

## The Predicted Structure of Immunoglobulin D1.3 and Its Comparison with the Crystal Structure

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Predictions of the structures of the antigen-binding domains of an antibody, recorded before its experimental structure determination and tested subsequently, were based on comparative analysis of known antibody structures or on conformational energy calculations. The framework, the relative positions of the hypervariable regions, and the folds of four of the hypervariable loops were predicted correctly. This portion includes all residues in contact with the antigen, in this case hen egg white lysozyme, implying that the main chain conformation of the antibody combining site does not change upon ligation. The conformations of three residues in each of the other two hypervariable loops are different in the predicted models and the experimental structure.

ANTIBODIES ACHIEVE THEIR RANGE of specificity in antigen recognition through the modulation of the conformation of specific loops by changes in the amino acid sequence. An understanding of how the conformations of these loops are determined is essential for an appreciation of the basic mechanism of antibody affinity and specificity, and for the design of modifications of immunoglobulin structures.

We describe here the prediction of the conformation of the antigen-binding domains of immunoglobulin G (IgG) D1.3, and the comparison of the predicted models with the experimental structure determined from the 2.8 Å resolution electron density map (1). Two methods of prediction were used. One was based on analysis of the known structures of the variable (V) domains of light (L) and heavy (H) chains ( $V_L$  and  $V_H$ ) (2) of the myeloma proteins NEWM, McPC603, KOL, REI, and RHE (Table 1). From these structures, we identified the relatively few residues that, through packing, hydrogen bonds, or ability to assume unusual conformations (Gly, Pro, Asn, Asp), are primarily responsible for the conformations of the regions in these domains involved in antigen binding: the hypervariable loops. In contrast, de la Paz *et al.* have based predictions on the length and overall sequence homology of the hypervariable regions (3). Our other predictions were based on conformational energy calculations (4).

Comparisons of the atomic structures of variable domains confirm and refine the distinction between a conserved framework consisting of the  $\beta$  sheets of the individual

domains and the  $V_L$ - $V_H$  packing, and hypervariable loops in the regions that link  $\beta$  sheets (5-11). In the five known  $V_L$  structures, 69 residues in two  $\beta$  sheets have the same conformation. The root-mean-square differences in position ( $\Delta$ ) of their main

Table 1. Immunoglobulin variable domains known to atomic resolution from x-ray crystal structure analysis. A Fab is a fragment of an immunoglobulin containing variable domains  $V_L$  and  $V_H$  and constant domains  $C_L$  and  $C_H1$ .

Protein	Chain type		Resolution (Å)	Reference
	L	H		
Fab' NEWM	$\lambda$ I	$\gamma$ II	2.0	(6)
Fab McPC603	$\kappa$	$\gamma$ I	2.7	(7)
Fab KOL	$\lambda$ I	$\gamma$ III	1.9	(8)
$V_L$ REI	$\kappa$		2.0	(9)
$V_L$ RHE	$\lambda$ I		1.6	(10)

Table 2. Residues buried in sheet-sheet packings in immunoglobulin variable domains.

At the interface between the two sheets in  $V_L$  domains:  
4, 6, 19, 21, 23, 25, 33, 35, 37, 47, 48, 62, 64, 71, 73, 75, 82, 84, 86, 88, 90, 97, 99, 101, 102, 104

At the interface between the two sheets in  $V_H$  domains:  
4, 6, 18, 20, 22, 24, 34, 36, 38, 48, 49, 51, 69, 78, 80, 82, 86, 88, 90, 92, 104, 106, 107, 109

At the interface between the  $V_L$  and  $V_H$  domains:  
 $V_L$ : 36, 38, 44, 87, 98  
 $V_H$ : 37, 39, 45, 47, 91, 93, 103

chain atoms are between 0.5 and 1.6 Å for different pairs of  $V_L$  domains. The  $\beta$ -sheet residues adjacent to the hypervariable regions differ in position by less than 1.5 Å (2). Similarly, 79 residues in the  $\beta$  sheets of three  $V_H$  domains have the same conformations. The relative geometry of the  $V_L$ - $V_H$  packing is also conserved (12). This conserved framework structure is produced by conservation of residues buried between the sheets and at the  $V_L$ - $V_H$  interface. In the monoclonal antibody IgG D1.3 (13), 56 of the 62 buried framework residues (Table 2) are identical to those in known structures; the other six differ in size by no more than a methyl group. We predicted the framework structure of D1.3 to be the same as that of the known structures.

