor is shown in lines of intermediate thickness and the P<sub>1</sub> and P<sub>-1</sub> phosphates are labeled. The SO<sub>4</sub>(B) anion of rEDN is more than 3Å from the nearest anion, the P<sub>-1</sub> phosphate. The Asn70 and Asp112 residues of rEDN are within the B<sub>2</sub> subsite and Thr42 and Gln40 are in the B<sub>1</sub> subsite.

Figure 5.12 A schematic representation of a short RNA substrate. The base, ribose and phosphate moieties are referred to using the nomenclature of Richards and Wykoff, 1971. The P<sub>1</sub> phosphate binds to the active site rEDN and the scissile phosphodiester bond is the P<sub>1</sub>-O(5'). Note that the P<sub>1</sub>, R<sub>1</sub> and B<sub>1</sub> moieties together are equivalent to a 3'-monophosphate nucleotide. The base, ribose and phosphate numbering increases on the 5' side of the scissile phosphate (P<sub>1</sub>) of the RNA substrate and decreases on the 3' side.

1991; Titani et al., 1987). These loops may play a role in the function of EDN. Interestingly, EDN and Onconase share little structural homology other than their general fold yet EDN, ECP and Onconase represent the only neurotoxins within this superfamily. Additionally, the large insertion unique to the non-secretory RNases (eg. EDN: Asp115 to Tyr123) may contribute to their unique properties. At present these residues have no known biological role.

rEDN has two discrete SO<sub>4</sub> anion binding sites. The active site SO<sub>4</sub>(A) and the SO<sub>4</sub>(B) anions observed in the rEDN structure are wholly bound by a single rEDN molecule and do not make contacts with symmetry related molecules. In the structures of RNase A-oligodeoxynucleotide complexes (Birdsall et al., 1993; Fontecelli-Camps et al., 1994), phosphate, ribose and base subsites have been identified for extended substrates (Figure 11). Phosphate subsites corresponding to P<sub>1</sub> and P<sub>2</sub> have been observed while the P<sub>0</sub> and P<sub>3</sub> phosphate moieties do not bind to specific subsites. The SO<sub>4</sub>(A) (P<sub>1</sub>) and SO<sub>4</sub>(B) anions are 9.4Å apart in rEDN structure and this distance is considerably larger than distance between consecutive phosphates of RNA or DNA when in an extended conformation. This would indicate the SO<sub>4</sub>(B) binding site of rEDN is not P<sub>0</sub> or P<sub>2</sub>. In the RNase A-d(Tp)4 structure, the P-1 phosphate makes van der Waals contacts with the side chain of Pro42. The residue that is structurally equivalent to Pro42 of RNase A is Asn39 and it donates hydrogen bonds to SO<sub>4</sub>(B) through its main-chain and side-chain atoms (Figure 5.10). In Figure 11, the SO<sub>4</sub>(B) anion of rEDN is nearest to the P-1 phosphate of the RNase A-d(Tp)4 model (>3.0 Å between sulfur to phosphate). As a result, the SO<sub>4</sub>(B) binding site likely correspond to the P<sub>-1</sub> subsite of rEDN. Within the superfamily, the residue equivalent to Pro42 of RNase A is not highly conserved. It is present in pancreatic enzymes and angiogenins but not in amphibian or non-secretory RNases. Thus, the P-1 phosphate binding in RNase A may not be representative of the P-1 phosphate binding in all related enzymes. Instead, relatives that do not have an imino acid at this position may bind the P-1 phosphate similar to the observed SO<sub>4</sub>(B) binding mode of rEDN.

