The Structural Basis of Repertoire Shift in an Immune Response to Phosphocholine

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Abstract

The immune response to phosphocholine (PC)-protein is characterized by a shift in antibody repertoire as the response progresses. This change in expressed gene combinations is accompanied by a shift in fine specificity toward the carrier, resulting in high affinity to PC-protein. The somatically mutated memory hybridoma, M3C65, possesses high affinity for PC-protein and the phenyl-hapten analogue, p-nitrophenyl phosphocholine (NPPC). Affinity measurements using related PC-phenyl analogues, including peptides of varying lengths, demonstrate that carrier determinants contribute to binding affinity and that somatic mutations alter this recognition. The crystal structure of an M3C65-NPPC complex at 2.35 Å resolution allows evaluation of the three light chain mutations that confer high-affinity binding to NPPC. Only one of the mutations involves a contact residue, whereas the other two have indirect effects on the shape of the combining site. Comparison of the M3C65 structure to that of T15, an antibody dominating the primary response, provides clear structural evidence for the role of carrier determinants in promoting repertoire shift. These two antibodies express unrelated variable region heavy and light chain genes and represent a classic example of the effect of repertoire shift on maturation of the immune response.

Key words: immunoglobulin • somatic mutation • X-ray crystallography • antibody affinity • affinity maturation

Introduction

Affinity maturation of serum antibodies occurs during the immune response to T cell-dependent antigens (1) and can be critical for protective immunity. In many antibody responses, much of the diversity exhibited by secondary and memory antibody populations is provided by somatic mutation of primary gene combinations (2-5). However, another level of diversity can also be provided by recruitment of new germline VH and VL genes during the response, termed repertoire shift (6). These new gene combinations are frequently modified by mutation, providing high affinity for the immunogen (4, 6-8). The molecular basis of this selective process is not entirely clear, but may depend in part on a shift in specificity toward recognition of the hapten plus carrier determinants.

The antibody response to phosphocholine (PC) is biologically interesting because of the frequent expression of PC on pathogenic bacteria and nematodes where the haptenic epitope is coupled to a variety of carrier structures, including carbohydrates and proteins (9-11). The murine response to PC-protein is a well-characterized model system used to study the shift in specificity to include the carrier as the immune response progresses (7, 12-15). After immunization with PC coupled to KLH via a diazophenyl linker, the antibody population specific for PC at the onset of the primary response (group I antibodies) shifts to one that includes the carrier as part of a more complex epitope. Thus the response matures, evolving toward a population that has a strong requirement for PC in the context of the nitrophenyl linker (group II antibodies). The emergent group II antibody population requires recruitment and ex-
expansion of B cell clones expressing novel V gene combinations rarely seen in the primary response (7). The expansion and eventual dominance by a minor component of the initial antibody repertoire points to the importance of carrier determinants in selecting and stimulating lymphocytes of the group II phenotype that predominate in the memory pool.

We have shown previously that somatic mutation in a prototype group II hybridoma, M 3C 65, results in a dramatic increase in affinity for antigen. High affinity for p-nitrophenyl phosphocholine (NPPC), a hapten that mimics the diazophenyl linkage between PC and protein, is attributable to mutations at three positions in CDR2 of the M 3C 65 λ1 L chain (16). Accumulation of replacement mutations at these key positions in the L chain occurs repeatedly in B cells undergoing affinity maturation in response to PC–KLH (17).

In this study, we use crystallographic and affinity analysis to reveal the molecular basis for the repertoire shift that occurs after immunization with PC–KLH. We determine the intrinsic affinity of M 3C 65 in its mutated and germline forms for compounds representing various portions of the carrier protein coupled to PC. We also solved the crystal structure of the single-chain Fv (sFv) of this antibody complexed to NPPC. The M 3C 65 combining site has an unusual requirement for direct interaction of the LCDR2 with NPPC, a finding that is rare for binding of small haptens molecules (18). The structure of the complex provides a stereochemical basis for emergence of group II dominance in a secondary response, and allows evaluation of the direct and indirect effects of L chain modification by somatic mutation in generating high affinity in the M 3C 65 hybridoma.