The antigen-binding site contains three hypervariable loops from  $V_L$  and three from  $V_H$ , denoted L1, L2, L3, and H1, H2, H3 (11). The residues, numbered according to Kabat *et al.* (5), are:

$V_L$		$V_H$	
Region	Residues	Region	Residues
L1	26-32	H1	26-32
L2	50-52	H2	53-55
L3	91-96	H3	96-101

These residues are outside the common  $\beta$ -sheet framework of the molecules of known structure. [The complementarity-determining regions (CDR's) defined by Kabat *et al.* (5) on the basis of sequence comparisons are more extensive.] The homology of the hypervariable loops of D1.3 with those of known structures of the same class is shown in Table 3. Fewer than 10 of the 32 residues in the hypervariable regions of IgG D1.3 are the same as those in homologous positions in any single known structure.

We analyzed the determinants of the conformation of the hypervariable loops.

L1: The structure of the L1 region is characteristic of the class of the domain (14). The  $V_\kappa$  class includes the  $V_L$  domains of REI, McPC603, and the target structure D1.3 (Table 1). In McPC603, L1 is 13 residues long and 7 in REI. The common residues have the same main chain conformation: for 26 to 29 and 32,  $\Delta = 0.5$  Å.

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They adopt an extended conformation, across the top of the  $V_L$  domain. The six additional residues in McPC603 form a loop extending away from the domain. The L1 conformation is determined primarily by packing of residue 29 against residues 2, 25, 33, and 71 in the interior of the framework. In D1.3, L1 has the same length as in REI, and the residues packing against the framework are conserved:

Residue:	2	25	29	33	71
D1.3	Ile	Ala	Ile	Leu	Tyr
REI	Ile	Ala	Ile	Leu	Tyr
McPC603	Ile	Ser	Ile	Leu	Tyr

We predicted L1 of D1.3 to have the REI conformation.

L2: In four  $V_L$  structures L2 is a three-residue turn (in NEWM, L2 is deleted). Their structures are similar, as conformations of short turns are restricted by the allowed values of the main chain torsion angles (15). We predicted that L2 in D1.3 would also have this common structure.

L3: The L3 regions in REI and McPC603 are each six residues long and have the same structure:  $\Delta = 0.4 \text{ \AA}$ . The conformations usually observed for such hairpin loops would involve main chain hydrogen bonds between residues 92 and 95. The residue Pro 95, present in most  $V_L$  domains including REI, McPC603, and D1.3, precludes this conformation (Fig. 1). In REI and McPC603, Pro 95 has a *cis* peptide, and the side chains of Asn 90 (REI) and Gln 90 (McPC603) form hydrogen bonds to L3 main chain atoms. In D1.3, L3 is also six residues long and includes Pro 95 and His 90. If His 90 could form the same hydrogen bonds as Gln or Asn do in REI and McPC603, we should expect a similar L3 conformation in D1.3.