The enzymatic activity and specificity of rEDN towards diribonucleotide substrates are comparable to those of RNase A. rEDN has a slight preference for cytidine in the B1 subsite and prefers adenosine in the B2 subsite. In the superposition of rEDN and d(Tp)4 (Figure 12) the N2 and O3 atoms of the B<sub>1</sub> base can hydrogen bonded to the invariant Thr42. In RNase A, O4 of B<sub>1</sub> hydrogen bonds to a solvent molecule that hydrogen bonds to Ser123 (Zegers et al., 1994). The equivalent residue of rEDN is Ile133 and it cannot make analogous interactions. The nearest potential protein hydrogen bond donor or acceptor of rEDN is Gln40. This is interesting as Gln40 can adopt conformations that allow it to hydrogen bond to SO<sub>4</sub>(B) yet does not. In the X-ray structure of rEDN, the Gln40 side chain is solvent exposed and disordered beyond CY. Perhaps Gln40 is responsible for binding the O4 of uracil or N4 of cytosine. Like Ser123 of RNase A, Gln40 can either accept or donate hydrogen bonds. This suggests both cytosine and uracil can be accommodated within the B<sub>1</sub> subsite and explains the observed specificity of rEDN (Table 1.4). In RNase A complex structures, the B2 base hydrogen bonds to Asn71 and Glu111. The equivalent residues of rEDN are Asn70 and Asp112. In Figure 11, Asn70 of rEDN can hydrogen bond to O4 atom while Asp112 is directed at the C5 methyl group. The B<sub>2</sub> base of the RNase A-d(Tp)<sub>4</sub> structure is rotated by 180° when compared to the high resolution RNase A-d(CpA) structure. The pseudorotation angle of the sugars R1 and R<sub>2</sub> and the glycosyl torsion angle of the bases B<sub>1</sub> and B<sub>2</sub> are comparable in these structures. The conformational difference arises from large differences in the  $\beta$  (P-O5'-C5'-C4') and y torsion angles (O5'-C5'-C4'-C3') of the nucleic acid backbone of these structures. If the B2 base binds as seen in the high resolution d(CpA) structure, Asp112 will be in close proximity to the N3 atom. Arg68 is an additional rEDN residues that may be near enough to contribute to the binding the B2 base.

In a recent comparative molecular modeling assessment, the loops 12, 13 and the omega loop 14, 16 and 17 (the large insertion of rEDN) were not successfully modeled by

the participating groups (Mosimann et al., 1995). These large loops are the sites of insertions and deletions in comparisons of the primary sequences of rEDN, RNase A and Onconase and correspond to the regions of the greatest divergence in structural superpositions of these enzymes (Figure 5.9). In rEDN, it appears the loops of RNase A and Onconase represent poor models of the conformations of loops in rEDN as a result of the low sequence identity and the presence of insertions and/or deletions. The conformation of the large insertion of rEDN is exceedingly difficult as it packs against additional inserted residues at the N-terminus (Figure 5.6). A successful prediction of the conformation of this loop would require the correct simultaneous prediction of the conformation of the N-terminus as well. In many ways this is a small version of the folding problem. These difficulties suggest the prediction of large, divergent loop structures is not yet possible using current methods.

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### CHAPTER 6

#### DISCUSSION

ONC and EDN are cytotoxic members of the pyrimidine specific RNase superfamily of vertebrates. They have therapeutic potential and play a role in various human pathologies, respectively. To further our understanding of the biological properties of these enzymes, the molecular structures of ONC, rONC and EDN were determined and refined at high resolution using X-ray crystallographic techniques. They represent the first reported structures of amphibian and non-secretory pyrimidine-specific RNases. This is significant as the homologous RNase A and angiogenin enzymes are not cytotoxins.

The overall folds of ONC and rEDN are related to those observed in crystal structures of RNase A (Wlodawer et al., 1983; Zegers et al., 1994) and angiogenin (Acharya et al., 1994). These enzymes are composed of an N-terminal  $\alpha$ -helix and two similarly folded lobes. Lobe 1 contains strands  $\beta$ 1,  $\beta$ 3 and  $\beta$ 4 and helix  $\alpha$ 2 while lobe 2 contains strands  $\beta$ 2,  $\beta$ 5 and  $\beta$ 6 and helix  $\alpha$ 3. Each lobe forms a three-stranded antiparallel  $\beta$ -sheet and the  $\alpha$ -helix packs against the  $\beta$ -sheet. In addition, the  $\alpha$ -helix of each lobe contributes a conserved disulfide bridge to the central strand of their respective  $\beta$ -sheets. The similarity in the structure of these two  $k_{\beta}$  er suggests a gene duplication event gave rise to the original RNase.

