# TESS: A geometric hashing algorithm for deriving 3D coordinate templates for searching structural databases. Application to enzyme active sites

# ANDREW C. WALLACE, 1 NEERA BORKAKOTI, 2 AND JANET M. THORNTON 1.3

<sup>1</sup>Biomolecular Structure and Modelling Unit, Department of Biochemistry and Molecular Biology, University College, Gower Street, London WC1E 6BT, England

<sup>3</sup>Department of Crystallography, Birkbeck College, Malet Street, London WC1E 7HX, England

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#### Abstract

It is well established that sequence templates such as those in the PROSITE and PRINTS databases are powerful tools for predicting the biological function and tertiary structure for newly derived protein sequences. The number of X-ray and NMR protein structures is increasing rapidly and it is apparent that a 3D equivalent of the sequence templates is needed. Here, we describe an algorithm called TESS that automatically derives 3D templates from structures deposited in the Brookhaven Protein Data Bank. While a new sequence can be searched for sequence patterns, a new structure can be scanned against these 3D templates to identify functional sites. As examples, 3D templates are derived for enzymes with an O-His-O "catalytic triad" and for the ribonucleases and lysozymes. When these 3D templates are applied to a large data set of nonidentical proteins, several interesting hits are located. This suggests that the development of a 3D template database may help to identify the function of new protein structures, if unknown, as well as to design proteins with specific functions.

Keywords: catalytic triad; database; 3D structure; protein function; template

The detection of recurring structural motifs in proteins is already well documented; indeed, they exist in all levels of protein structure, from primary to tertiary. At the primary level, there are now comprehensive protein sequence databases such as SWISS-PROT (Bairoch & Boeckmann, 1994) and OWL (Bleasby et al., 1994), that have been analyzed, using both automatic and manual patternmatching and sequence-alignment techniques, to produce databases of recurring sequence motifs or templates such as PROSITE (Bairoch & Bucher, 1994) and PRINTS (Attwood et al., 1994). At the tertiary level, protein structure analysis, using both automatic (Orengo et al., 1993) and manual techniques, has enabled the creation of classifications such as CATH, SCOP, (Murzin et al., 1995) and DALI (Holm & Sander, 1993), which cluster proteins with similar folds. These databases are useful for the identification of biological function and fold recognition (Lemer et al., 1995). Neighborhood relationships can also be identified, using structural similarity methods, as exemplified by MMDB (Ohkawa et al., 1997).

Reprint requests to: Janet M. Thornton, Biomolecular Structure and Modelling Unit, Department of Biochemistry and Molecular Biology, University College, Gower Street, London WC1E 6BT, England; e-mail: thornton@biochem.ucl.ac.uk.

Many investigations have taken place at the substructural level of proteins, for example, analysis of 3D topologies of metal-binding sites in proteins and small molecules (see reviews by Glusker, 1991; Jernigan et al., 1994). PROMOTIF (Hutchinson & Thornton, 1996) provides automatic assignment and analysis of structural motifs in proteins, such as secondary structure,  $\beta$ -and  $\gamma$ -turns,  $\beta$ -hairpins, and disulfide bridges. In addition, there are already various algorithms to detect similar 3D arrangements of secondary structure in proteins. Some are comparison techniques that require the linear order of the amino acid sequences to be conserved (Matthews & Rossmann, 1985); others allow some degree of insertion/deletion in the protein sequence (Alexandrov et al., 1992), whereas others match by secondary structure elements (Mitchell et al., 1990).

However, as yet, a database of recurring 3D templates or motifs in proteins does not exist; these can be thought of as the 3D equivalent of the 1D templates found in the PRINTS and PRO-SITE databases. The number of protein 3D structures being solved by X-ray crystallography and NMR spectroscopy techniques is increasing rapidly; there are expected to be more than 20,000 by the turn of the century. This suggests that the need for a 3D equivalent of PROSITE is also growing; this would enable us to suggest functions of proteins whose roles are unknown as well as allow us to locate automatically functional regions and catalytic residues

<sup>&</sup>lt;sup>2</sup>Roche Discovery Wellwyn, 40 Broadwater Road, Welwyn Garden City, Hertfordshire AL7 3AY, England

within the protein structure. Such databases could address many different substructural aspects of proteins, such as enzyme active sites, ligand binding sites, loop conformation, and metal binding sites.

Previously, Artymiuk et al. (1994) have used a graph-theoretic approach for searching 3D patterns of side chains in protein structures. For example, they constructed a search template from the side-chain atoms of the Ser 195-His 57-Asp 102 catalytic triad of chymotrypsin and, depending on the allowed interatomic distance tolerances, different numbers of catalytic triads were identified from their data set. The template also revealed the existence of a second Ser-His-Asp triad in trypsinogen and chymotrypsinogen. A different structural comparison of the serine proteinases, using a less specific technique, has been performed by Fischer et al. (1994). Their method, derived from geometric hashing methods first described by Lamdan et al. (1988) for use in computer vision research, treats all  $C^{\alpha}$  atoms in a protein as points in space and compares proteins purely on the geometrical relationships between these points. It can detect recurring substructural 3D motifs and was able to identify the structural similarities of the active sites of the trypsin-like and subtilisin-like serine proteases based solely on the similarities of the  $C^{\alpha}$  geometries of their constituent residues.

We have previously shown that a Ser-His-Asp 3D enzyme active site template can be defined that will identify all the serine proteinases and lipase active sites in a set of PDB structures with the exclusion of all other noncatalytic Ser, His, and Asp interactions (Wallace et al., 1996). However, this method is not applicable to the case where the search atoms lie far apart in the protein structure because the search procedure becomes combinatorially intense. Here, an algorithm called TESS (TEmplate Search and Superposition) is described that is also based on the geometric hashing paradigm. TESS searches through a data set of Protein Data Bank structures (PDB; Bernstein et al., 1977) for any user-defined combination of atoms in space, from single atoms to multiple residues. The method is faster yet just as accurate as the method implemented by Wallace et al. (1996).

The new method is described together with its application to deriving enzyme active site consensus templates. First, we investigate those enzymes with a His-based "catalytic triad," namely the serine proteinases (Blow et al., 1969; Wright et al., 1969), cysteine proteinases (Drenth et al., 1968), lipases (Brady et al., 1990), and  $\alpha/\beta$ -hydrolase (Ollis et al., 1992) enzymes. In addition, the active sites of eukaryotic and prokaryotic lysozymes (Johnson & Phillips, 1965; Matthews & Remington, 1974) as well as ribonucleases A and  $T_1$  (e.g., Beintema et al., 1990; Arni et al., 1992) are explored.

In addition, using the TESS program, we have created a database of enzyme active site templates called PROCAT that is available using the World Wide Web at: http://www.biochem.ucl.ac.uk/ bsm/PROCAT/PROCAT.html.

### Results

# The TESS algorithm

The TESS algorithm must address the following problem. Given the coordinates of the 3D query template and a protein structure, is there a match between them and what is the transformation, if any, that will best superimpose the template onto the relevant part of the protein molecule? This means that the atoms, residue types, and 3D coordinates of the template need to overlap with those of the stored PDB structure within a user-defined distance cut-off. The

method we have devised to solve this problem is summarized in Figure 1 and consists of two stages: pre-processing and search with query template.

# Pre-processing

In the pre-processing stage, a set of hash tables is compiled, containing geometrical information about the atoms of all the structures in the PDB. This avoids having to recalculate the data each time TESS is run and speeds up the comparison process considerably.

The atoms comprising a given template need to be defined with respect to a reference frame. For example, in the Ser  $O^{\gamma}$ -His sidechain-Asp  $O^{\delta}$  consensus template of the serine proteinases and lipases, the His side chain defines the reference frame (Wallace et al., 1996). In fact, only three atoms of the side chain are used  $(C^{\delta 2}, C^{\gamma}, \text{ and } N^{\delta 1})$ . Table 1 shows the three reference atoms used for each of the 20 standard amino acid side chains. The method used to create a hash table is summarized in Figure 2, using a His reference frame residue as an example. For each of the His residues in the protein structure, all atoms within a user-defined distance are identified; we chose 18 Å. The transformation matrix is calculated that places the His residue reference frame  $C^{\gamma}$  atom at the origin, with the  $C^{\delta 2}$  and  $N^{\delta 1}$  either side of the positive x direction. The same transformation matrix is applied to the atoms surrounding each His residue. A grid of 1 Å separation is placed around the His and the surrounding atoms and the position within the grid that each atom occupies is noted. To enable rapid searching through the hash table with the query template, the atoms in each grid are reordered according to their grid position number and are assigned an atomic label according to the residue and atom type. This information, along with the PDB file atom numbers, is stored in the hash table.