H1: In KOL and McPC603 the seven residues have similar conformations ( $\Delta = 0.4 \text{ \AA}$ ). In KOL and McPC603, Gly 26 produces a sharp turn. Residues 28 to 32 are a helix with Phe 29 packing in a cavity formed by the side chains of residues 34 and 94 and the main chain of 72 and 77. In D1.3, H1 also has seven residues, and the important side chains are similar but not all identical:

Antibody	Residue				
	26	27	29	34	94
D1.3	Gly	Phe	Leu	Val	Tyr
KOL	Gly	Phe	Phe	Met	Tyr
McPC603	Gly	Phe	Phe	Met	Tyr

In particular, two important residues are both smaller in D1.3. The total volume of Leu 29 and Val 34 in D1.3 is  $65 \text{ \AA}^3$  less than

the total volume of Phe 29 and Met 34 in KOL and McPC603.

H2: In KOL, NEWM, and McPC603, H2 differs in size and conformation. In NEWM, the three residues in H2 are part of a seven-residue hairpin turn with a Gly in the fourth position. Such turns have the same conformation whenever they occur in known structures (15). As H2 in D1.3 also has three residues and a Gly at this position, we predicted it to have the NEWM conformation.

H3: The length and conformation of H3 differ in KOL, NEWM, and McPC603. Medium and large H3 loops extend from the surface of the domain. In McPC603 the

conformation of the stem of H3 is determined primarily by the packing of Tyr 100b into a cavity between the  $V_L$  and  $V_H$  domains, its OH group remaining accessible. The homologous residue in D1.3, Arg 99, is also large. We predicted it would pack in the same manner to produce a similar H3 conformation.

We began the construction of the model based on the analysis of known structures with the framework regions of  $V_L$  REI and  $V_H$  KOL. Of those domains of known structure of the same classes as those of D1.3, these had been determined at the highest resolution. The overall sequence homologies with D1.3 are 66 percent ( $V_L$ )

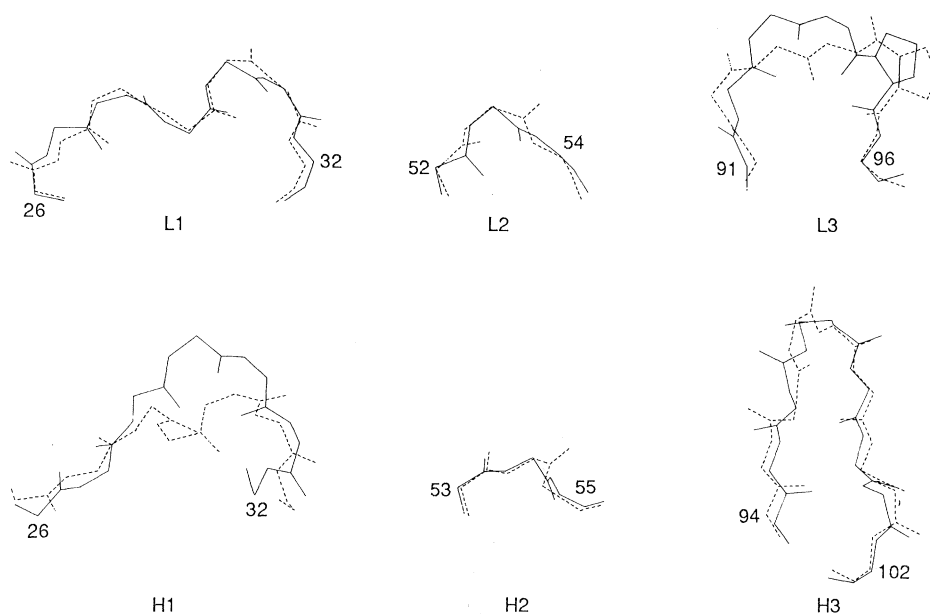


Fig. 1. The main chain atoms of the hypervariable regions in the experimental structure (---) and in the predicted model based on analysis of the known structures (—). In a 2.8 Å electron density map, the exact orientations of peptides outside secondary structures may not be well determined. This figure contains independent pictures of these regions, each shown in a viewpoint perpendicular to the mean plane of the loop.