While these enzymes have the same overall fold, their detailed molecular structures are significantly different. In particular, the N-terminus, the loop structures following the three  $\alpha$ -helices and the omega loops following the central strands of the two  $\beta$ -sheets have very different conformations in each of these enzymes. In addition, there are a number of non-conservative amino acid replacements among structurally equivalent residues. These differences are expected to be responsible for the biological properties of these enzymes.

At present, our lack of knowledge regarding the identity and nature of the receptors of ONC and EDN prevent us from identifying the residues involved in the cell binding activity of these enzymes. Future studies using site directed mutants of rONC and rEDN will be aimed at identifying residues that facilitate the cytotoxic mechanism of these enzymes.

# B<sub>1</sub> Subsite of Pyrimidine Specific RNases

The active site of the pyrimidine specific RNases is located at the junction of the Nterminal α-helix, lobe 1 and lobe 2. Together, they form a cleft that runs parallel to strand β6. At the base of the cleft are two invariant residues; a histidine and a threonine. In the B1 subsite the N and  $O^{\gamma l}$  atoms of the invariant Thr complement the hydrogen bonding potential of the O(2) and N(3) atoms of the pyrimidine base, respectively. As a consequence, the differences in the pyrimidine-base specificity of these two enzymes arise from differences in the binding of the N(4) (cytosine)/O(4) (uracil) atom. In the RNase Ainhibitor structures, direct hydrogen bonds between N(4) and Ser123OYl (Richards & Wykoff, 1971) and water-mediated interactions between O(4) and Ser123Oγ1 (Wlodawer et al., 1983; Borah et al., 1985) have been reported. Since Ser123091 and water molecules can either accept or donate hydrogen bonds, they account for the pyrimidine base preference of RNase A. The residues of ONC, EDN and angiogenin that are equivalent Ser123 of RNase A are Val101, Ile133 and Ser124, respectively. The ONC-d(UpG) model (Chapter 4) and the rEDN-d(Tp)3 model (Chapter 5) suggest Lys33 of ONC and Gln40 of EDN replace the Ser123 interaction with the B<sub>1</sub> base observed in RNase A complex structures. Lys33 of ONC and Gln40 of EDN are structurally equivalent residues that account for the observed specificity of their respective enzymes. In RNase A, the counterpart to these residues is Val43. As a result, two non-conservative substitutions are required to change the specificity of ONC to that of RNase A or vice versa. It is interesting that angiogenin has an Arg that is equivalent to Lys33 of ONC and retains a Ser that is equivalent to Ser123 of RNase A. Both angiogenin residues can participate in hydrogen bonding interactions with

the O(4) (uracil)/N(4) (cytosine) at ms of the pyrimidine base and angiogenin has a specificity that is intermediate to that observed for ONC and RNase A (Talle 1.4).

# B<sub>2</sub> Subsite of Pyrimidine Specific RNases

The B2 subsite of all pyrimidine specific RNases includes a Glu (RNase A; Glul 11) or an Asp (EDN: Asp112) that are structurally equivalent. In RNase A, EDN and angiogenen there are several residues from the loop 14 that can participate in substrate binding. In ONC a large portion of this loop has been deleted (Chapter 2, 3). As a consequence, the loop 14 and the B2 subsite of ONC have very different conformation when compared to other pyrimidine specific RNases. In angiogenin, residues of the loop 14 contribute to its angiogenic potential. The loop 14 of angiogenin has been replaced with the equivalent loop of RNase A and resulting enzyme shows increased enzymatic activity and a greater RNase A-like character. Moreover, the chimeric enzyme loses its angiogenic activity (Hallahan et al., 1991). As a result, it has been proposed that the loop 14 is a module that controls the enzymatic activity and biological function of pyrimidine specific RNases. If this is correct, the deletion of this loop may confer the guanine specificity of ONC to other members of the superfamily.

