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Conformations of immunoglobulin hypervariable regions

Cyrus Chothia*†, Arthur M. Lesk*‡, Anna Tramontano‡, Michael Levitt*§, Sandra J. Smith-Gill, Gillian Air, Steven Sheriff***, Eduardo A. Padlan*. David Davies*, William R. Tulip**, Peter M. Colman**, Silvia Spinelli**. Pedro M. Alzari# & Roberto J. Poljak#

- * MRC Laboratory of Molecular Biology, Hills Road, Cambridge CB2 2QH, UK
- ‡ European Molecular Biology Laboratory, Meyerhofstrasse 1, Postfach 1022.09, D-6900 Heidelberg, FRG
- § Department of Cell Biology, Stanford University Medical School, Stanford, California 94305, USA
- † Christopher Ingold Laboratory, University College London, 20 Gordon Street, London WC1H0AJ, UK
- National Institute of Cancer and # National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, Maryland 20892, USA
- ¶ Department of Microbiology, University of Alabama, Birmingham, Alabama 35294, USA
- †† CSIRO Division of Biotechnology, 343 Royal Parade, Parkville 3052, Australia
- ‡‡ Unité d'Immunologie Structurale, Département d'Immunologie, Institut Pasteur, 25 rue du Dr Roux, 75724 Paris, France

On the basis of comparative studies of known antibody structures and sequences it has been argued that there is a small repertoire of mainchain conformations for at least five of the six hypervariable regions of antibodies, and that the particular conformation adopted is determined by a few key conserved residues. These hypotheses are now supported by reasonably successful predictions of the structures of most hypervariable regions of various antibodies, as revealed by comparison with their subsequently determined structures.

THE relationships between the amino-acid sequences of immunoglobulins and the structures of their antigen-binding sites are important for understanding the molecular mechanisms of the generation and maturation of the immune response and for designing engineered antibodies. Antigen-binding sites are formed by six loops of polypeptide, the hypervariable regions; three from the variable domain of the light chain (VL) and three from the variable domain of the heavy chain (VH), denoted L1, L2, L3, and H1, H2, H3, respectively (Fig. 1a). Within the domains, the loops are connected to a β -sheet framework whose structure is conserved^{1,2}. The specificity and affinity of the binding sites are governed by the structures of the six hypervariable regions^{3,4}.

Two models can be proposed for the relationship between the amino-acid sequence and structure of the binding-site loops. In one model, different sequences produce different conformations for both the main chain and side chains of the loops. Because hypervariable regions have different sequences in different antibodies, this model implies that each region adopts a different conformation in different antibodies. In the other

** Present address: The Squibb Institute for Medical Research, PO Box 4000, Princeton, New Jersey 08543-4000, USA.

model, antibodies have only a few main-chain conformations or 'canonical structures' for each hypervariable region. Most sequence variations would only modify the surface provided by the side chains on a canonical main-chain structure. Sequence changes at a few specific sets of positions would switch the main chain to a different canonical conformation.

Canonical structure model

Experimental evidence indicates that the canonical structure model describes the relationship between amino-acid sequence and structure for at least five of the six hypervariable regions⁵⁻⁹. Kabat et al.5 found conserved residues at sites within certain sets of hypervariable regions and suggested that they had a structural role. Padlan and Davies⁶, and more recently de la Paz et al.⁷, showed that some of the hypervariable regions in the immunoglobulins of known structure have the same main-chain conformation in spite of several differences in sequence.