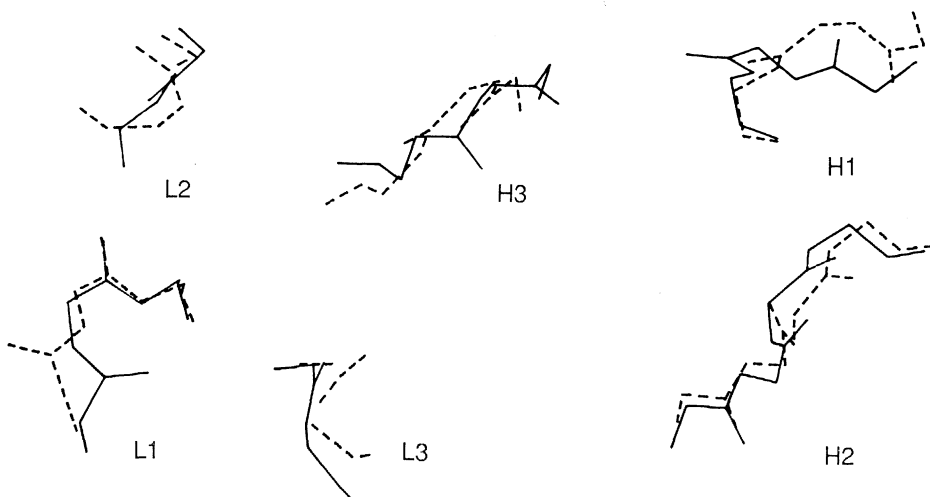


Fig. 2. The main chain atoms of the part of the binding site that makes contact with the antigen (I), in the experimental structure (---) and in the predicted model based on analysis of the known structures (—). The actual relative positions of the regions in the structure are shown from a viewpoint roughly perpendicular to the plane of the antigen-antibody interface.

and 49 percent ( $V_H$ ). To combine the domains we superposed  $V_L$  REI onto  $V_L$  KOL by fitting the conserved  $V_L$  residues of the domain-domain interface. We took L1 and L2 from  $V_L$  REI and H2 from  $V_H$  NEWM. For L3, noting the conserved Pro, and believing on the basis of model building that His 90 could make the same hydrogen bonds as Gln, we used the loop from REI. In the H1 region of D1.3, two crucial residues are smaller than those in the known structures. Nevertheless, we did not expect this loop to change its conformation radically, and took H1 from  $V_H$  KOL.

No known structure has an H3 region of the same length as D1.3. Faced with alternative conformations in known structures, we first chose to build a loop based on NEWM H3 (for reasons we now believe to be incorrect). Informed that this initial prediction was qualitatively incorrect, we then based our prediction on McPC603, which proved to be the proper model. To correct the length we removed three residues from the apex of the loop—these were chosen to preserve the pattern of hydrogen bonding—and closed the chain using the interactive graphics program FRODO (16). Side chains of D1.3 were substituted wherever its sequence differed from the parent structures, and bad steric contacts relieved by changing their conformations.

Two other models were constructed independently with the use of conformational energy calculations. One was based exclusively on McPC603, the molecule of known structure having the highest homology with D1.3 in both V regions. The side chains of McPC603 were replaced with their D1.3 homologs, and insertions and deletions made without, initially, preserving the continuity of the backbone. Equivalent atoms were taken from the McPC603 structure whenever possible; 125 missing atoms (out of 1977) were placed at random within 1 Å of the atom of known position to which they are bonded. Energy minimization closed the gaps in the chain; eliminated close contacts; formed strong hydrogen bonds; and brought all bond lengths, bond angles, and torsion angles close to their standard values (4). After 600 conjugate gradient cycles, the total energy was -1655 kcal/mol.

A second model merged features from each of the three known Fab structures, by selecting seven stretches of residues locally most homologous in sequence to D1.3: From McPC603  $V_L$ , residues 1 to 17 and 27 to 107; KOL  $V_L$ , 18 to 26; NEWM  $V_H$ , 1 to 67 and 90 to 101; and KOL  $V_H$ , 68 to 89 and 102 to 115. Only 100 atoms were missing. Minimization reduced the total energy to -1681 kcal/mol.