# PREPROCESSING QUERY TEMPLATE Dataset of PDB structures Liser defined

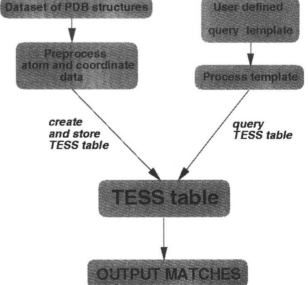


Fig. 1. Summary of the process involved when TESS searches through a data set of PDB structures for a user-defined 3D template.

Table 1. Side-chain atoms	used to define the reference frames for each	h
standard amino acida	•	

Residue	1	2	3	Residue	1	2	3	Residue	1	2	3
Ala (A)	N	Cα	$C^{\beta}$	Gly (G)	N	$C^{\alpha}$	С	Pro (P)	N	$C^{\alpha}$	$C^{\beta}$
Arg (R)	$N^{\eta_1}$	N€	$N^{\eta_2}$	His (H)	$C^{\delta_2}$	$C^{\gamma}$	$N^{\delta_1}$	Ser (S)	$C^{\alpha}$	$C^{\beta}$	$O^{\gamma}$
Asn (N)	$O^{\delta_1}$	$C^{\gamma}$	$N^{\delta_2}$	Ile (I)	$\mathbf{C}^{\gamma_1}$	$C^{\beta}$	$C^{\gamma_2}$	Thr (T)	$O^{\gamma_1}$	$C^{\beta}$	$C^{\gamma_2}$
Asp (D)	$\mathbf{O}^{oldsymbol{\delta}_1}$	$C^{\gamma}$	$\mathbf{O}^{\delta_2}$	Leu (L)	$C^{\delta_1}$	$C^{\gamma}$	$\mathbf{C}^{\delta_2}$	Trp (W)	$C^{\delta_1}$	$C^{\gamma}$	$C^{\delta_2}$
Cys (C)	$C^{\alpha}$	$C^{\beta}$	$S^{\gamma}$	Lys (K)	C <sup>\epsilon</sup>	$C^{\delta}$	Nζ	Tyr (Y)	$C^{\delta_1}$	$O^{\eta}$	$C^{\delta_2}$
Gln (Q)	$O^{\epsilon_1}$	$C^{\delta}$	$N^{\epsilon_2}$	Met (M)	C۴	$C^{\gamma}$	$S^{\delta}$	Val (V)	$C^{\gamma_1}$	$C^{\beta}$	$C^{\gamma_2}$
Glu (E)	$O^{\epsilon_2}$	$C^{\delta}$	$O^{\epsilon_1}$	Phe (F)	$C^{\epsilon_1}$	$C^{\gamma}$	$C^{\epsilon_2}$				

<sup>&</sup>lt;sup>a</sup>Atoms in column 2 are transformed to the origin with atoms in columns 1 and 3 either side of the positive x direction.

# Searching with a 3D query template

A query template is defined in terms of a side-chain reference frame and the atoms or other side chains in its vicinity. The coordinate template file is in standard PDB format and each atom can be specified in terms of an atom type (e.g., C, N, S, O) or a particular side-chain atom (e.g., Asp  $O^{\delta}$ ).

An example of a typical query template, taken from the active site of  $\alpha$ -lytic proteinase, 1lpr, is shown in Table 2. The reference frame residue, in this case a His, is defined by putting a "-1" in the atom number column corresponding to  $C^{\delta 2}$ ,  $C^{\gamma}$ , and  $N^{\delta 1}$ . This enables comparison of the relative positions of the atoms surrounding the query template with those of the protein structures stored in the hash tables. For the nonreference frame atoms in the query template, it is possible to define any combination of both atom and residue types that are to be searched for at that point. This is in contrast to the method of Fischer et al. (1994), which is limited to only  $C^{\alpha}$  atom searches.

To define which residues are to be located at a given template atom position requires the one-letter amino acid code corresponding to that amino acid to be placed after the coordinates of the query template. The atom type to be searched for is defined by placing one of the numbers in listed in Table 3 in the atom number column of

Read in protein coordinates

Extract all His residues

Mark all atoms less than 18Å from each His residue

Transform each His and surrounding atoms

Place grid around His and surrounding atoms

Calculate position of each atom relative to grid

Store atom and grid positions in TESS table

Fig. 2. Flow diagram showing the steps required in creating a hash table in the pre-processing stage, using the His reference frame residue as an example.

the query template. For example, there will be a search for any noncarbon side-chain atom of residue type Ser or Asp at the coordinate position in the first row of the query template in Table 2.

The relative position of the atoms around the query His residue are calculated in the same way as the pre-processing stage; a grid is placed around the 3D template atoms, with the His reference frame at the origin, and the grid positions of each of the atoms are calculated. First the grid positions and then the atomic labels (atom and residue type) of the query template are compared to those of the proteins stored in the hash table; a hit occurs when all the atoms of the query template have a corresponding match with atoms in the protein structure. When comparing the grid positions, the situation is slightly complicated because it is not possible to tell where the atoms are lying in the grid boxes; they may be very near to one of the sides. Therefore, we have to search all neighboring grid boxes stored in the hash table to the central query atom grid box. Of course, the larger the distance cut-off, the more layers of neighboring grid boxes that will have to be searched. For each match, the matching PDB file is opened and the relevant atomic coordinates are transformed to the same reference frame and the RMS distance of the match is calculated.

Once the search is completed, a consensus template may be calculated by taking the mean coordinates of all the hits located.

# Performance of the TESS algorithm

TESS takes around 0.25 CPU seconds to locate a template match for a typical protein structure on a SGI R4400 Challenge. The run

**Table 2.** An example of a typical query template taken from the active site of  $\alpha$ -lytic proteinase, Ilpr

Search option	Residue	Residue number	Atom	х	у	z	Residue search
1	Ser	195	Ογ	16.3	30.6	14.7	D
1	Asp	102	$\mathrm{O}^{\delta_2}$	18.2	31.5	20.8	Е
0	His	57	$C^{\beta}$	14.5	28.8	20.9	
-1	His	57	$C^{\gamma}$	15.0	29.3	19.5	
-1	His	57	$N^{\delta_1}$	16.2	30.0	19.3	
-1	His	57	$C^{\delta_2}$	14.3	29.1	18.3	
0	His	57	$C^{\epsilon_1}$	16.2	30.3	18.0	
0	His	57	$N^{\varepsilon_2}$	15.1	29.8	17.4	

**Table 3.** Search parameter numbers placed in the atom number column of the query PDB format file <sup>a</sup>

Search option	Atom numbe
Template atom	-1
Search by atom type, i.e., $O^{\delta_1}$ , $O^{\delta_2}$ , $O^{\gamma}$	0
Search by non-carbon sidechain atom	1
Search by non-carbon atom	2
Search by specified atom, i.e., C, O, N	3
Search by non-carbon mainchain atom, i.e., O, N	4
Search by any mainchain atom	5
Search by any sidechain atom	6
Search by any atom type	7

<sup>a</sup>One of these numbers is placed against each of the atoms in the query template. This defines which atom types are to be searched for at the corresponding atom position. To search for different residue types at a given atom point requires the one-letter code of that amino acid to be placed after the coordinates in the query template file.

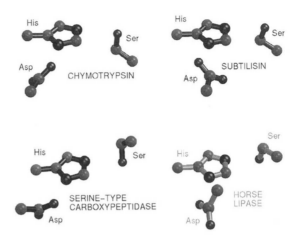
time is proportional to the order, O(nh), where n is the number of atoms present in the template and h the number of hits. Each hash table uses around 70 megabytes of memory for the 3,019 structures found in the January 1995 PDB. The amount of memory used is proportional to  $l^3$ , where l is the length of the grid side, in our case, 36 Å.