Materials and Methods

Synthesis and Expression of M 3C 65 sFv. cDNA copies of the VH and VL genes of the hybridoma M 3C 65 (16) were synthesized from mRNA using random hexamer primers (Amersham Pharmacia Biotech). PCR amplification was carried out using a primer for the 5' portion of Va1 (5'-agatgcatgagcatggGACA-GGGTGGTGACTCATGGAA-3') and a primer for the 3' portion of Jα1 plus the 5' part of the (Gly3Ser)1 linker (5'-cgaggaccacc-acgcccgagccaccgccaccACCTAGGACAGTCAGTTTGGT-3'). Amplification of the VH M 141/D/JH3 gene was carried out using a primer for the 3' portion of the linker plus the 5' VH (5'-tcg-gcctgctgctgctgctgcatgatcaGGTGACTCATGGAAAGAGTCAG-3') and a primer for the 3' end of JH3 plus two stop codons and an N col site (5'-agatgcatgagcatggGACA-GGGTGGTGACTCATGGAAAGAGTCAG-3'). N col sites are underlined, and region coding sequences are capitalized. The VL and VH genes were joined by PCR with the linker (Gly3Ser)1 in the order VL-linker-VH. The amplified sFv DNA was ligated into the N col site of the pET3d vector (Novagen), sequenced, and transformed into Esherichia coli B121(DE3). DNA sequencing of clone 2-1.2 showed that the deduced protein sequence was identical to the M 3C 65 hybridoma. The induced protein sFv (24.9 kD) were denatured and refolded as described (19). R enatured sFv fragments were purified by affinity chromatography on PC-sepharose as described for intact antibody (12).

Transfected Antibodies and Kd Determination. Antibodies expressing mutations at λ1 L chain positions 52, 53, or 55 produced by site-directed mutagenesis and the method for determination of binding constants by fluorescence quenching have been described (16). Stable transfecants cotransfected with the M 3C 65 H chain and the various mutant L chain constructs were made in the SP2/0 cell line and purified as described previously (16). The synthesis and structure of monoconjugates of PC coupled to tyrosine, histidine, and the Gly-Tyr-Ala tripeptide have been described; PC is coupled to Tyr on the tripeptide (20). These compounds were provided by Drs. D. Peyton, E. Barbar, and H. Moulton (Portland State University, Portland, O R.). A 15- amino acid peptide, acetylated at the NH2 terminus (Peptide Express), was designed to contain one haptenation site and to have α-helical secondary structure (21). The peptide, Ser-Asp-Ala-Leu-Ala-Glu-M et-Tyr-Glu-Leu-M et-Ala-Val-Asp-Gly, was coupled to PC at the Tyr residue as described previously (22). PC-histone was coupled by the same method and has a PC/protein ratio of 2:1.

Crystallization of M 3C 65 sFv Complexed to NPPC. The purified sFv at 400 μg/ml was dialyzed against 20 mM Tris, pH 7.6, concentrated to 5.6 mg/ml, and NPPC was added to a final concentration of 2 mM. Crystals were grown at room temperature by vapor diffusion from a reservoir buffer that contained 1.5 M NaH2PO4/K2HPO4, pH 6.5, and 0.1 M Hepes buffer (GIBCO BRL). The crystals are orthorhombic, take the space group P212121, (a = 130.5 Å, b = 35.9 Å, c = 50.5 Å), and contain one monomer in the asymmetric unit.