We compared the predicted structures

with the experimental structure of D1.3 derived from the 2.8 Å electron density map of a crystal containing the Fab fragment of D1.3 binding the antigen hen egg white lysozyme (1). In Fig. 1, the experimental hypervariable regions and those predicted from structural analysis are superposed. The observed antigen-binding site (1) and the homologous regions of this predicted structure are shown in Fig. 2. Deviations in atomic positions of observed and predicted structures are shown in Tables 4 to 7. Results are given for the framework structures, the residues adjacent to the hypervariable regions, the hypervariable regions themselves, and a maximal set of well-fitting residues that includes the lysozyme binding site. Atoms included in superposing the individual loops are N, C $\alpha$ , C, and C $\beta$ . The

carbonyl O is omitted because in a 2.8 Å electron density map the orientations of peptides outside secondary structures may not be well determined. The C $\beta$  is included to evaluate the general positioning of side chains.

In this model, the framework and the folds of L1, L2, H2, and H3 were predicted correctly (Table 6). Three residues in the middle of L3 are different. In the current interpretation of the electron density map, the peptide of Pro 95 has a *trans* conformation; in the predicted structure it is *cis* (Fig. 1). Although the *trans* conformation is favored by the current map, a *cis* conformation cannot be ruled out. Three residues at the center of H1 are very different; in particular, in the experimental structure the side chain of Leu 29 does not lie in the cavity occupied

Table 3. The homology of the hypervariable loops of IgG D1.3 and other immunoglobulins of known structure and the same class. References to the structures are given in Table 1. Residue numbers for the hypervariable regions are given in the text.

L1	REI	Ser Gln Asp Ile Ile Lys	Tyr
	McPC603	Ser Gln Ser Leu Leu Asn Ser Gly Asn Gln Lys Asn Phe	
	D1.3	Ser Gly Asn Ile His Asn	Tyr
	REI	Glu Ala Ser	
L2	McPC603	Gly Ala Ser	
	D1.3	Tyr Thr Thr	
	REI	Tyr Gln Ser Leu Pro Tyr	
L3	McPC603	Asp His Ser Tyr Pro Leu	
	D1.3	Phe Trp Ser Thr Pro Arg	
H1	KOL	Gly Phe Ile Phe Ser Ser Tyr	
	NEW	Gly Thr Ser Phe Asp Asp Tyr	
	McPC603	Gly Phe Thr Phe Ser Asp Phe	
	D1.3	Gly Phe Ser Leu Thr Gly Tyr	
H2	KOL	Asp Asp Gly Ser	
	NEW	Tyr His Gly	
	McPC603	Asn Gly Lys Asn Lys	
	D1.3	Gly Asp Gly	
H3	KOL	Gly Gly His Gly Phe Cys Ser Ser Ala Ser Cys Phe Gly Pro Asp	
	NEW	Leu Ile Ala	Gly Cys Ile Asp
	McPC603	Tyr Tyr Gly Ser	Thr Trp Tyr Phe Asp
	D1.3	Arg Asp	Tyr Arg Leu Asp

by Phe 29 in KOL and McPC603, but is external. Although the experimental structure may have to be modified, as the three residues have main chain conformations usually disallowed because of steric hindrance, the predicted structure is incompatible with the electron density. In retrospect, it appears that the reduced volume of two

Table 4. Fit of  $V_L$ - $V_H$  framework. Comparisons of predicted and observed structures. In Tables 4 to 7, the model SA is based on structure analysis, and model CE on conformational energy calculations starting from a structure derived initially from McPC603.  $\Delta$  is the root-mean-square deviation in angstroms after superposition.