# The His-based catalytic triad

# Template definition

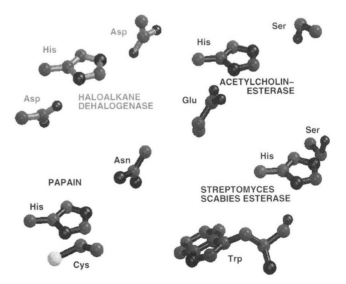
The His-based catalytic triad was first identified in the serine proteinases (Blow et al., 1969; Wright et al., 1969) and comprises the residues Ser-His-Asp. Subsequently, other enzymes in the PDB, the  $\alpha/\beta$ -hydrolase fold enzymes (Ollis et al., 1992), the esterase from Streptomyces scabies (Wei et al., 1995), and the cysteine proteinases (e.g., papain, Drenth et al., 1968) were shown to have a His-based catalytic triad. In fact, these triads can be generalized as Nu:-His-ELEC, where Nu: is the nucleophilic group and ELEC is the electrostatic group (equivalent to Asp in the Ser-His-Asp triad). Figures 3 and 4 show the heterogeneity in the orientation of the residues comprising these catalytic triads. We have divided these enzymes into five classes according to the residues that comprise their catalytic triads, which are Ser-His-Asp, Ser-His-Glu, Asp-His-Asp, Ser-His-Trp, and Cys-His-Asn. These are summarized in Table 4, which lists all PDB entries in the June 1996 release in each class. The  $\alpha/\beta$ -hydrolase fold (Ollis et al., 1992) occurs in three of the five classes and, despite a low sequence identity, it suggests that the enzymes in this fold group have evolved from a common ancestor and have preserved the positions of the key catalytic residues.

Figure 5 is a 3D representation of the catalytic triads from classes 1, 2, 3, and 4. The His side chains have been superimposed, enabling us to compare the relative position of the nucleophilic and electrostatic side chains. Chymotrypsin 1*cho* (Fujinaga & James, 1987) represents class 1, although the dispositions of the side chains of the other class 1 members—namely subtilisin, serine-type carboxypeptidase, and lipase—are quite different to those of chymotrypsin and have been left out for clarity (Wallace et al., 1995). The catalytic triad of the cysteine proteinases (class 5) is



**Fig. 3.** Relative conformation of the catalytic triads from chymotrypsin 1*cho* (Fujinaga & James, 1987), subtilisin 2*sic* (Takeuchi et al., 1991), serine-type carboxypeptidase 3*sc*2 (Liao et al., 1992), and lipase 1*hpl* (Bourne et al., 1993). Diagram produced using MOLSCRIPT (Kraulis, 1991) and Raster3D (Bacon & Anderson, 1988)

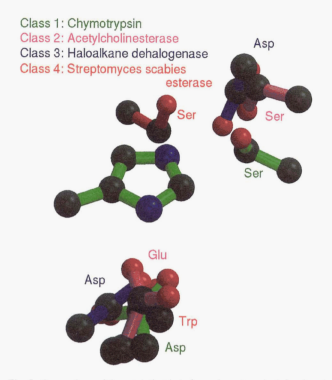
different from those of classes 1, 2, 3, and 4 because the Cys 25 nucleophile interacts with the His  $N^{\delta 1}$  rather than the  $N^{\epsilon 2}$  (Fig. 4). The diagram shows that, although both side chains originate from different orientations in classes 1, 2, and 3, there is clustering of the functional atoms; the nucleophilic oxygens are all in the proximity of the acid/base catalyst His  $N^{\epsilon 2}$  and the electrostatic residues are in a hydrogen bonding position with the His  $N^{\delta 2}$ . In contrast, the serine in the esterase (class 4) lies in a distinctly different position. We have shown previously that a Ser  $O^{\gamma}$ -His side-chain-Asp  $O^{\delta}$  template can identify all serine proteinases and lipase catalytic triads of class 1 with the exclusion of all other interactions between Ser, His, and Asp side chains. Figure 5 suggests that this template might be extended to include classes 2 and 3, giving a class 1-2-3 consensus template. Sussman et al. (1991)



**Fig. 4.** Relative conformation of the catalytic triads from haloalkane dehalogenase (Verschueren et al., 1993), acetylcholinesterase (Sussman et al., 1991) and *Candida rugosa* lipase (Grochulski et al., 1993), papain (Drenth et al., 1968), and *S. scabies* esterase (Wei et al., 1995).

**Table 4.** Five different classes of Nu:-His-ELEC catalytic triads found in the PDB (June 1996 release), where ELEC acts to perturb the  $pK_a$  of the acid/base His and Nu: is a nucleophilic group

					β-Sandwich t	Asp catalytic t rypsin-like fold					
lacb E	legi E	legj E 2cha	1chg	Ch Icho E 2gct A	nymotrypsin E.C 1gcd 2gmt	2.3.4.21.1 31 ch 1gct A 3gch	lgha E	Ighb E 4cha A	Igmc A 4cha B	Igmd A 4gch	1gmh 5cha A
cga A icha B	2cga B 5gch	6cha A	2gch 6cha B	6gch	7gch	8gch 4.21.4 55 chain	3gct A	Tella A	-ciia ib	4gen	Jena A
bit mct A	lbra Ippc E	lbrb E lppe E	lbrc E lpph E	lbtp lsgt	lbtw A lsmf E	lbtx A Itab E	lbty ltbs	lbtz A ltgb	lept A ltgc	lept B ltgn	lgbt ltgs Z
tgt	Itld	ling	1 tnh	ltni	1tnj	Itnk	l tnl	Itpa E	ltpo	Itpp 2tgd	1tps 2tgp Z
trm A !tgt	1trm B 2trm	1trn A 3ptb	1trn B 3ptn	1try 3tpi Z	ltyn 4ptp	2ptc E 4tpi Z	2ptn	2tbs	2tga	2 (gu	zigp Z
labi H	labj H	labj L	laht H	1bbr H	1bbr E	3.4.21.5 58 chai 1bbr K	1bbr N	ldwb H	ldwc H	Idwd H	1dwe I Ihdt H
letr H lhgt H	lets H lhlt H	lett H lhit K	lfpc H lhrt H	lfph H lhut H	lhag E lihs H	lhah H liht H	Ihai H Inrn H	Ihao H Inro H	lhap H Inrp H	Ihbt H Inrq H	Inrr H
lnrs H 2hnt F	1ppb H 2hpp H	1thr H 2hpq H	1ths H 2hpq P	1tmh H 2hpq L	1tmt H 2pf1	1tmu H 2pf2	2hat H 2spt	2hgt H 3hat H	2hgt L 4htc H	2hnt C	2hnt E
lcea A	lcea B	Iceb A	Iceb B	1pk4	lpkr	.4.21.7 9 chain: Ipmk A	lpmk B	2pk4			
lgba A	lgbb A	lgbc A	1gbd A	α-ly 1gbe A	lgbf A	.C.3.4.21.12 34 1gbh A	1gbi A	lgbj A	lgbk A	Igbl A	Igbm A
llpr A 2alp	lp01 A 2lpr A	1p02 A 2p07	1p03 A 3lpr A	1p04 A 4lpr A	lp05 A 5lpr A	lp06A 6lpr A	1p08 A 7lpr A	1p09 A 8lpr A	lp10 A 9lpr A	Ipl1 E	1p12 E
Iton	2kai A	2kai B	2pka A	Tis: 2pka B	sue kallikrein E 2pka X	.C.3.4.21.35 7 c 2pka Y	chains	-	-		
Ibma A	leas	leat	leau			E.C.3.4.21.36 26 Telc A	chains leld E	Iele E	lelf	lelg	lelt
lesa Best E	lesb 9est	lest	linc	ljim	lnes E	2est E	3est	4est E	5est E	6est	7est E
lhne E	Ippf E	Ippg E		Leul	ocyte elastase	E.C.3.4.21.37 3	chains				
	тррг Б	TPPS L		Comp	lement factor D	E.C.3.4.21.46	1 chains				
l dst	toro			Lysyl	endopeptidase	E.C.3.4.21.50 2	2 chains				
larb	lare	2E	Arm E		eptogrisin A E.	C.3.4.21.80 5 c	hains				
lsgc	2sga	3sga E	4sga E		eptogrisin B E.	C.3.4.21.81 5 c	hains				
lsgp E	1sgq E	lsgr E	3sgb E	Sub		subtilisin-like f C.3.4.21.62 32					
lese E Ised	Imee A Iscn B	Impt Isel A	1s01 1sel B	1s02 1sib E	lsbc lspb P	lsbh lspb S	1sbi 1st2	1sbn E 1st3	lsbt Isub	1sca 1suc	1scb 1sud
lsup	2sbt	2sec E	2sic E	2sni E Enc	2st1 lopeptidase K E	3sic E E.C.3.4.21.62 5	5sic E chains				
lpek E	lptk	2pkc	2prk	3prk E	• •	3.4.21.66 4 cha					
Itec E	lthm	2tec E	3tec E			3,4.21.68 10 ch					
lpk2	Ipml A	Ipml B	Ipml C	ltpg	ltpk A	1tpk B 2.3.4.21.76 4 ch	1tpk C	ltpm	1tpn		
lfuj A	Ifuj B	lfuj C	1fuj D	IV	•		latits				
Ісру	1whs A	lwhs B	1wht A	Serine-ty 1 wht B		frolase fold idase E.C.3.4.16	6.5 6 chains				
-				Itah B		.1.1.3 10 chains Itah C	s Itah D	3tgl	4tgl		
Ihpl A	1hpl B	Hpa A	Ilpa B		β-Ba	rrel fold		Jigi	41g1		
1apo	lccf	lheg A	lheg B	v		A E.C.3.4.21.6					
lkdu	11mw B	Ilmw D		-	_	or E.C.3.4.21.7					
lhpg A				Glutam	yl endopeptidas	e II E.C.3.4.21.	82 1 chain				
				С		-Glu catalytic t frolase fold	triad				
Lala A	tala P	lerl		11pn		.1.1.3 10 chains	s Hpm	11po	Ithg	ltrh	
Icle A	1cle B			Ace	tylcholinesteras	e E.C.3.1.1.7 9	chains	Imah F	7.11.6		
lace	lacj	lack	lacl		$\alpha/\beta$ Hy	lfss B -Asp catalytic drolase fold		Illian I			
ledb	ledd	lede	2dhc	2dhd	2dhe	nase E.C.3.8.1.5 2eda	2edc	2had			
				С		<b>-Trp catalytic</b> ( 3 Fold	triad				
lesc	lesd	1ese		Strep		s E.C.3.1.1.50	3 chains				
	1000	. 030		C	$\alpha + \beta$ Cysteir	-Asn catalytic ne proteinase fo	ld				
lpad	lpe6	lpip A	1pod A	lppd	¹1ppn	4.22.2 12 chair 1ppp	1stf E	2pad	4pad	5pad	6pad
laec	2act					3.4.22.14 2 chai					
					Caricain E.C.	3.4.22.30 1 cha	ın				