Data Collection and Structure Solution. X-ray intensity data were collected at room temperature using a R AXIS IV (R igaku) imaging plate system and a Rigaku RU 300 rotating anode X-ray generator equipped with double-focusing mirrors and operating at 50 kV and 100 mA. Data were processed with B IOTEX (M olecular Structure Corporation, Inc.). The structure was solved by molecular replacement (M R ) using data from 8.0- to 4.0-Å resolution and the λ1 L chain from antibody HC19 (23), 1G1G, as the search model. The MR was carried out using EPM R (24), which resulted in a solution with a correlation coefficient of 0.372. Search models using structures of several antibody H chains failed to produce a solution. Rigid body refinement of the L chain using T N T (25), using data from 10.0 to 3.0 Å, reduced the R factor to 44.6%. Phases from this partial structure were used to calculate an electron density map that revealed some interpretable density for the H chain. The D1.3 H chain (26), 1VFA, was then manually rotated into the density, and rigid body refinement was carried out. This reduced the R factor to 35.6%, at which point the correct side chains were substituted in the H chain. Positional (xyz) refinement, using data from 10.0 to 2.8 Å, was then initiated, and reduced the R factor to 22.9%. The resulting electron density map revealed very clear density for the NPPC hapten, which was then included, and the resulting model was refined via xyz, and subsequently via positional and thermal parameter (xyz) refinement. Xyz refinement was carried out initially using data from 10.0- to 2.6-Å resolution. The final refinement included data extending from 10.0- to 2.35-Å resolution. The current structure includes residues 1-109 of the L chain, 1-112 of the H chain, the NPPC molecule, and 88 solvent molecules. P R O C H E C K analysis (27) revealed 97.9% of residues in allowed regions (78% in most favored), and 2.1% in generously allowed regions. See Table II for selected crystallographic refinement statistics. Coordinates have been deposited in the Protein D a tabase Bank with accession code 1DL7.
Results

Affinity of M3C65 Wild-Type and Mutant Antibodies.

The M3C65 hybridoma antibody (y2b, l1) is a member of a set of clonally related antibodies produced in a secondary response to PC–protein. High affinity for the hapten analogue NPPC is attributable to mutations at positions 52, 53, and 55 in CDR2 of the l1 L chain (16). Mutations at these key positions occur repeatedly in M3C65 clonal relatives, as well as in unrelated l-beariing anti–PC protein hybridomas, indicating that LCDR2 replacements are vital to expansion and selection of these memory antibodies (16, 17). To assess whether the hapten–carrier linkage beyond the diazophenyl moiety forms part of the immunogenic structure of PC–KLH, we have analyzed binding to a number of structures representing distinct carrier epitopes. We also determined whether these structures are recognized differently as mutations accumulate. A comparison was made of the affinity of antibody constructs expressing zero (germline), one, or two mutations with the binding of M3C65, which contains all three pertinent mutations in LCDR2. The sequences of the H and L chain V regions and the positions of mutations are indicated in Fig. 1. The PC ligands used to measure affinity by fluorescence quenching are shown in Fig. 2.

Diazophenyl links preferentially to tyrosine or histidine residues in proteins (28). Because both KLH isoforms contain multiple tyrosine and histidine residues (29), both amino acids are likely to be haptenated during conjugation of PC to KLH via a diazophenyl linkage. To analyze the contribution of a single amino acid linked via diazophenyl to PC, we compared the affinity for PC–tyrosine or PC–histidine to the affinity for NPPC in antibodies expressing germline or mutated L chains. As shown in Table I, the relative affinity for ligand of antibody expressing the unmutated L chain is PC–histidine > PC–tyrosine > NPPC, with the affinity for PC–histidine about ten times that of the affinity for NPPC. Although the affinity for NPPC increased as mutations accumulated in the L chain, all the mutant antibodies had higher affinity for PC–tyrosine and PC–histidine compared with antibody expressing germline L chain. It is noteworthy that ligand recognition shifts toward PC–tyrosine with the accumulation of mutations in the M3C65 antibody. Thus, binding to PC–tyrosine or PC–histidine allows for detection of changes in fine specificity, with the affinity of the mutated M3C65 antibody for PC–tyrosine being fivefold greater than for PC–histidine. These results demonstrate that carrier determinants, as represented by tyrosine and histidine, contribute to binding, and that somatic mutation can modulate carrier recognition.

