Paratope Plasticity in Diverse Modes Facilitates Molecular Mimicry in Antibody Response

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Molecular mimicry is the antithesis to the specificity of recognition processes as exemplified by the immune system. The hallmark of the acquired immune system is its ability to distinguish between distinct molecules. Specificity at the molecular level enables immune responses to be mounted against the invading pathogens while ensuring self-nonself discrimination. Molecular mimicry, in essence, is the breakdown of the specificity of recognition of humoral and cellular responses. The physiological equivalence of disparate molecular entities has substantive implications in immunological processes that confront the repertoire of antigenic diversity. Although molecular mimicry has been the central premise in the etiology and pathogenesis of self-reactive Abs and T cells (1, 2), it also provides an archetypal model for mechanistic understanding of the specificity and complexity of immune recognition. Exploration of diverse facets of molecular mimicry would provide insights into the genesis of autoimmunity and aid vaccine design strategies (3).

We have addressed the molecular basis of mimicry by analyzing chemically dissimilar ligands which show equivalence to an impressive degree (4–12). Con A, a mannose-specific lectin, has been shown to recognize Tyr-Pro-Tyr motif-containing peptides (13, 14). However, in the crystal structures of the complexes of a number of such peptides with the lectin, well-defined structural similarity could not be deciphered as the carbohydrate and peptide ligands did not bind at a common site (5, 6, 8, 9). Similarly, functional mimicry between the 12-mer (DVFPYPYPASGS) and methyl-α-d-mannopyranoside was demonstrated in polyclonal responses in terms of cross-reactivity wherein immunization with the peptide gave rise to carbohydrate-recognizing Abs and vice versa (4). In addition, during immune maturation, a booster with the peptide Ag could also enhance the anti-mannopyranoside Ab response, thus establishing equivalence between the peptide and carbohydrate moieties (7). However, precise molecular description of the functional mimicry, as seen by the immune system, between these otherwise chemically independent Ags remained elusive.

Previous investigations had suggested that the ability to recognize mimicking Ags arose from the conformational flexibility in the Ab paratope through adaptation of structure appropriate for complementing the Ag (12). The present study addresses the physicochemical basis of mimicry in humoral responses against a peptide and a carbohydrate at the monoclonal level. Molecular mimicry between 12-mer and mannanpyranoside was analyzed in terms of recognition specificity and affinity of interaction with the mimicking Ags. Quantification of the changes that occur upon binding in terms of kinetic and thermodynamic parameters and the effect of temperature on these would shed light on the modes of binding of diverse yet mimicking Ags. Furthermore, a systematic understanding of the energetics of Ag binding and its correlation with the structural features of the combining site of the Ab would facilitate better understanding of the phenomenon of molecular mimicry in the humoral response. Our studies highlight the role of Ab paratope in the generation of diverse mimotope-recognition modes, revealing a new facet of the evolving paradigm in molecular mimicry. Modulation of mimicry by Ab-dependent properties suggests mechanisms by which the host can evolve strategies for minimizing the consequences of antigenic variation by invading pathogens.
Materials and Methods
Preparation of Ags and generation of hybridomas
The 12-mer (DVFYPYPYASGS) peptide was synthesized and conjugated to BSA and diphtheria toxoid (DT)\(^1\) as reported previously (12). Mannopyranoside-BSA was prepared after activation of p-aminophenyl-\(\alpha\)-mannopyranoside (Sigma-Aldrich) with an equimolar amount of glutaraldehyde in 0.1 M sodium carbonate buffer (pH 9.0), for 30 min at room temperature, and then mixing with BSA. The reaction mixture was incubated at 4°C overnight, after which it was extensively dialyzed against normal saline.

Previously standardized protocols for immunizations were followed wherein each mouse was injected i.p. with 200 \(\mu\)g of conjugated 12-mer-DT emulsified with CFA (Difco) and given a booster of the Ag with IFA (Difco) on day 21. The highest responder mouse was sacrificed for Mannopyranoside-BSA was prepared after activation of goat anti-mouse IgG (Jackson ImmunoResearch Laboratories) was used as the reference system (Amersham Biosciences). Both 12-mer-BSA and mannopyranoside and 12-mer. The level of inhibition was calculated by comparison of binding signal of the Ab incubated with different ligand concentrations with respect to signal of Ab containing no soluble ligand and expressed as a percentage.