**Fig. 5.** Comparison of the catalytic triads from chymotrypsin 1*cho* class 1 (Fujinaga et al., 1987), acetylcholinesterase 1*ace* class 2 (Sussman et al., 1991), haloalkane dehalogenase 2*dhc* class 3 (Verschueren et al., 1993), and *S. scabies* esterase 1*esc* class 4 (Wei et al., 1995). The reference frame His residues are superimposed, allowing us to compare the relative conformations of the nucleophilic and electrostatic side chains.

noted that the main-chain atoms of both the Ser and His residues of the acetylcholinesterase triad originate from different sides when compared to the class 1 Ser-His-Asp catalytic triad. However, the difference in the handedness of these triads is not relevant to our consensus template because we are interested in the functional atoms and not the main-chain scaffold.

We derived such a template by averaging the functional consensus templates created for classes 1, 2, and 3. This class 1-2-3 template was compared with the consensus templates derived for each of the five classes and the RMS differences are summarized in Table 5. Figure 6 shows the distribution of individual RMS deviations (RMSDs) for each of the triads in all class 1, 2, 3, and 4 proteins from the class 1-2-3 template. The majority of triads in classes 1-3 have an RMSD between 0.3 Å and 1.3 Å-triads above this value are the class 1 triads with inhibitors bound to their active sites. All three triads in class 4 cluster around 2.0 Å RMS distance despite the fact that two of the structures are crystallized with inhibitors. Figure 5 indicates that, in class 4, the S. scabies triad Ser 14 residue is in a different position with respect to the His residue and this is reflected in the high RMSD. Given the different tertiary fold, it would seem likely that this different conformation is a consequence of convergent evolution. Whether the catalytic mechanism differs in this enzyme compared to classes 1, 2, and 3 is a matter for speculation (Wei et al., 1995).

# Class 1-2-3 template search through the PDB

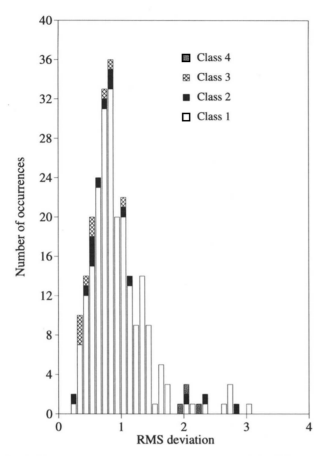
It is interesting to see if the arrangement of the atoms in the class 1-2-3 consensus template, in terms of the functional oxygens sur-

**Table 5.** Coordinates of the functional templates for the class 1-2-3 template and those of the five separate classes <sup>a</sup>

Residue	Atom	X	У	z
	Class 1-2-3 funct	tional consensu	s template	
Ser/Asp	$O^{\gamma}/O^{\delta_1}$	4.9	0.9	-0.3
Glu/Asp	$O^{\epsilon_1}/O^{\delta_1}$	-0.4	-3.8	0.1
		1: Ser-His-Asp	068	
		ss 1-2-3 templat	e 0.6 A	
Ser	$O^{\gamma}$	4.9	0.8	-0.3
Asp	$O^{\delta_2}$	0.4	-3.7	0.1
		2: Ser-His-Glu	061	
	RMS from cla	ss 1-2-3 templa	te 0.6 A	
Ser	$O^{\gamma}$	4.9	1.3	0.3
Glu	$O_{\epsilon_1}$	-0.6	3.7	0.3
		3: Asp-His-Asp		
		ss 1-2-3 templa	te 0.2 A	
Asp	$O^{\delta_1}$	4.8	0.8	-0.3
Asp	$\mathrm{O}^{\delta_2}$	-0.5	-3.6	0.2
		4: Ser-His-Trp		
		ss 1-2-3 templa		
Ser	$O^{\gamma}$	3.2	3.1	-0.4
Trp	О	-0.4	-3.4	-0.9
		5: Cys-His-Asn	0	
Cys	$S^{\gamma}$	0.5	-4.0	2.1
Asp	$\mathrm{O}^{\delta_2}$	4.4	2.3	0.5
	His te	emplate residue		
His	$C^{\beta}$	-1.4	-0.1	-0.0
His	$C^{\gamma}$	0.0	0.0	0.0
His	$N_{\hat{\alpha}}^{\delta_1}$	0.8	-1.1	0.0
His	$C^{\delta_2}$	0.8	1.1	0.0
His	C <sup>€1</sup>	2.1	-0.7	-0.0
His	$N^{\epsilon_2}$	2.1	0.6	-0.0

<sup>a</sup>Each 3D template is superimposed onto the same His residue. The RMS distances of the individual templates from the class 1-2-3 template are also given.

rounding the His side-chain reference frame, occurs elsewhere in the PDB. We searched through a data set of 639 representative protein structures in the January 1995 PDB. Some of the structures in classes 1, 2, and 3 were present in this data set, others having been excluded on the basis of having greater than 95% sequence identity. A triad located in this nonidentical data set was considered interesting if its RMSD was less than 2.0 Å from the class 1-2-3 consensus template and there was an Asp or Ser side-chain atom in the position equivalent to Nu:, the nucleophilic group, and an Asp or Glu side-chain atom in the position equivalent to ELEC, the electrostatic group. We found a number of proteins, not members of classes 1, 2, and 3, having the characteristic Nu:-His-ELEC triad conformation. Two of the triads found resemble the Ser-His-Asp catalytic triad of the class 1 enzymes; cyclophilin 2cpl and immunoglobulin 2ig2. These triads have already been discussed in detail



**Fig. 6.** Distribution of RMSDs of observed catalytic triads in all the structures in classes 1, 2, 3, and 4 from the class 1-2-3 template in the June 1996 PDB

previously (Wallace et al., 1996). A further six proteins have triads resembling the Asp-His-Asp triads of the class 3 enzymes. These are listed in Table 6 in order of increasing RMSD from the template. We take the first of these, the nitrogenase molybdenum-iron protein, as an example.

# Nitrogenase molybdenum-iron protein E.C.1.18.6.1

The MoFe protein from Azotobacter vinelandii is an  $\alpha_2/\beta_2$  tetramer and the X-ray structure has been determined to 2.7 Å resolution, PDB code 1min (Jongsun & Rees, 1992). It is part of the nitrogenase enzyme system, which consists of two metalloproteins, the molybdenum-iron (MoFe) protein and the iron Feprotein. There is another cofactor, the P-cluster pair, which is thought to transfer electrons between the 4Fe:4S cluster Fe-protein and the MoFe cofactor. The precise catalytic mechanism is unknown, but the  $N_2$  substrate is proposed to bind directly to the MoFe cofactor. The MoFe cofactor is bound to homocitrate and is surrounded by water molecules; it may be a source of protons for the formation of the NH<sub>3</sub> product.