How much of the PC–protein carrier provides determinants recognized by group II antibodies? Previous experiments involving inhibition of binding of M3C65 to PC–histidine.
The results in Table I show that affinity of antibody expressed with PC (discussed in Materials and Methods), whereas the (and no histidine) in the sequence, and are monosubstituted and the 15–amino acid peptide each have only one tyrosine for PC coupled to carriers of varying sizes. The tripeptide questions, we determined the intrinsic affinity of antibody with an increase in size of the carrier? To answer these

Is the superior inhibitory capacity of PC–protein due solely to the effects of avidity, or does effective binding correlate with the effects of avidity, or does effective binding correlate to the effects of avidity, or does effective binding correlate with an increase in size of the carrier? To answer these questions, we determined the intrinsic affinity of antibody for PC coupled to carriers of varying sizes. The tripeptide and the 15–amino acid peptide each have only one tyrosine (and no histidine) in the sequence, and are monosubstituted with PC (discussed in Materials and Methods), whereas the histones have multiple tyrosine and histidine residues (30). The results in Table I show that affinity of antibody expressing germline γ1 for the PC–GY A tripeptide is higher than for PC–tyrosine, PC–histidine, or PC–histone. These findings indicate that neither size nor avidity alone provides the advantage, but that qualitative characteristics of each individual carrier can be critical. After the introduction of the single mutation at position 53, binding to the monosubstituted 15–amino acid PC–peptide and to PC–histone is at levels near or equal to that of the fully mutated antibody. Thus, it appears that an increase in size of the carrier would benefit binding to germline-encoded antibodies up to a certain point (PC–GY A > PC–histidine > NPPC), but a further increase in size represented by the protein conjugate, PC–histone, confers little affinity advantage until mutation occurs.

The cumulative effects of somatic mutation on binding of small haptens in several systems indicate that high affinity can result from incremental improvements as mutations accumulate (31, 32). In M 3C65, acquisition of high affinity for NPPC, PC–tyrosine, or PC–histidine by introduction of L chain mutations follows a similar pattern (Table I). In contrast, introduction of a single mutation at LCDR 2 position 53 accounts for high-affinity binding of the monosubstituted 15–amino acid PC–peptide and results in a 200–fold increase in affinity for PC–histone. Thus, binding to the haptenic compounds mimics the stepwise acquisition of high affinity for small haptens seen with other antibodies, whereas high affinity to the larger PC–carrier structures may result from a single substitution. Production of a high-affinity antibody by a single mutation emphasizes the adaptive potential of the group II combining site (33). The surprising difference in the effect of mutation on binding to the various PC–containing compounds raises a cautionary note concerning the extrapolation of antihapten binding results to the in vivo situation, where the complete hapten conjugate may be the structure relevant to shaping of the immune repertoire. In addition, as demonstrated here and by others, somatic mutation not only increases affinity for hapten, but may also shift antigenic specificity (34–36).

Structure Determination and Overall Structure of the M 3C65 sFv–NPPC Complex. To determine the molecular basis of PC–carrier recognition and the contribution of somatic mutations, the crystal structure of an M 3C65 sFv fragment complexed with NPPC was determined at a resolution of 2.35 Å (atom labeling of NPPC shown in Fig. 2 A). The sFv fragment of M 3C65 was used for crystallographic analysis because the sFv binding data were in good agreement with that of intact antibody (Table I). Attempts to analyze the sFv fragment bound to the higher-affinity ligand, PC–tyrosine, were unsuccessful because the complex did not yield data-quality crystals. The crystallographic data collection and refinement statistics for the sFv–NPPC complex are shown in Table I.

The combining site of M 3C65 is a long narrow groove, ~15 Å long by 5 Å wide by 7 Å deep, created by residues from LCDR 1, LCDR 2, LCDR 3, HCDR 2, and HCDR 3 (Fig. 3). The binding pocket is unusual in that it is enclosed at one end by LCDR 2 residue Thr55, which seals off the p-nitrophenyl group of the hapten from solvent. Contact with LCDR 2 is rare in antibodies recognizing small haptens (18), but has been seen in the anti-DNP antibody.
ANO2, which has a similarly shaped combining site (37).