Chothia and Lesk⁸ identified the residues that through packing, hydrogen bonding, or the ability to assume unusual values of the torsion angles ϕ , ψ or ω , are primarily responsible for the main-chain conformations of the hypervariable regions in the structures then known—the Fab fragments of NEW (ref. 10), McPC603 (ref. 11), KOL (ref. 12) and J539 (ref. 13) and the VL domains of REI (ref. 14) and RHE (ref. 15). The conformations are determined by the interactions of a few residues at specific sites in the hypervariable regions and, for certain loops, in the framework regions. Hypervariable regions that have the same conformations in different immunoglobulins have the same or very similar residues at these sites (Fig. 1 and Table 1). Examination of the amino-acid sequence of the antibody D1.3 showed that its hypervariable regions are the same size as those in known structures and contain the same or similar residues at the sites responsible for known conformations. On the basis of these observations the atomic structure of the VL-VH dimer of D1.3 was predicted before its experimental determination. Comparison of this predicted structure with the preliminary crystal structure showed that the conformations of four of the hypervariable regions had been predicted correctly; the conformation of L3 was significantly different from that predicted, and

TABLE 1 Sequences and conformations of V_K and VH hypervariable regions of known structure

L1 Regions†																		
Canonical																		
Structure	Protein	26	27	28	29	30	31	а	b	С	d	е	f	32	2	25	33	71
					*										*	*	*	*
1	J539	S	S	S	V	S		_		_	_	_	_	S	1	Α	L	Υ
	HyHEL-5	S	S	S	V	N				_		_		Υ	1	Α	M	Υ
	NQ10	S	S	S	V	R	_	_	_			-	_	Υ	1	Α	М	Υ
2	REI	S	Q	D	1	1	K	_						V	ı	Α	L	Υ
	D1.3	S	Ġ	N	- 1	H	N				_	_	_	Υ	1	Α	L	Υ
	HyHEL-10	S	Q	S	1	G	N	_	_	_	_	_		N	1	Α	L	F
	NC41	S	Q	D	V	S	Τ	-				_	_	Α	1	Α	L	Υ
3	McPC603	S	Ε	S	L	L	N	S	G	N	Е	K	N	F	1	S	L	F
4	4-4-20	S	Q	S	L	V	Н	S	_	N	G	N	Т	Υ	V	S	L	F

Total no. of sequences known for L1 regions: human, 95; mouse, 299. Canonical structure 2 3 4 1 Human sequences that fit (%) 5

Mouse sequences that fit (%)

20

10

51

52

L2 Regions

Canonical Structure

REI McPC603 J539 D1.3 HvHEL-5 HvHEL-10

Protein

NC41

NO10

D Y

50

w

D

S S S S S

G G G G G G G G

48

64

G

4-4-20 S Total no. of sequences known for L2 regions: human, 69; mouse, 183.

Canonical structure Human sequences that fit (%)

Human sequences that ht (70)	30
Mouse sequences that fit (%)	95
L3 Regions	

Canonical Structure	Protein	91	92	93	94	95	96	90
						*		*
1	REI	Υ	Q	S	L	Р	Υ	Q
	McPC603	D	Н	S	Υ	Р	L	N
	D1.3	F	W	S	Т	Р	R	Н
	HyHEL-10	S	N	S	W	Р	Υ	Q
	NC41	Н	Y	S	Р	Р	W	Q
	4-4-20	S	Т	Н	V	Р	W	Q
	NQ10	W	S	S	N	Р	L	Q
2	J539	W	Т	Υ	ř	L	1	Ž
3	HyHEL-5	w	G	R	N	*		*

Total no. of sequences known for L3 regions: human, 52; mouse, 152.

Canonical structure Human sequences that fit (%) 2 3

Mouse sequences that fit (%)

2 1

H1 had a very different fold from that predicted⁹. (We report below that the refined conformation of D1.3 corresponds more closely to the predicted structure.)

An examination of the library of the known immunoglobulin sequences shows that many immunoglobulins have hypervariable regions that are the same size as those in the known structures and contain the same or closely related residues at the sites responsible for the known conformations⁸. These observations indicate that for at least five of the hypervariable regions there is only a small repertoire of canonical main-chain conformations and that the conformation actually present can often be predicted from the sequence by the presence of specific residues.