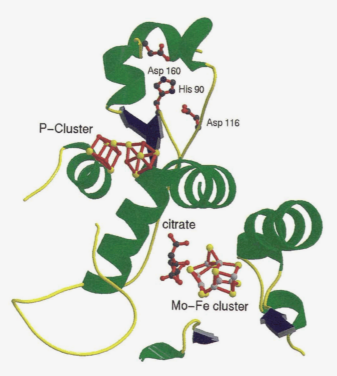
A close-up view of the Asp-His-Asp triad and the MoFe and P-cluster cofactors is shown in Figure 7. The Asp 160-His 90-Asp 116 triad and the P-cluster are located within about 8 Å of each other. The triad has not been mentioned or implicated in the reaction course and whether it is involved in proton or electron transferral is unknown. Closer inspection reveals that the Asp 116

**Table 6.** List of the potential catalytic triads found when the 95% nonidentical data set of PDB structures was searched with the class 1-2-3 catalytic triad template

	Putativ	e Asp-His-Asp c	atalytic tria	ds	
	Res.				
Residue	number	Atom	X	У	z
	Niti	rogenase molybd	enum-iron		
		Protein RMSD			
		1min(D)			
Asp	160	$\mathrm{O}^{\delta_2}$	5.0	1.4	-0.
Asp	116	$\mathrm{O}^{\delta_2}$	-0.5	-3.4	0.
His	90	Side chain	_	_	_
	Py	ruvate oxidase E	.C.1.2.3.3		
		RMSD 0.5			
		1pox(D)			
Asp	69	$\mathrm{O}^{\delta_1}$	4.7	1.8	-0.
Asp	27	$\mathrm{O}^{\delta_2}$	-0.4	-3.6	0.
His	28	Side chain	_	_	_
	ľ	Macromycin RMS 2mem	SD 1.08		
Asp	100	$\mathrm{O}^{\delta_2}$	5.8	1.9	-0.
Asp	53	$O^{\delta_1}$	0.2	-3.5	-0.
His	32	Side chain	_	_	_
		R2 of ribonucleo E.C.1.17.4.1 RMS 1rib(A)		ase	
Asp	237	$O^{\delta_2}$	4.3	2.1	-0.
Asp	84	$O^{\delta_1}$	-1.4	-4.8	0.
His	118	Side chain	_	_	_
	Super	roxide dismutase RMSD 1.5 1sos(D)		1	
Asp	124	$\mathrm{O}^{\delta_1}$	4.1	2.1	-0.
Asp	83	$O^{\delta_1}$	1.6	-3.7	0.
His	71	Side chain	_	_	_
		dehyde-3-phospha E.C.1.2.1.12 RMS 1gdl(O)		genase	
Asp	312	$\mathrm{O}^{\delta_2}$	6.2	1.8	-1.
Asp	47	$O^{\delta_1}$	-1.3	-4.7	0.
His	50	Side chain	_	_	

is, in fact, accessible to the surface of the protein, indicating that it could have access to solvent or ligands.

We checked the sequence of the MoFe protein (PDB code 1min) against the SWISS-PROT (Bairoch & Boeckmann, 1994) database using the automatic sequence alignment program BLAST (Altschul et al., 1990) and obtained several MoFe protein sequences, all derived from nitrogen-fixing bacterium, although the sequence identity with 1min may be as low as 25%. The His residue is conserved in all but one case, and there are several instances of a Ser or a Thr in the position of Asp 116. The Asp at position 160 is conserved in all cases.



**Fig. 7.** A view of the MoFe protein from *Azobacter vinelandii* showing the positional relationship of the P-cluster, the Asp-His-Asp triad and the MoFe cofactor associated with the nitrogenase enzyme (Jongsun & Rees, 1992).

These results show this to be a region worthy of further investigation, which may give insight into the catalytic mechanism of MoFe proteins.

# The ribonucleases

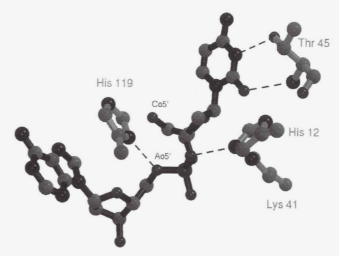
Ribonucleases are found in both prokaryotes and eukaryotes (Beintema et al., 1990); they catalyze the hydrolysis of phosphodiester bonds in RNA chains. Structurally, the best understood ribonucleases are bovine pancreatic ribonuclease (RNase A, E.C.3.4.27.5) and ribonuclease  $T_1$  (RNase  $T_1$ , E.C.3.4.27.3) from the fungus *Aspergillus oryzae*. Both enzymes have an  $\alpha+\beta$  roll architecture, but adopt a different topology.

The crystal structures of ribonuclease H and barnase have also been solved, but they will not be considered here.

# Ribonuclease A

Ribonuclease A has 124 amino acids and is a member of a large superfamily of homologous bovine RNases (Beintema et al., 1988). RNase A is a monomer, with an  $\alpha+\beta$  fold and four intrachain disulfides. Ribonuclease S (RNase S, e.g., 1rbc; Varadarajan & Richards, 1992) is a product of the cleavage of RNase A by subtilisin between residues 20 and 21.

RNase A is specific for the pyrimidines uridine and cytidine; this specificity is achieved by hydrogen bonding from the backbone NH and side-chain -OH atoms of the sequentially conserved residue Thr 45. Chemical modification studies (Crestfield et al., 1963) have shown that the residues His 12 (acid/base catalyst), His 119 (acid/base catalyst), and Lys 41 (stabilizes transition state) are involved in the catalytic mechanism (Blackburn & Moore, 1982).



**Fig. 8.** 3D representation of the active site of ribonuclease A complexed with D(CPA) (Zegers et al., 1992). The catalytic residues are His 119, His 12, and Lys 41.

Figure 8 is a 3D representation of the inhibitor deoxycytidyl-3',5'-deoxyadenosine, D(CPA) bound to ribonuclease A 1*rpg* (Zegers et al., 1992). The His 12 interacts with the 2'-oxygen (equivalent to CO5' in the inhibitor in Fig. 8) and Lys 41 is in the vicinity.

Borkakoti et al. (1982) noticed that in a phosphate complex of RNase A, there are two distinct conformations of His 119 (A and B) that are related by a 180° rotation about the His 119  $C^{\beta}$ - $C^{\gamma}$  ( $\chi_2$ ) bond. There is still controversy as to which of these conformations is catalytically active (Borkakoti et al., 1982; deMel et al., 1992; Zegers et al., 1992).

Two consensus templates have been created describing both the A and B conformations (Table 7). The seed atoms used were the His 12 side chain, His 119  $N^{\delta 1}$ , and Lys 41  $N^{\zeta}$  taken from the structure 3rn3 (Borkakoti et al., 1982), which has coordinates for both the A and B His 119 conformations. Figure 9 is a 3D representation of the distribution of the His 119  $N^{\delta 1}$  atoms relative to His 12 for all structures in the ribonuclease A and ribonuclease S entries in the PDB. There are two distinct clusters of His 119  $N^{\delta 1}$  atoms (in blue) representing the A and B conformations; Table 8 lists the PDB structures responsible for these clusters. Figure 10 is a histogram of the number of hits versus RMSD from the B conformation of the RNase template for all the ribonuclease A and S PDB structures (Table 9); it confirms that the A and B conformation occupy distinct positions for the His 119  $N^{\delta 1}$  atom.

# Template search through the PDB

The 95% by sequence nonidentical protein data set was searched for other proteins with residues and atoms in a similar conformation to the two RNase A consensus templates. An RMS distance cut-off of 2 Å was used when defining matches to either template.