### rONC

In the X-ray crystallographic structure of ONC (Chapter 3), the N-terminal pyroglutamyl residue folds back against the N-terminal α-helix and forms a hydrogen bond with Lys9 at the enzymes active site. An equivalent pair of residues is found in each of the amphibian RNases and in the iguana RNase (Zhao et al., 1994). In all mammalian members of this superfamily the residue equivalent to Lys9 is a Gln. This suggests the Pyr1 interaction with Lys9 is an important feature of the non-mammalian enzymes. The discovery that rONC with an N-terminal sequence fMet-1Glu1Asp2··· lacks significant RNase activity and cytotoxicity confirms Pyr1 has a role in the RNase activity of the

enzyme (Wu, Y. & Youle, R.J., unpublished results). At present several approaches to restoring the activity of rONC are being explored. These include; the generation of Pyr1 by chemical means after CNBr or enzymatic removal of  $^f$ Met.1 and the construction of point mutants that would replace the function of Pyr1 in the R2 subsite (Chapter 4). In the structure of ONC, Pyr1 packs against Thr5 of the N-terminal  $\alpha$ -helix. In all mammalian pyrimidine specific RNases the residues equivalent to Thr5 of ONC are lysine (RNase A), tryptophan (EDN) or histid (angiogenin). The relatively large side chains of these amino acids sterically clash with Pyr1 when modeled in place of Tm3. As a result we have suggested a mutation of Thr5 to Trp to replace Pyr1 in the R2 subsite and an additional mutation, Asp2 to Trp. By changing Asp2, the negative charge of the  $\beta$  carboxyl is removed from the R2 subsite and potentially unfavorable electrostatic interactions with adjacent phosphate subsites are avoided (Chapter 4).

### P.<sub>1</sub> Subsite of rEDN

The structure also contains two discreet solfate anion binding sites. The active site anion is non-covalently bound in a similar position and conformation in each of these structures. The SO<sub>4</sub>(B) anion occupies a site that may correspond to the P<sub>-1</sub> subsite. SO<sub>4</sub>(B) hydrogen bonds to the main-chain and side-chain atoms of Asn39. In RNase A, the residue equivalent to Asn39 is Pro42. Clearly, Pro42 can make neither of the Asn39 interactions observed in the rEDN structure. In a tetra deoxythymidylic acid-RNase A complex (Birdsall et al., 1992) the P<sub>-1</sub> phosphate makes van der Waal's contacts with Pro42. In the absence of Pro42, RNase A may bind the P<sub>-1</sub> phosphate as observed at rEDN. Since the position equivalent to Pro42 (RNase A) is not highly conserved within the superfamily, the SO<sub>4</sub>(B) binding in rEDN may be more representative of how other RNase bind extended substrates.

# Comparative Molecular Modeling

The comparative molecular modeling study of ONC (Chapter 2) demonstrates ONC can adopt a fold related to that observed in RNase. The model correctly predicted the conformation of residue Ser54-Leu58 justifying a revision in the existing automated sequence alignment. The original sequence alignment predicted the deletion of strand  $\beta 2$ and aligned these residues with a large loop structure (Ardelt et al., 1991). The comparative molecular modeling did not correctly predict the structures of loops with large insertions and deletions and failed to predict the conformation of the N-terminus and helix α2. As evidence of the differences in the model and observed structures of ONC, the model was unable to provide a molecular replacement solution. The correct rotation function solution does not appear among the top 100 rotation function peaks when using the model. Previous and subsequent comparative molecular model assessments have suggested these loop structures are exceedingly difficult to predict when they are the sites of insertions and deletions and have little or no amino acid sequence identity (Greer, 1985; Mosimann et al., 1995). The helix a2 is flanked loops with large deletions and is more than 2Å from its predicted position. The modest amino acid sequence identity shared by these enzymes is restricted to disulfide bridges that help define the tertiary structure of these enzymes, residues known to be important for catalysis and several residues of the hydrophobic core (Table 1.2). As a result, the loop structures of RNase A are a poor model for the observed conformation of the ONC and rEDN loops.

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