The accuracy of the canonical structure model for immunoglobulin binding sites depends on (1) the correct determination of the sets of residues responsible for the observed conformations and (2) changes in the identity of residues at other sites not significantly affecting the conformations of the canonical structures. The model can be tested, refined and extended by using it to predict the atomic structures of binding sites in immunoglobulins before their structures have been determined by X-ray crystallography.

We have now tested the canonical structure model by using it to predict the structures of four immunoglobulins before their structures had been experimentally determined. These immunoglobulins are HyHEL-5 (ref. 16), HyHEL-10 (ref. 17), NC41 (ref. 18) and NQ10 (S.S., P.M.A. and R.J.P., manuscript in preparation). The analysis of the amino-acid sequences of these immunoglobulins indicated that 19 of their 24 hypervariable regions should have conformations close to known canoni-

H1 Regions‡												
Canonical												
Structure	Protein		26	27	28	29	30	31	32		34	94
			*	*		*					*	*
1	McPC603		G	F	T	F	S	D	F		M	R
	KOL		G	F	- 1	F	S	S	Υ		M	R
	J539		G	F	D	F	S	K	Υ		M	R
	D1.3		G	F	S	L	Τ	G	Υ		V	R
	HyHEL-5		G	Υ	Т	F	S	D	Υ		1	R
	NC41		G	Υ	T	F	Т	N	Υ		M	R
	NQ10		G	F	T	F	S	S	F		M	R
	4-4-20		G	F	T	F	S	D	Υ		М	G
1'	NEW		G	S	Т	F	S	N	D		Υ	R
	HyHEL-10		G	Đ	S	1	Т	D	D		W	Ν
Total no. of sequer Canonical structure Human sequences Mouse sequences	that fit (%)	1 50 80										
H2 Regions§												
Canonical												
Structure	Protein		52a	ı b	С	53	54	55		71		
								*				
1	NEW		_	_	_	Υ	Н	G				
	D1.3					G	D	G				
	HyHEL-10		_		_	Υ	S	G				
			*					(*)		*		
2	HyHEL-5		Р		_	G	S	G		Α		
-	NC41		T	_	_	N	Т	G		L		
							(*)			*		
3	KOL		D			D	G	S		R		
3	J539		P	_	_	D	S	G		R		
	NQ10		S			G	S	S		R		
	INOTO		5			G						
							(*)	*		*		
4	McPC603		N	K	G	N	K	Y		R		
	4-4-20		N	K	Р	Υ	N	Υ		R		
Total no. of sequer	nces known for H2	2 region	ns: humai	n, 54; m o	ouse, 248.							
Canonical structure				3	4							
Canonical Structure	?	1	2	3								
Human sequences		15	1	40	15 20							

The residues listed here (single-letter code) are those that form the hypervariable regions and those in the framework regions that are important for the observed conformations of these regions. The hypervariable regions are taken as those outside the framework β -sheet. Except for H2, they are similar to, but not identical with the regions that show high sequence variations and which Kabat $et~al.^{26}$ use to define hypervariable regions. The sequences are grouped so that those that have the same main-chain conformation, or canonical structure, are adjacent. The canonical structure numbers used below refer to the conformations shown in Fig. 1. The residues in the hypervariable and framework regions that are mainly responsible for these conformations are indicated by an asterisk. The classification and sequence requirements of the H2 conformations have been revised in the light of work described here and elsewhere²⁸. For each hypervariable region the number of human and mouse sequences listed by Kabat $et~al.^{26}$ are given. We also give the percentage of these sequences that are the same size as the known canonical structures and have the same residues at the positions marked by an asterisk.

† Canonical structure 4 is illustrated in Fig. 4. Although the size of the known L1 structures varies between 6 and 13 residues, they have closely related folds with residues 26–19 and 32 packed against the framework in the same conformation⁸. The remaining residues form a turn or loop on the surface (Figs 1 and 4). The ends of the long loops have some flexibility. There are another 25% of the human sequences and 20% of the mouse sequences that have one more residue than structure 2, or one fewer than structure 4, and whose sequences satisfy the requirements listed above. It is expected that these differ only in the conformations of the tips of the surface loops.