In addition to the steric complementarity of the binding pocket to the hapten, electrostatic components appear to be crucial for selective binding and orientation. Specifically, at one end of the combining site, a negatively charged patch created by H chain CDR3 residue Asp95 attracts and orients the positively charged trimethylammonium portion of the choline moiety of NPPC (Fig. 3).

Antibody–Hapten Interactions. Shape complementarity and electrostatic contributions anchor NPPC in the binding site, allowing for optimal interactions with both H and L chain residues. The absence of water molecules around the ligand contributes to the tight fit in the binding site, and calculations using GRASP (38) reveal that NPPC binding buries 644 Å$^2$ of combining site surface area from solvent. The contacts from the H chain are provided by just two residues, HCDR2 residue Trp52 and HCDR3 residue Tyr96 (Table III; Fig. 4). Notably, Tyr96 also forms part of a “tyrosine sandwich” involving both itself and LCDR1 residue Tyr32 (Figs. 3 and 4). These tyrosines play a critical role in binding by effectively encapsulating the central part of the hapten. HCDR2 residues Trp52 and Met50 combine with residues from LCDR3 (Trp91 and Trp96) to form a hydrophobic well at the base of the binding site that interacts with the choline moiety.

Most of the contacts with NPPC involve hypervariable regions of the M3C65 L chain (Fig. 4; Table III). Asn34 from LCDR1 hydrogen bonds to O1 of the phosphate group of the hapten, whereas Tyr32, the second half of the “tyrosine sandwich,” enforces the phosphate group along with H chain Tyr96. LCDR3 residues Trp91 and Trp96 interact with the choline moiety, completing the aromatic, hydrophobic well at the base of the hapten docking site. There is also a cation–π interaction between the positively charged nitrogen of the choline moiety and the indole ring of Trp96. Cation–π complexes have been shown to be important in other biological systems, including a group I PC-binding antibody M603, where these interactions with aromatic amino acid side chains enhance recognition of choline-containing ligands (39).

LCDR2 is the site of the three mutations at positions 52, 53, and 55 that confer increased binding affinity, and the crystal structure reveals the key roles of His53 and Thr55 in the formation of the combining site. His53 forms two hydrogen bonds, via the side chain Nδ1, to the carbonyls of L chain Gly49 (Nδ1-O, 3.1 Å) and Gly50 (Nδ1-O, 3.1 Å). These hydrogen bonds stabilize the combining site and anchor the side chain of His53 in a position to stack against and interact with NPPC. The side chain of His53 directly

**Table II. Data Collection and Refinement Statistics**

| Description                              | Value          |
|------------------------------------------|----------------|
| Resolution range (Å)                     | 10.0–2.35      |
| No. of reflections                       | 8,983          |
| No. of measurements                      | 47,930         |
| Completeness (%)                         | 84             |
| I/σ(I)                                   | 8.5            |
| R sym*                                   | 7.3            |
| Data for highest shell (2.42 to 2.35 Å)   |                |
| Completeness (%)                         | 47             |
| I/σ(I)                                   | 1.5            |
| R sym*                                   | 26.5           |
| R factor (%)                             | 19.1           |
| R free (%)                               | 26.5           |
| No. of atoms                             | 1,680          |
| No. of solvent molecules                 | 88             |
| Rms deviations                           |                |
| Bond lengths (Å)                         | 0.014          |
| Bond angles (°)                          | 1.901          |
| B factors (Å$^2$)                        | 3.1            |

*R sym = ΣI_{o} / ΣI_{a}, where I_{o} = observed intensity and ΣI_{a} = average intensity obtained from multiple observations of symmetry-related reflections.

*R factor = Σ|F_{o} - F_{c}| / ΣF_{o}; The lower resolution limit of refinement for all structures was 10.0 Å.

*R free = Σ|F_{o} - F_{c}| / ΣF_{o}, where all reflections belong to a test set of 10% randomly selected data.