All the hits located for the A conformation, other than the RNase structures in this data set, have RMSDs greater than 2 Å; they can therefore be discounted as unlikely to have RNase A catalytic activity. The B conformation consensus template located His 285, His 321, and Lys 191 from the A and B chains of the same enzyme RUBISCO, ribulose-1,5-bisphosphate carboxylase, E.C.4.1.1.39 (5rub, Schneider et al., 1990) with RMSDs from the consensus templates of 1.51 Å and 1.86 Å, respectively. Figure 11 is a 3D

**Table 7.** Coordinates of the consensus templates that describe the two conformers, A and B, for the active site of ribonuclease A as well as the coordinates of the consensus template for ribonuclease  $T^a$ 

Residue	Number	Atom	X	У	z
		Ribonuclea	se A		
	Templa	te conformer	A coordinate	es	
Lys	41	$N^{\zeta}$	5.3	-1.1	-2.8
His	119	$N^{\delta_1}$	6.2	3.2	3.8
		Ribonuclea	se A		
	Templa	te conformer	B coordinate	es	
Lys	41	$N^{\zeta}$	5.1	-1.4	-2.9
His	119	$N^{\delta_1}$	5.8	5.8	2.0
		Ribonuclea	se T		
His	92	$\begin{array}{c} N_2^\epsilon \\ C^\delta \end{array}$	9.7	2.5	-2.1
Glu	58	$C^{\delta}$	3.4	2.8	-2.6
Glu	58	$O^{\epsilon_2}$	3.6	3.3	-1.5
Glu	58	$O_{\epsilon_1}$	4.3	2.3	-3.2
	R	eference fram	e atoms		
His	12/40	$C^{\gamma}$	0.0	0.0	0.0
His	12/40	$N^{\delta_1}$	0.8	-1.1	0.0
His	12/40	$C^{\delta_2}$	0.8	1.1	0.0
His	12/40	$C^{\epsilon_1}$	2.1	-0.7	0.0
His	12/40	$N^{\epsilon_2}$	2.1	0.6	0.0

<sup>&</sup>lt;sup>a</sup>The two templates share the same reference frame His 12/40 coordinates.

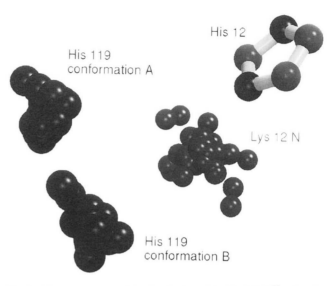
representation of the active site of RUBISCO from 5rub (Lundqvist & Schneider, 1989) with substrate ribulose-1,5-bisphosphate. Also shown is the active site Mg that is coordinated to Asp 193 (yellow bonds) and the substrate. The residues located, His 285, His 321, and Lys 191 (red bonds), are found in the large L chain and are also part of the active site. In addition, they are all conserved in the RUBISCO sequence (Schneider et al., 1990). Lys 191 is the site of carbamylation during activation, and His 321 is involved in binding the phosphate group of the substrate (Lundqvist & Schneider, 1989). There is no clear functional role assigned to His 285. When compared to ribonuclease A, the position of the active site residues with respect to the substrate is different and the equivalent residues also have different roles. However, it is interesting that the RNase template has identified the catalytic residues in RUBISCO even though they have different mechanisms, different primary E.C. numbers, and a totally different function.

# Ribonuclease T<sub>1</sub>

The bacterial RNase T<sub>1</sub> is isolated from *A. oryzae*. The enzyme has two isoforms containing Lys or Gln at position 25 of the polypeptide chain, denoted Lys 25-RNase T1 and Gln 25-RNase T1.

# Specificity and catalytic mechanism

RNase T<sub>1</sub> is specific for the purine nucleotide guanosine (as opposed to pyrimidines in RNase A) and is strictly limited to hydro-



**Fig. 9.** 3D representation of the distribution of the His 119  $N^{\delta 1}$  active site atom conformations A and B for all the RNase A and RNase S structures in the PDB. Also shown is the side chain of His 12 and the distribution of the Lys 41  $N^{\zeta}$  atoms.

lysis at 3'-phosphate groups in RNA. The reason for this specificity is not fully understood.

The catalytic residues are Glu 58, which takes a proton from  $\rm H_2O$  in the first step, and His 92, which donates a proton to the leaving group and activates the water molecule used in hydrolysis. The guanosine-specific recognition occurs in the loop Tyr 42-Asn 43-Asn 44-Tyr 45-Glu 48.

# The consensus template

The atoms used to generate a consensus template were the side chain of His 40, His 92 N $^{\epsilon 2}$ , and Glu 58 C $^{\rm gd}$ , O $^{\epsilon 1}$ , O $^{\epsilon 2}$  taken from the RNase T $_1$  X-ray crystal structure 1m1 (Arni et al., 1992); the distance cut-off was set to 3.0 Å. Table 9 gives the data set of RNase T $_1$  PDB codes and their RMSDs from the resultant consensus template.

There are two structures in the ribonuclease  $T_1$  data set, 2aae (Zegers et al., 1992) and 5rnt (Lenz et al., 1991), whose active site residues are not identified by the consensus template. 2aae has its His 40 mutated to a lysine and 5rnt's structure is only refined to 3.2 Å and has the inhibitor guanosine-3',5'-bisphosphate bound to its active site.

No potential ribonuclease  $T_1$  active sites were located when the 95% nonidentical-by-sequence data set of PDB structures was searched.

# Comparison of ribonuclease A and $T_1$ active sites

It is interesting to compare the three different templates, RNase A conformations A and B and RNase  $T_1$ . Figure 12 is a 3D representation of these templates with the atoms superimposed according to their proposed role in the catalytic mechanism. As Zegers et al. (1992) have shown, the general base catalysts, His 12 and Glu 58, the electrostatic stabilizing groups, Lys 41 and His 40, and the general acid catalysts, His 119 and His 92, of RNase A and RNase  $T_1$ , respectively, all superimpose, indicating convergent evolution of these enzyme active sites.

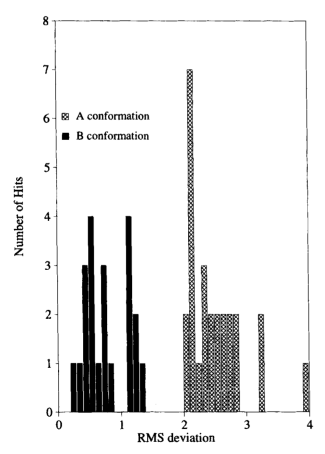
**Table 8.** Summary of the ribonuclease PDB structures and their RMSDs from their respective consensus templates that adopt either the A or B conformation of their active site His 119 residue<sup>a</sup>

	Te	mplate conform	ner A: Ribonucl	ease E.C.3.1.27	7.5	
		1	Ribonuclease A			
1bsr B 0.45	3rn3 1.16	1ras 1.50	1rat 0.90	2rat 0.74	3rat 0.92	4rat 0.79
5rat 0.31	6rat 0.43	7rat 0.48	8rat 0.44	9rat 0.41	1rbn 0.62	1rcn E 0.71
1rnc 0.35	1rnd 0.46	1rar 1.03	1rob 0.49	1rpg 0.33	1rph 0.53	1rtb 1.56
5rsa 0.34	6rsa 0.53	7rsa 0.48	1rtb 1.56		_	
	Te	mplate conforn	ner B: Ribonucl	ease E.C.3.1.2	7.5	
			Ribonuclease A	\		
1bsr B 0.64	1rbn 0.97	3rn3 1.05	1rpf 1.29	1rph 0.78	9rsa B 0.71	1srn A 0.65
3srn A 0.49	4srn A 0.32	1ssa A 0.78	1ssb A 0.50	-		
			Ribonuclease S	3		
1rbc S 1.00	1rbd S 0.30	1rbe S 0.74	1rbf S 1.50	1rbg S 0.49	1rbh S 0.44	1rbi S 0.40
2rln S 0.29	1rnu 1.36	1rnv 0.54	2rns 1.38	•		

<sup>&</sup>lt;sup>a</sup>PDB codes in bold have coordinates describing both conformations.

# Lysozyme

Lysozyme kills certain bacteria by cleaving the polysaccharide component, N-acetylglucosamine (NAG) and N-acetylmuramate (NAM), of their cell wall. There are two catalytic residues, Glu 35



**Fig. 10.** Histogram of the number of hits against RMS distance from the B conformation of the RNase A consensus template. It shows that the A and B conformations of the His 119 residues are in distinct positions.

and Asp 52, in the eukaryotic lysozyme (Blake et al., 1965; Johnson & Phillips, 1965) and Glu 11 and Asp 20 in the prokaryotic lysozyme from bacteriophage T4 (Matthews & Remington, 1974).

The prokaryotic and eukaryotic lysozyme structures have low sequence identity, but have a similar  $\alpha+\beta$  tertiary fold (Remington & Matthews, 1978) and may have evolved from a common precursor. They can, however, be divided into eukaryotic and prokaryotic groups on the basis of the geometry of their catalytic residues.

Two other lysozymes, characterized by a low sequence identity and different tertiary fold to those mentioned above, are found in the goose (154l, Weaver et al., 1995) and black swan (1gbs, Rao et al., 1995). In fact, they also lack a catalytic Asp and for this reason will not be considered further.