‡ The H1 hypervariable regions with canonical structure 1 have very similar conformations: the r.m.s. differences in the coordinates of their main-chain atoms are 0.3–0.8 Å. The H1 regions in NEW and HyHEL-10 only partly satisfy the sequence requirements for structure 1 and have a distorted version of its conformation.

§ The H2 region here comprises residues 52a-55. The region with high sequence variation is 50-65 (ref. 26). In the known structures the main-chain conformation of 50-52 and 56-63 do not differ significantly (Fig. 1b). (*), The residues at positions 55 or 54 in the canonical structures 2, 3 and 4 have residues with positive values for ϕ and ψ , and usually, but not in all cases, Gly, Asn or Asp is found at these sites. For a sequence to match that of canonical structure 2, 3 or 4 the presence of these residues at sites 54 or 55 is required.

cal structures. We then compared the predicted structures of these hypervariable regions with the subsequently determined structures. Another immunoglobulin structure, 4-4-20 (ref. 19) has recently been reported. We did not have the opportunity to predict the structure of 4-4-20 before its experimental determination, and we discuss here only how its hypervariable regions have the conformations expected from the known canonical structures. Also, we report that the refined conformation of D1.3 (ref. 20) corresponds more closely to the predicted structure.

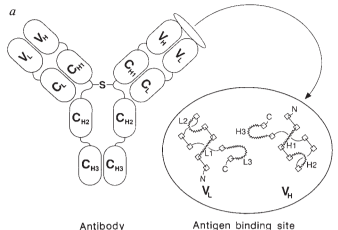
Model building procedure

The main-chain conformations of the hypervariable regions in the V_K and VH domains of known structure are shown in Fig. 1. The residues responsible for these conformations are listed

in Table 1. Each hypervariable region in the immunoglobulins of unknown structure was examined to determine (1) whether it has the same size as any homologous hypervariable region of known structure and (2) whether its sequence contains the set of residues responsible for a known conformation. Except for L3 in HyHEL-5, all the light-chain regions correspond to a known canonical structure, as do all the H1 regions and the H2 region in HyHEL-10 (Table 1). The conformation predicted for the H2 regions in NC41 and HyHEL-5 was based on the analysis of the H2 region in the preliminary structure of J539 (ref. 8). In all three of these antibodies the H2 region is a four-residue turn with Gly at the fourth position and the predicted conformation is that almost always found for such turns²¹. (Below we present a more accurate analysis of H2 regions.) For H3 regions in HyHEL-5, HyHEL-10, NC41 and NQ10, no prediction of

conformation could be made on the basis of the known canonical structures.

The sequences of the VL and VH domains were compared to see which of the known framework structures have sequences close to those of the unknown structures. From the comparisons



of the hypervariable and framework regions, one known VL and one known VH structure were taken as the starting points—parents—for the model of the predicted structure. If the conformation predicted for a hypervariable region was not present in the parent, but was present in another known structure, the hypervariable region in the parent was replaced by that in the other structure. Side chains in the parent that were different from those in the unknown structure were replaced^{22,23} and the resulting model subjected to a very limited energy refinement²⁴.

HyHEL-5, HyHEL-10 and NC41 hypervariable regions

The atomic structures of Fab fragments of immunoglobulins HyHEL-5, HyHEL-10 and NC41 in complexes with their protein antigens were determined by X-ray crystallography¹⁶⁻¹⁸. The resolution of the X-ray data used to determine the structures and value of the residual (R) after refinement are (complex, resolution, R value): HyHEL-5-lysozyme, 2.5 Å, 20%; HyHEL-10-lysozyme, 3.0 Å, 24%; and NC41-neuraminidase, 2.9 Å, 19%. These structures have been determined at medium resolution. The tracing of the polypeptide chain of the hypervariable regions is unambiguous, although the orientation of some of