Residues	$\Delta$ (N, C $\alpha$ , C, and O atoms)	
	SA	CE
$V_L$ : 4-6, 19-25, 33-48, 52-54, 61-76, 84-90, 97-107	1.0	1.0
$V_H$ : 3-12, 17-25, 33-52, 56-60, 68-82, 88-95, 102-113		

Table 5. Shifts of framework residues adjacent to hypervariable regions. Shift is the difference (angstroms) in position of residue after superposition of the framework.

Residues	Shift			
	SA		CE	
$V_L$ : 25 33	0.6	0.3	2.5	0.8
48 52	0.6	0.8	0.2	0.6
90 97	0.8	0.4	0.9	0.8
$V_H$ : 25 33	0.9	1.8	0.6	1.2
52 56	1.5	1.8	1.7	0.7
95 102	1.2	1.0	1.3	0.6

Table 6. Fits of hypervariable loops.

Residues	$\Delta$ (N, C $\alpha$ , C, and C $\beta$ atoms)	
	SA	CE
L1 26-32	0.85	3.76
L2 49-51	0.63	0.47
L3 91-96	0.97	1.10
H1 26-32	2.07	1.68
H2 53-55	0.50	0.89
H3 99-104	0.86	0.87

Table 7. Fit of maximal well-fitting portion, including framework and all lysozyme contact residues (model SA).

Residues	$\Delta$ (N, C $\alpha$ , and C atoms)
$V_L$ : 2-6, 19-65, 69-92, 96-104	1.0
$V_H$ : 3-7, 14-27, 30-59, 67-75, 78-81, 84-113	

important side chains (29 and 34) may have changed the packing to produce a different fold.

None of the residues at which the model differs from the observed structure is involved in antigen contacts. The main chain conformation of the binding site itself is predicted accurately (Fig. 2).

Both the models based on conformational energy calculations correctly predicted the framework and the folds of L2, H2, and H3. Loop L1 is very different from the experimental structure because it was built from the extended part of McPC603 L1 rather than from the part common to other  $V_K$  chains. L3 and H1 are similar to those in the model based on structure analysis.

The comparisons show that the prediction of the main chain conformation of D1.3 was largely successful (Tables 4 to 7 and Figs. 1 and 2). They support the premise that the binding site conformation is determined principally by specific interactions of a few residues, and that these residues can be identified and used to formulate rules valid for structure prediction. A structure of D1.3 at higher resolution will permit a more detailed evaluation of the predictions, as well as the refinement and extension of the rules.

The predictions treated an isolated  $V_L$ - $V_H$  dimer. The observed D1.3 structure is contained in a Fab-antigen complex. The close similarity of the observed and predicted D1.3 binding site and framework implies

that the association with antigen does not significantly alter the main chain conformation of the antibody.

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## Cambrian River Terraces and Ridgetops in Central Australia: Oldest Persisting Landforms?

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**Fluvial sediments in paleovalleys cut in the Ashburton surface of the Davenport province of central Australia form terrace remnants that appear to retain their original depositional tops and have probably existed as subaerial landforms since their inception. Marine fossils in sediments conformable with the fluvial sediments near the southeast margin of the province give a Cambrian age for the terraces; the Ashburton surface forming the ridgetops between the paleovalleys is Cambrian or older.**

INLAND AUSTRALIA IS WELL KNOWN for the flatness of its landscape and the antiquity of its erosion surfaces (1, 2). The highest surface in the Tennant Creek region of central Australia, the Ashburton surface, has long been thought to be Cretaceous or older (2). We report evidence from the Davenport province of the Tennant Creek region (Fig. 1) that suggests that, during the Cambrian, fluvial sediments were deposited in valleys between ridges whose tops are remnants of the Ashburton surface. Subsequent dissection of the sediments

formed terraces and mesas, but where preserved, the relation of the terraces to the adjoining ridges indicates that the terraces and mesatops have existed as subaerial landforms since the Cambrian sedimentation. The Ashburton surface itself is therefore Cambrian or older.

The Davenport province (3) is a broad

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