# Eukaryotic: Mammalian and avian lysozyme

There are 76 eukaryotic lysozyme structures in the PDB from several species, although many of these originate from mutational studies of the same structure. Figure 12 shows 3D representations of the conformation of the Glu 35 and Asp 52 catalytic residues in the mammalian and avian lysozymes, together with the corresponding catalytic residues in the bacteriophage T4 lysozymes. In all cases, the Glu residues have been superimposed so that the relative conformation of the Asp side chain can be compared. Although the conformations of the Asp side chains are similar, in all the lysozymes there is no obvious clustering of the functional Asp  $O^{\delta}$ atoms. This is in contrast to the situation in the Ser-His-Asp catalytic triads, where we have shown that, although the relative position and orientation of the catalytic Asp shows great variability, the location of the functional Asp  $O^{\delta}$  is highly conserved. In the lysozyme catalytic mechanism, the negative charge on Asp 52 stabilizes the formation of the positive charge on the sugar in the D-site (Phillips, 1966). This suggests that the Asp 52 side-chain carboxyl group only needs to be in the vicinity of the D-site sugar and its precise position is not so important.

A consensus template was constructed for the mammalian and avian data set using the seed template atoms of the Glu 35 side chain and Asp 52  $C^{\gamma}$ ,  $O^{\delta 1}$ , and  $O^{\delta 2}$  from the PDB structure 135*l* (Harata et al., 1993) of human lysozyme with a 3.0 Å cut-off; the

**Table 9.** Summary of the ribonuclease  $T_1$  PDB structures and their RMSDs from the ribonuclease  $T_1$  consensus template<sup>a</sup>

1fus 1.08 1rnlA 1.53 9rnt 1.77	1fut 1.28 1rnlB 1.53 1trpA 0.85	1rga 1.28 1rnlC 0.62 1trpB 1.00	1rgeB 1.40 1rnt 0.99 1trgA 0.87	1rgcA 1.92 2rnt 0.47 1traB 0.83	1rgk 1.78 3rnt 0.76 <b>2aae</b>	1rgl 0.66 6rnt 0.41 <b>5rnt</b>	1rls 1.71 7rnt 0.90	1rms 1.56 8rnt 1.78
--------------------------------------	---------------------------------------	---------------------------------------	---------------------------------------	---------------------------------------	---------------------------------------	---------------------------------------	------------------------	------------------------

<sup>&</sup>lt;sup>a</sup>PDB codes in bold are missed by the RNase T<sub>1</sub> template using a 3.0-Å distance cut-off.

coordinates are given in Table 10. Nine hen-egg lysozyme structures, which are different refinement models based on the original Blake et al. (1965) lysozyme structure (Diamond, 1975), were removed. Although they gave extraordinary insight into lysozyme structure and function at the time, they are poorly refined compare to the lysozyme structures solved in the last 15 years. Table 11 summarizes the number of catalytic Glu 35–Asp 52 diads located using the template for each species.

There are three hen-egg white lysozyme structures whose Glu 35-Asp 52 catalytic residues are not located. Two of these, 1lym (Rao et al., 1983) and 3lyt (Young et al., 1993), are only refined to 2.5 Å, and the third, 2hfl (Sheriff et al., 1987), has been crystallized in a complex with an immunoglobulin molecule. This also applies to three structures of mouse lysozyme (e.g., Fischmann et al., 1991), which are not identified. The catalytic residues in horse lysozyme (Tsuge et al., 1992) have an RMSD of 2.52 Å from the consensus

template; Figure 12 shows that the relative position of the Asp 52 is shifted with respect to the other mammalian Asp residues.

# Prokaryotic: Bacteriophage T4 lysozyme

T4 lysozyme is produced late in the infection of *Escherichia coli* by T4 bacteriophage. The structure was first determined by Matthews and Remington (1974). This enzyme has been used as a model to study the effects of mutations on protein stability and function and there are 160 different mutant forms deposited in the PDB (e.g., Alber et al., 1987; Weaver & Matthews, 1987).

A consensus template was constructed from the wild-type T4 structure 2lzm (Alber et al., 1987) using the same atoms as for the mammalian template; the distance cut-off was set to 3.0 Å. The coordinates of the resultant template are given in Table 10.

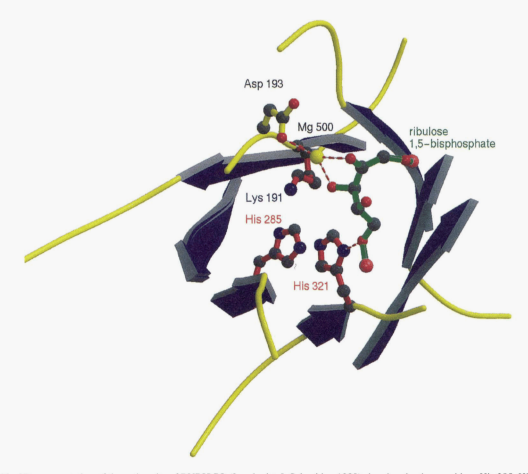
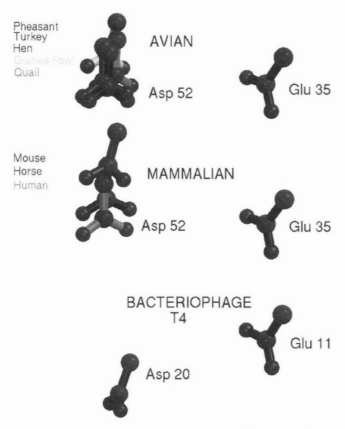


Fig. 11. 3D representation of the active site of RUBISCO (Lundqvist & Schneider, 1989) showing the three residues His 285, His 321, and Lys 191 (red bonds) that have the same conformation as the active site residues of ribonuclease A.



**Fig. 12.** Relative conformations of the catalytic Glu 35-Asp 52 residues from avian and mammalian lysozymes as well as the bacteriophage T4 catalytic Glu 11-Asp 20.

Of the 160 T4 structures in the PDB, 7 are not located by the consensus template. All "missed" structures have mutations of the active site residues or of regions around the active site, perturbing the conformation of the catalytic residues.

# Comparison of prokaryotic and eukaryotic lysozymes

Due to the similarity of the tertiary fold of both the prokaryotic and eukaryotic lysozyme structures, and because they both have a Glu and Asp residue as their catalytic residues, it would not be surprising if the conformation of the catalytic residues is also the same. In fact, as Figure 12 indicates, when the Glu residues of the two templates are superimposed, the Asp residues lie around 4.5 Å apart. It is proposed that the general mechanism of catalysis of these two lysozyme is the same. As for the mammalian and avian lysozymes, Asp 20 acts as an electrostatic stabilizer of the positively charged sugar in the D-site, so it is only required to be in the vicinity of the D-site.

When the prokaryotic and eukaryotic lysozyme consensus templates are used to search through the representative structures of the PDB, there are about 100 hits located for each template. This large number of matches is not surprising given that the template consists of only two residues. This shows that, although the lysozyme templates are able to extract the majority of catalytic Glu-Asp diads, there are many false hits involving noncatalytic interactions, and so the template has a low specificity.

**Table 10.** Coordinates of the consensus template describing the active site of the prokaryotic T4 as well as mammalian and avian lysozymes present in the PDB

Residue	Residue number	Atom	X	y	z
	Mam	malian and a	vian Asp 52		
Asp	52	$C^{\gamma}$	4.0	-4.3	4.1
Asp	52	$O^{\delta_1}$	4.4	-3.8	5.1
Asp	52	$\mathrm{O}^{\delta_2}$	4.4	-4.3	3.1
		Phage t4 A	sp 20		
Asp	20	$C^{\gamma}$	7.5	-3.6	1.5
Asp	20	$O^{\delta_1}$	4.4	-3.8	5.1
Asp	20	$\mathrm{O}^{\delta_2}$	4.4	-4.3	3.1
		Template re	esidue		
Glu	11/35	$C^{\gamma}$	-1.5	0.1	-0.0
Glu	11/35	$C^{\delta}$	0.0	0.0	0.0
Glu	11/35	$O^{\epsilon_1}$	0.6	-1.0	0.0
Glu	11/35	$O^{\epsilon_2}$	0.6	1.1	0.0