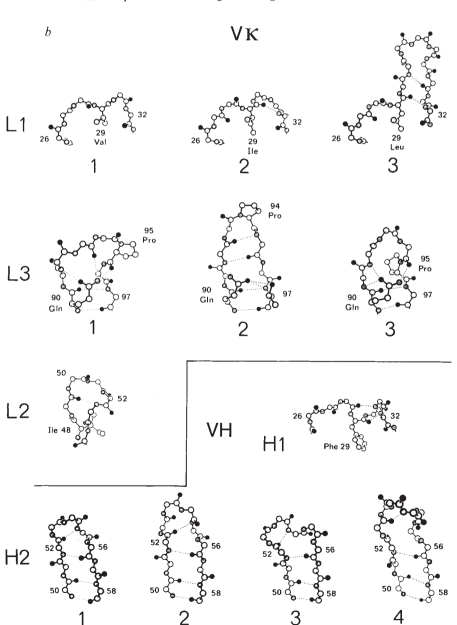


FIG. 1 a, Antigen-binding sites of immunoglobulins are formed by six loops of polypeptide, three from the VL domain L1, L2 and L3 and three from the VH domain H1, H2 and H3 (wavy lines). These loops are attached to strands (\square) of a conserved β -sheet. b, Canonical structures for the hypervariable regions of Vk and VH domains. In each drawing the region is viewed so that the accessible surface is at top and the framework region below. The main-chain conformation and some of the side chains that determine this conformation are shown. For the definition of the hypervariable regions used here, see Table 1. The immunoglobulins in which the different canonical structures occur are listed in Table 1.

the peptide groups is uncertain. Most side chains are unequivocally placed.

Figure 2 shows the predicted and observed structures of each of the hypervariable regions, superposed by a least-squares fit of their main-chain atoms. Table 2a gives, for each predicted and observed hypervariable region, the r.m.s. difference in position of the main-chain atoms.

The main-chain conformations of the predicted and observed hypervariable regions are very similar (Fig. 2, Table 2a). The only exception is the H1 region of HyHEL-5. Although the observed and predicted conformations of residues 26-29 and 32 are the same, residues 30 and 31 are in quite different positions. The recently refined structure of Fab J539 (T. N. Bhat, E.A.P. and D.D., manuscript in preparation) shows that these differences were inherited as a result of an error in the original determination of the J539 structure used as the parent for this region. Rebuilding the predicted model with the refined J539 structure puts residues 30 and 31 in the correct position and gives an r.m.s. difference between the predicted and observed H1 regions of 0.6 Å.

Given the medium resolution of the structures used to derive the models and of the experimental structures, the agreement of the predicted and observed loop conformations is excellent.

Relative positions of the hypervariable regions

Figure 3 shows the positions of the hypervariable regions relative to each other and to the framework for the predicted and observed structures. To produce this figure the predicted and observed structures were superposed by a least-squares fit of framework residues. In Table 2b, the differences in position of the hypervariable regions are reported.

Small differences in the relative positions of the hypervariable regions in the predicted and observed structures might be expected because of two factors not corrected for in the model build-

ing. First, the predicted structures were built using parent V domains that have some residues in the framework and VL-VH interface that are different from those in the final predicted structure. These differences have small but significant effects on the main-chain structure of the individual domains and the way they pack together 2,25,26. Second, differences between the predicted and observed structures could arise from the effects of the association with the antigen 18. In other proteins, ligand binding can result in the movement of close-packed segments of polypeptide relative to each other by 1-2 Å, and the ends of loops are able to move somewhat more 27.

The differences in the positions of the predicted and observed H2 regions in HyHEL-5 and NC41 (Table 2b) are larger than expected from these factors. The interactions that the H2 regions make with the rest of the VH domain were therefore examined.