**Figure 3.** The electrostatic surface potential of the M3C65 Fv. Negative and positive potentials are red and blue, respectively, and neutral areas are white. The NPPC hapten is colored by atom type with blue nitrogen, yellow phosphorus, red oxygen, and white carbon. Positions of the three CDR2 L chain mutations (K52, H53, and T55) and other amino acids important in binding NPPC are indicated. The antibody chain designation, H or L, is in parentheses. The figure was generated using GRASP (38).
contacts the p-nitrophenyl group of NPPC (Table III), and also stacks against its aromatic group. The critical role played by His53 corroborates the finding that CDR2 position 53 is a specificity-determining residue in L chains (40).

Residue Thr55 plays two key roles in the architecture of the binding site near the p-nitrophenyl group. First, this b-branched residue closes off the end of the binding groove, allowing a tighter fit with the ligand by exclusion of water from the site, which is exposed in antibodies using the germline L chain. Second, Thr55 engages in a hydrogen bond from the Oe atom to the carbonyl oxygen of H chain Gly97. This interaction locks the L and H chains together in the specific conformation required for tight binding, and is therefore critical. Notably, this interaction is made possible by the presence of a proline at H chain position 98, which has adopted a cis conformation.

The role of mutated residue Lys52 is less clear, as the side chain is oriented away from the binding pocket. However, comparison of the M 3C65 structure to the germline structure suggests that the importance of this mutation may lie in the replacement of the germline asparagine residue. In the germline \( \lambda \)1 structure, Asn52 hydrogen bonds to Asn53. Such a hydrogen bond would also be possible between germline Asn52 and the mutated M 3C65 His53 side chain, a contact that would disrupt the crucial contacts provided by His53. In contrast, a lysine at position 52 would be unlikely to hydrogen bond to His53, allowing His53 to make critical architectural contacts necessary for optimal binding of NPPC. Mutation away from asparagine may be aided by the high mutability index of the germline codon sequence (41).

The structural analyses of several other antibody–hapten interactions indicate that amino acid replacements resulting from somatic mutation may directly affect contact residues, but more frequently elicit their effects indirectly via conformational changes in the binding site (42–46). However, in M 3C65, both effects are seen. The His53 and Thr55 replacements, along with the indirect effects of the Lys52 replacement, optimize and fix the shape of the combining cavity, whereas His53 also contacts the hapten directly.

Comparison of M 3C65 \( \lambda \)1 Structure with Germline \( \lambda \)1. To assess structural changes in combining site shape resulting from somatic mutation, we compared the M 3C65 antibody with that of an unmutated \( \lambda \)1 structure. Superpo-
tion of hybridoma HC19 germline λ1 (23) with the mutated M3C65 sFv shows that the Cα backbone tracings are quite similar, with a root mean square deviation of 0.50 for 108 corresponding residues (Fig. 5). Strikingly, the only notable differences between the two structures are observed in LCDR2 and involve residues 52–56, the region containing the mutated residues, with the major shifts occurring in residues Thr55 and Pro56. As noted, this significant shift results from contacts made by the mutated LCDR2 residues His53 and Thr55, thus providing the necessary conformation of the hapten binding pocket. The Cα backbone tracings of germline λ1 and M3C65 λ1 are very similar. However, one notable shift in amino acid side chain placement occurs in the binding pocket, where L chain residue Tyr32 has rotated ~90° from the germline, orienting the ring face so that it is more perpendicular to the ligand. This rotation of Tyr32 also places the Cδ of Tyr32 closer to the O4 of the ligand phosphate group, a shift from 4.0 to 2.8 Å, such that its positive edge complements the negatively charged phosphate group. This close approach is indicative of an unusually strong C-H···O hydrogen bond (47).