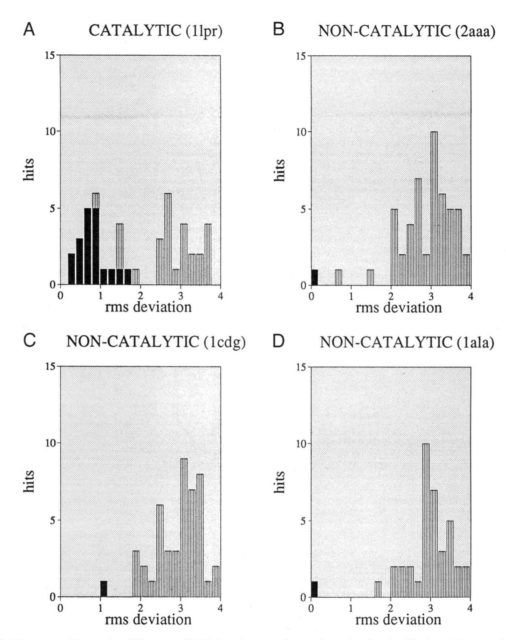
# Discussion

We have developed an algorithm called TESS that is based on the geometric hashing paradigm. The program enables us to search through a data set of 3D PDB structures for any user-defined, sequence order-independent 3D template. The seed for the 3D template consists of atoms or residues extracted directly from a PDB file, and it is possible to define explicitly for each coordinate which atoms or residue types should be included. This work enables us to create a collection of 3D enzyme active site or functional templates, that, by analogy to the 1D templates present in secondary protein sequence databases such as PROSITE or PRINTS, may allow the assignment of the biological function and evolutionary origins of a new protein structure. Of course, TESS could

**Table 11.** Numbers of Asp and Glu catalytic residues located for lysozymes originating from different species

	Total no. PDB	
Name	structures	Hits
	Avian lysozymes	
Guinea fowl	1	1
Hen	25	22
Pheasant	1	1
Quail	1	1
Turkey	4	4
	Mammalian lysozymes	
Mouse	4	1
Horse	1	1
Human	17	17

# SER-HIS-ASP SEARCH (O-HIS-O)



**Fig. 13.** Histograms of the number of hits versus RMSD from the respective templates when four Ser-His-Asp triads are used to search through the nonidentical data set. The 1*lpr* triad is the catalytic consensus template for the serine proteinases and lipases. The other three are randomly chosen noncatalytic triads. Black bars are diads originating from the same protein by function as the template diad.

be used to produce databases of other recurring 3D templates, such as metal or nonenzyme ligand binding sites. In addition, the templates are useful as a purely practical method to identify automatically the catalytic triad in, for example, a serine proteinase. It also allows us to orient all serine proteinases into the same reference frame, to facilitate simple visual comparisons.

Having studied all the enzymes in the PDB with the catalytic residues of type Nu:-His-ELEC, the conservation of these triads is striking. Although the residue types of the Nu: and ELEC groups

can vary according to enzyme type, we find that the positions of the functional atoms in the triads are conserved. Indeed, one template is able to describe the active site of all the serine proteinases, acetylcholinesterase, and haloalkane dehalogenase proteins. This suggests that convergent evolution has drawn the functional atoms into optimal catalytic positions. In addition, these triads are confined to the active sites of enzymes, although we did find several examples of other unidentified Nu:-His-ELEC triads that may have biological relevance. This suggests

that, as the number of protein structures deposited in the PDB increases, other common 3D templates, functional in more than one enzyme type, will occur.

Conversely, the Asp-Glu catalytic diad found in all lysozyme structures in the PDB does not have such a conserved structure. The Asp is thought to act as an electrostatic stabilizer of the transition state and therefore its precise position in the active site is not so critical. In addition, when the lysozyme Glu 35-Asp 52 consensus templates are used to search through the representative structures of the PDB, there are about 100 hits for each of the templates. This larger number of hits occurs simply because there

is greater chance of two rather than three residues being located randomly in the same conformation as the catalytic pair in a given data set of protein structures.

To investigate this, we have taken three randomly picked noncatalytic Ser, His, and Asp interactions and compared the number of hits located when these triads are used to search our representative nonidentical data set of PDB structures with those for a catalytic triad. Secondly, we took three randomly picked noncatalytic Asp and Glu residues and compared the number of hits located in the protein data set with the catalytic Asp and Glu from lysozyme. The representative nonidentical data set is different from

# **ASP-GLU SEARCH**

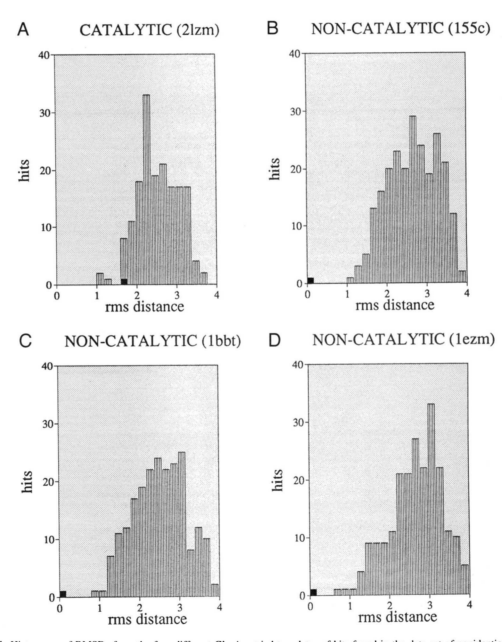


Fig. 14. Histograms of RMSDs from the four different Glu-Asp triad templates of hits found in the data set of nonidentical protein structures. Black bars represent diads located from the same protein by function as the template triad.

the 95% nonidentical data set used previously and comprises protein structures that all have an SSAP score (Orengo et al., 1993) less than 80 when compared and less than 25% sequence identity (i.e., it excludes homologous proteins). This was extracted from the June 1996 release of the PDB.

The histogram in Figure 13 (top left) represents the search with the catalytic triad consensus template derived from the Ser 195  $O^{\gamma}$ -His 195 side-chain-Asp  $102^{\delta 2}$  catalytic triad from 1lpr (Bone et al., 1991). The other three histograms are searches using noncatalytic Ser-His-Asp interactions that have an RMSD between 3 Å and 6 Å from the 1lpr consensus template; these are Ser 202  $O^{\gamma}$ -His 205 side-chain-Asp 172  $O^{\delta 2}$  from chicken annexin 1ala (Bewley et al., 1993), Ser 13 O $^{\gamma}$ -His 503 side-chain-Asp 518 O $^{\delta 2}$ from cyclodextrin glycosyltransferase 1cdg (Lawson et al., 1994), and Ser 104 O<sup> $\gamma$ </sup>-His 108 side-chain-Asp 201 O<sup> $\delta$ 1</sup> from  $\alpha$ -amylase 2aaa (Boel et al., 1990). For the catalytic triad search, 20 other catalytic triads are located from the data set of structurally nonidentical proteins, compared to only one for each of the noncatalytic Ser, His, and Asp interactions. In addition, for the catalytic triad, there are only a few hits below 2.0 Å RMS distance that are noncatalytic. although other factors, such as accessibility to substrate, are important, this does suggest that when Ser, His, and Asp are found in this conformation in a protein structure, we can be confident that they play a catalytic role.

A similar test was performed on a template consisting of only two residues; here we compared the nonidentical data set search of the T4-lysozyme catalytic Asp 11-Glu 20 diad with three other randomly chosen noncatalytic Glu-Asp interactions. These diads were the side chains of Glu 11-Asp 109 from cytochrome C550 155c (Timkovich & Dickerson, 1976), Glu 128-Asp 169 from foot-and-mouth virus 1bbt (Parry et al., 1990) and Glu 172-Asp 189 from the elastase structure 1ezm (Thayer et al., 1991). Figure 14 shows the number of hits located for searches with these templates. There are around the same number of hits located for the catalytic as the noncatalytic diads.

This proves that, as expected, the number of hits located when searching a data set of protein structures with a consensus template depends on the number of atoms and residues in that template. If hits are located below the chosen RMS cut-off, it does not necessarily mean that they are functionally significant, but merely provides a possible starting point for further experimental investigation. Indeed, when such a hit is located, other factors should be considered, such as locality with respect to potential ligand binding site or accessibility to the protein surface.

These results are borne out by the searches using the ribonuclease and lysozyme templates. Although there were several hits below the defined RMSD cut-off, there is no example of a hit revealing a new catalytic site. It should be noted that the functions of the vast majority of proteins in the PDB are understood and the PDB represents a small fraction of all proteins in the genome.

One of the major problems in genome sequence analysis is the lack of information available on functional sites. Even for proteins of known structure, the relevant information is often hidden in the literature and cannot be extracted easily. These catalytic templates represent one approach to encapsulating functional information, and relating it back from structure to sequence. As the number of templates grows, it will become a useful source of additional information on functionally important residues in a sequence, which may be useful in recognizing distant homologues and so assigning function.

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