Residue 71 and position and conformation of H2

At the same time as the structures of HyHEL-5 and NC41 became available, the refinement of the atomic structure of J539 was completed (T. N. Bhat, E.A.P. and D.D., manuscript in preparation). The conformation of H2 in the refined structure is not like that in HyHEL-5 and NC41 but is the same as that in KOL. This was quite unexpected. The main determinant of the conformation of small turns is usually the position of glycine residues²¹: in KOL, Gly occurs at position 54, and in J539, HyHEL-5 and NC41, Gly occurs at position 55.

An examination of the environments of the H2 regions²⁸ shows that in KOL and J539 the side chain of framework residue Arg 71 packs between, and forms hydrogen bonds to, H1 and H2. In HyHEL-5, residue 71 is Ala, and here the cavity that would be created by this smaller side chain is filled by the insertion of a residue from the H2 region—Pro 52a. In KOL and J539 the side chain at position 52a is on the surface. The relative movement of position 52a involves a change in the

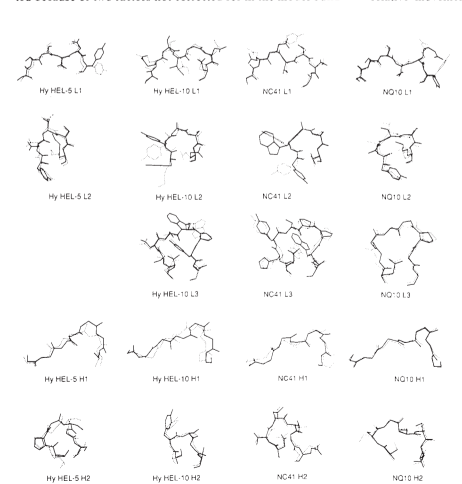


FIG. 2 The predicted (broken line) and observed (continuous line) conformations of the hypervariable regions. The structures have been superposed by a least-squares fit of their main-chain atoms. Residue numbers and the r.m.s. difference in position of the superposed atoms are given in Table 2a. Predicted and observed side-chain conformations are shown for all regions except H1 and L3 in NQ10 where they obscure the main chain. After our prediction of the NC41 structure several revisions were made to the sequence and some of the differences can be seen here.

TABLE 2 Differences in structure of predicted and observed hypervariable regions

(0)	Differences	in	local	confe	rmation	/ Å)
(a)	Differences	ın	locat	conto	irmation	(A)

Hypervariable		erence in atomic coms after optim		
region	HyHEL-5	HyHEL-10	NC41	NQ10
L1:26-32	0.8	1.1	0.7	0.4
L2:49-53	0.9	8.0	0.4	0.9
L3:90-97		0.3	0.5	0.6
H1:26-32	1.4	1.3	0.9	0.3
H2:52-56	1.1	1.0	0.7	0.4

(b) Differences in position relative to framework (Å)

Range of differences in positions of C_{α} atoms after

	superposition of frameworks residues (Å)							
	Hypervariable	Original	New structure					
Protein	region	prediction	library					
HyHEL-5	L1	2.0-3.8	0.8-2.1					
	L2	1.6-1.6	1.2-2.3					
	H1	0.8-4.1	0.7-2.1					
	H2	3.0-7.2	0.5-2.1					
HyHEL-10	L1	0.7-1.6						
	L2	0.613	_					
	L3	0.8-1.5	_					
	H1	1.3-3.5	_					
	H2	0.6-2.9	_					
NC41	L1	1.4-2.4	1.5-2.6					
	L2	1.1-1.8	1.0-2.0					
	L3	1.6-2.3	1.8-3.0					
	H1	1.3-2.0	1.0-2.1					
	H2	2.1-4.4	0.4-1.9					
NQ10	L1	0.4-2.7	_					
	L2	0.5-1.4	_					
	L3	0.6-1.5	_					
	H1	0.6-1.2	_					
	H2	0.6-0.9	_					

Superposition of a are illustrated in Fig. 2. The differences in the positions of the hypervariable regions in the original predictions and the observed structures of b are shown in Fig. 3. The predictions with a new structure library involve rebuilding the HyHEL-5 model using the refined VL J539 and the VH NC41 structures, and rebuilding the NC41 model using VL McPC603 and VH HyHEL-5; see text.

conformation of H2. It also tilts the H2 loop so that the positions of residues at the top of the loop in HyHEL-5 differ in position, relative to those in KOL and J539, by \sim 4.5 Å. In NC41, the residue at position 71 is Leu and that at position 52a is Thr, and the shift is not as large as that in HyHEL-5.