**Discussion**

A shift in antibody repertoire occurs after immunization with some hapten-protein conjugates, reflecting recruitment of alternative gene combinations at different stages of the immune response (6). The potential to shift the antibody repertoire could play an important role in determining the ultimate success of immunization, but the molecular forces driving this complex process are not clear. These forces are likely to be multifaceted and may include (a) the precursor frequency of cells expressing a given VH/VL gene combination (48), (b) intrinsic affinity and/ or binding kinetics of the immunoglobulin B cell receptor for ligand (3, 6), (c) adaptability of the immunoglobulin receptor to somatic mutation (8), (d) restrictions in entry into the memory cell compartment because of differences in B cell lineage (49), and (e) idiotypic interactions that may down-regulate dominant clonotypes and promote polyclonality of the developing memory B cell population (6). In addition, as discussed here, competitive advantages may lie in the use of alternative V’ gene combinations that allow accommodation of carrier structures, resulting in alteration of initial patterns of clonal dominance.

**Repertoire Shift in the Immune Response to PC-KLH.** Immunization of BALB/c mice with PC-KLH leads to expression of two antibody populations differing in fine specificity that predominate in the immune response at different times (7, 12–15). This repertoire shift occurs in the late primary and early secondary response, resulting in loss of initial dominance by antibodies bearing the T15 idiotype (group I) and their replacement by group II antibodies that employ different VH and VL combinations. We propose that the new gene combinations employed by group II antibodies allow recognition of complex PC-carrier epitope(s), and that the ability of these combining sites to be shaped by favorable somatic mutations allows emergence and dominance by group II-specific B cells.

T15 antibodies are restricted to use of the VH1/Vλ22 gene combination. These antibodies undergo little affinity matura-

Figure 5. Overlay of the M3C65 V region (blue) with the germline λ1 of antibody HC19 (yellow). The orientation of L chain residue Tyr32 is shown for germline λ1 and M3C65. The positions of mutated residues 52, 53, and 55 are indicated. The NPPC hapten (pink) is shown complexed to M3C65. The figure was generated using SYBYL (Tripos, Inc.).
mutations (56, 57). Thus, T15 may be unique in the failure of mutations to improve antigen binding (33).

Group II antibodies use a variety of H and L combinations unrelated to T15 (7). Although the κ L chain is expressed in many memory anti-PC–KLH antibodies, λ-bearing antibodies can dominate (>75%) some responses. To clarify factors conferring selective advantage to these antibodies, we focused on the antigen binding properties and combining site structure of a prototype group II memory antibody, M 3C65. We demonstrate that (a) carrier determinants extending from PC contribute to the affinity of M 3C65, (b) the shape of the M 3C65 combining site is well suited for accommodation of PC in the context of carrier residues, (c) somatic mutation can alter the specificity for carrier determinants, and (d) somatic mutation can increase affinity directly by affecting residues in the combining site that contact ligand and indirectly by modifying the shape of the combining site.

More than 35 years ago, it was suggested that antibodies elicited to conjugated haptens might recognize carrier residues in addition to the haptenic determinant (58). Defining the complete antigenic determinant has been difficult, but in many systems it is thought that the protein carriers of haptens make a significant contribution to binding by anti-hapten antibodies (37, 44, 59). However, T15 has approximately equal intrinsic affinities for PC ($K_a = 2.9 \times 10^7 \text{M}^{-1}$) and NPPC ($K_a = 1.8 \times 10^8 \text{M}^{-1}$), indicating the absence of a requirement for the diazophenyl linkage between PC and the protein carrier (60, 61). These results are in agreement with the earlier conclusion that T15 antibodies recognize only the PC portion of DPPC, another hapten that mimics the linkage between PC and protein (13). Thus, the carrier linkage does not appear to be an essential part of the epitope, in contrast to group II antibodies, where the carrier linkage is an essential feature.