This analysis implies that the conformation and position of four residue H2 regions are determined by the packing against the VH framework and by the identity of the residue at position 71 in particular, and not by the position in the sequence of H2 of Gly (or Asn or Asp) as was believed previously⁸. Thus VH domains of HyHEL-5 and NC41 should provide better parents for each other than the other structures do, for they contain similar determinants for the position of H2. The predicted structure of HyHEL-5 using the observed structure of VH of NC41 and the predicted structure of NC41 using the observed structure of VH of HyHEL-5 were thus rebuilt. In these new predicted structures the large differences between the predicted and observed positions of H2 are not present: most residues differ by no more than 2.1 Å and none differs by more than 3.0 Å (Table 2b).

The results of this analysis were used in the prediction of the structure of the H2 region of the antibody NQ10.

Hypervariable regions of NQ10 and D1.3

The amino-acid sequence of the hypervariable regions and associated framework sites of NQ10 are given in Table 1. For five of the hypervariable regions, the size and the residue conservation at the relevant sites clearly indicate particular canonical structures, and a model of the VL-VH dimer of NQ10 was made using the procedure described above. For H2, after the

analysis described above, canonical structure 2 was expected because of the Arg at position 71 (Table 1).

Recently a crystal structure has been determined for NQ10 (S.S., P.M.A. and R.J.P., manuscript in preparation). This structure is determined to a resolution of 2.8 Å and the present residual is 21%. The tracing of the chain in the hypervariable regions is unambiguous. Figure 2 shows the predicted and observed structures of each of the hypervariable regions, superposed by a least-squares fit of their main-chain atoms. Table 2a gives the r.m.s. differences in position of the main-chain atoms. Figure 3 shows the relative positions of the predicted and observed hypervariable regions.

There is close agreement between the predicted and observed main-chain conformations of the hypervariable regions: the r.m.s. differences in position are between 0.3 and 0.9 Å (Table 2a). There is also close agreement in the relative positions of the hypervariable regions (Fig. 3 and Table 2b). No residues differ by more than 2.7 Å in position and all but two are within 1.7 Å.

Recently, the atomic structure of antibody D1.3 complexed with lysozyme has been refined²⁰. The comparison of the preliminary experimental structure with the prediction had shown differences for two hypervariable regions⁹. The L3 regions had differences associated with Pro 95 having a *cis*-peptide in the predicted structure and a *trans*-peptide in the observed. The H1 regions differed because the predicted structure had this region folded into framework, whereas in the observed it folded out into the solvent. The refinement of the experimental structure has resulted in the rebuilding of these two regions²⁰, and their conformations now agree with the original predictions.

Canonical structures in immunoglobulin 4-4-20

The atomic structure of immunoglobulin 4-4-20 has recently been determined at a resolution of 2.7 Å (ref. 19). The aminoacid sequence of the hypervariable regions and associated framework sites of 4-4-20 are given in Table 1. Four of the hypervariable regions, L2, L3, H1 and H2, have the size and the residues at specific sites that clearly indicate particular canonical structures (see Table 1). These four hypervariable regions do have the expected main-chain conformations. In Fig. 4 the four hypervariable regions, superposed on examples of the same canonical structures taken from other immunoglobulins, are shown. The r.m.s. difference in the atomic

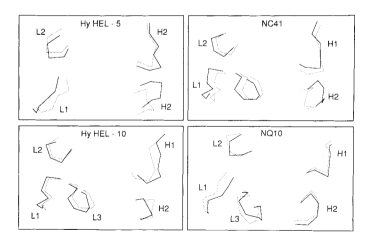


FIG. 3 The relative positions of the hypervariable regions in the predicted and observed structures. The positions of the hypervariable regions shown are those given by the superposition of the VL–VH framework regions of the observed and predicted structures. The C_α atoms of residues in the observed structure are joined by solid lines and those in the predicted structure by broken lines. The size of the differences in position are given in Table 2b.

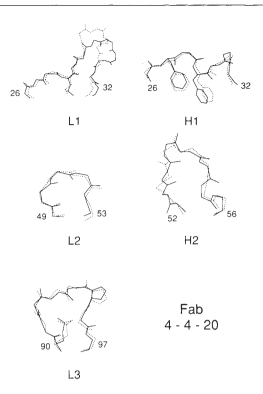


FIG. 4 The hypervariable regions in immunoglobulin 4-4-20. The observed conformations of the hypervariable regions in 4-4-20 are drawn with continuous lines. They are superposed on hypervariable regions from other immunoglobulins that have the same canonical structure-L2 and L3 from REI, H1 from KOL and H2 from McPC603-or in the case of L1, the closely related canonical structure from McPC603. The side chains shown are those that are important in determining the conformation of the region.

position of the main-chain atoms of these pairs are in the range 0.5-0.9 Å.

The L1 region of 4-4-20 has a different size to those found previously and represents a new canonical structure (see Table 1 and Fig. 4). As predicted in a previous discussion of the sequence patterns of this region⁸, its conformation is closely related to those of the other canonical structures known for this region. The main-chain atoms of residues 26-31 and 31e-32 fit the same region in McPC603 with a r.m.s. difference in position of 0.8 Å (Fig. 4).

Common occurrence of known canonical structures

The close agreement of the predicted and observed main-chain conformations of the hypervariable regions strongly supports the canonical structure model. It implies that a good estimate can be made of the extent to which hypervariable regions in other immunoglobulins have main-chain conformations close to those that are now known. This can be done by inspecting the known sequences to see whether their hypervariable regions correspond in size to the known canonical structures, and whether they contain one of the sets of residues that produce the observed conformations.

In Table 1 we give the results of examining the immunoglobulin sequences collected by Kabat et al.²⁹. About 90% of the hypervariable regions in V_K domains, and $\sim 70\%$ of the H1 and H2 regions in VH domains, are expected to have conformations close to those found in the immunoglobulin structures now known. These estimates are conservative in that they include only hypervariable regions that match the sequence requirements exactly, that is, those that contain the particular residues, or any combination of the small range of closely related residues, that are found in the known structures at the sites marked by asterisks in Table 1.

Discussion

The wide occurrence of the known canonical structures, the use of the much larger library of parent structures that will soon become available, and the more detailed understanding of the determinants of structure that will emerge from the analysis of the differences between predicted and observed structures, should allow, for many immunoglobulins, the prediction of the structure of five of the hypervariable regions with accurate local conformation and errors in position of 2 Å or less.

Predictions of the structures of the H3 regions in the immunoglobulins discussed here could not be made on the basis of the previously known structures. It was possible to predict correctly the H3 conformation in immunoglobulin D1.3 (ref. 9). In general, however, the various genetic mechanisms that produce the H3 regions usually result in medium or large surface loops with very different sequences and patterns of interaction. For such hypervariable regions it may be possible to predict their structure using the conformational search algorithms that have been developed by several groups³⁰⁻³⁵.

The results presented here have interesting implications for the molecular mechanisms involved in the generation of antibody diversity. For at least five of the six hypervariable regions of most immunoglobulins there seems to be only a small repertoire of main chain conformations, most of which are known from the set of immunoglobulin structures so far determined. Sequence variations within the hypervariable region modulate the surface that these canonical structures present to the antigens. Sequence variations within the framework and hypervariable regions shift the canonical structures relative to each other by small but significant amounts.

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