Comparison of Group I and Group II Combining Sites. Inspection of the binding pockets of two group I antibodies, T15 (43, 62) and the related antibody M 603 (63, 64), reveals structural differences that would allow better accommodation of carrier residues by group II antibodies. Fig. 6 shows comparative models of T15 bound to PC and M 3C65 bound to NPPC. In the T15 combining site, PC is bound in a deep pocket anchored by contacts between the antibody and the phosphate group. The cavity of T15 is deeper than M 3C65 as a result of longer LCDR1, HCDR2, and HCDR3 loops that protrude farther from the binding pocket. PC occupies only a small portion of the combining site, leaving room for carrier residues; however, the deeper wedge-shaped cavity may hinder accommodation of larger carrier structures. Because of the orientation of PC toward the H chain, the HCDR2 region of T15 would likely interact with carrier epitopes, but the extended length and protrusion of CDR2 may also restrict the carrier interactions by steric hindrance. Although recognition of carrier epitopes can be modulated by mutation of the T15 HCDR2 loop, the effect on binding is subtle (65). The shape of the M 3C65 binding site appears to be more suited for binding PC-associated carrier residues than that of T15, and selected mutations may favor such interactions. The elongated shallow groove of the M 3C65 combining site provides a flatter surface more akin to antibodies recognizing proteins than the deep pockets of many antibodies recognizing small antigens (18, 46, 66). Extension of carrier residues would continue over the surface of the antibody in such a way as to offer potential interaction with other residues of LCDR2. The more ideal shape of the M 3C65 binding site for accommodating carrier compared with T15 is reflected in the 1,000-fold better binding of PC–protein (7, 16). Thus, the basic geometry of the combining sites differs between these two primary and secondary anti-PC antibodies, unlike the response to phenyloxazolone, where it has been postulated that crucial contact residues would maintain the same mode of binding after recruitment of a new VH (67).

Biological Consequences of Repertoire Shift and Somatic Mutation. Repertoire shift is commonly observed during immune responses to hapten conjugates, as well as to viral and other protein antigens (55, 68–71). Selective pressure on B cells imposed by affinity for antigen is probably one key factor influencing the changing antibody repertoire (6). In the response to PC–protein, recruitment of new gene combinations produces a corresponding shift away from recognition of free PC toward higher-affinity recognition of PC–protein. The affinity of M 3C65 for PC–protein ($K_a = \ldots$)
1.2 \times 10^8 \text{ M}^{-1}

is in the range for affinities to protein antigens (K_a = 10^8-10^{10} \text{ M}^{-1}) (70, 72, 73), as predicted by Foote and Eisen (74), and as found for a catalytic antibody binding to a hapten–peptide conjugate (46). Moreover, the affinity of the mutated M 3C65 for PC-protein is well above the limit for cellular stimulation (K_a = 10^5 \text{ M}^{-1}) determined for a monovalent antigen (75). M 3C65 expressing germline L chain (K_a = 0.5 \times 10^6 \text{ M}^{-1}) is just below this threshold. Presumably, antigen excess and the multivalent nature of PC–KLH allow the repeated selection of this threshold. Antigen excess and the multivalent nature of PC–KLH is likely to lead to repertoire shift seen in the immune response to PC–KLH. The role of somatic mutation in optimizing binding site geometry during affinity maturation (3, 76) and charge characteristics imparted to antibodies by new VH/VL gene combinations provide a mechanism for the accommodation of carrier residues.

The crystal structure of the M 3C65 sFv has allowed us to correlate binding with structural features of the antibody that confer advantage to mutated antibodies forming the group II memory pool. The role of somatic mutation in optimizing binding site geometry during affinity maturation has been studied in previous crystallographic analyses of antibody–hapten complexes (44, 67, 78, 79). However, the structural basis of repertoire shift has remained more elusive because of the lack of crystal structures representing different stages of the maturing antibody response. Comparison of T 15 and M 3C65 binding sites illustrates how the accommodation of carrier residues provides a mechanism for the repertoire shift seen in the immune response to PC–KLH.

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It has been postulated that repertoire shift could be driven by kinetic parameters, such that the hydrophobic and charge characteristics imparted to antibodies by new VH/VL gene combinations provide high association rates leading to more efficient long-range attraction of antigen (3, 76). In addition, a combining site that is more accessible to antigen can lead to faster binding kinetics (48, 77). It will lead to more efficient long-range attraction of antigen (3, 76). In addition, a combining site that is more accessible to antigen can lead to faster binding kinetics (48, 77). It will lead to more efficient long-range attraction of antigen (3, 76). In addition, a combining site that is more accessible to antigen can lead to faster binding kinetics (48, 77).

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