Ab purification
The Ab was purified from the ascitic fluid of mice by an initial partial purification by salt fractionation with a 40% ammonium sulfate cut-off. The precipitated Ab was then solubilized and dialyzed in 10 mM Tris-Cl (pH 8.5), and subjected to anion exchange chromatography on a DEAE column. The purity of the Ab was checked by SDS-PAGE and concentration was estimated by protein assay (Bio-Rad) using BSA as the standard.

ELISAs
The binding of the Abs, from cell supernatant or in purified form, to the peptide and carbohydrate ligands was assayed by sandwich ELISA. 12-mer-BSA or mannopyranoside-BSA was used as the coating Ag at a concentration of 2 \(\mu\)g/well on 96-well immunosorbent plates and Ab was added at appropriate dilutions after blocking with 1% BSA. HRP-labeled goat anti-mouse IgG (Jackson ImmunoResearch Laboratories) was used as the secondary Ab and \(\alpha\)-phenylendiamine (Sigma-Aldrich) and \(H_2O_2\) as peroxidase substrates. Absorbance was recorded at 490 nm after addition of 1 M ethanolamine. The reference surface was treated in the same way except that no ligand was passed over this surface to normalize for the H chain.

Ab sequencing
Approximately 10\(^6\) to 10\(^7\) hybridoma cells were used for total RNA extraction with TRIzol reagent (Invitrogen Life Technologies). Protocols followed for synthesis of the first strand of cDNA were identical with that reported earlier (12) except that a chain 3' primer (Mouse Ig-Primer set; Novagen) was used in the case of L chain of mAb 7C4 and for H chain, 5'-GGCCAGTGGTATAGAC-3' primer was used as the 3' primer. Subsequent amplification of the single-stranded cDNA by PCR was conducted using the 5' and 3' primers (Mouse Ig-Primer set; Novagen) for the L chain and 5'-AGGTG(C/G)(A/C/A)(A/G)CTGCAG(C/G)AGTCT(A/T)GGG-3' as the 5' primer and 5'-GGCCATGGTATAGAC(T/C/A)GA-3' as the 3' primer for the H chain. A total of 3 \(\mu\)l of the PCR product was analyzed on 1% agarose gel. Subsequently, the PCR products were sequenced using their respective forward and reverse primers.

Molecular modeling
Homology modeling of proteins depends on the identification of a protein of known structure that shares sequence, properties, function, and/or evolutionary classification with the protein of interest. The search for structures with sequence similarities to our Ab was performed using the basic local alignment search tool (15) in the Protein Data Bank (PDB). As we were primarily studying the Ag-binding site, only the V regions of the Ab have been modeled. To optimize the juxtaposition of the \(V_L\) and \(V_H\), we selected the template for modeling based on highest homology with both chains of the Ab rather than selecting two different Abs that show higher identity, individually, with each of the chains.

The structure of the Fab of an anti-hapten catalytic Ab 34E4 (PDB ID: 1Y0L) that catalyzes the conversion of benzoxazoles to salicylnitriles was used as the template for homology based modeling of the mAb 7C4 Fv. Sequence alignment with 1Y0L showed a two-residue gap in CDR H3. Amber (12) except that a chain 3' primer (Mouse Ig-Primer set; Novagen) for the L chain and 5'-AGGTG(C/G)(A/C/A)(A/G)CTGCAG(C/G)AGTCT(A/T)GGG-3' as the 5' primer and 5'-GGCCATGGTATAGAC(T/C/A)GA-3' as the 3' primer for the H chain. A total of 3 \(\mu\)l of the PCR product was analyzed on 1% agarose gel. Subsequently, the PCR products were sequenced using their respective forward and reverse primers.

Synthesis and assay of the peptide analogs on solid surface
Sequentially mutating each residue of the 12-mer peptide to glycine and measuring the binding of the individual peptide analogs to the Ab would allow delineation of the Ab-specific epitope of the 12-mer. The 12-mer peptide analogs were synthesized on the surface of polystyrene pins.

\(^1\)Abbreviations used in this paper: DT, diphtheria toxoid; RU, resonance unit.
using F-moc chemistry as described previously (12). After completion of the synthesis, all peptide analogs were acetylated at the N terminus and side chains were deprotected. The pins with their irreversibly bound peptide analogs could be reused by removing the bound Ab by sonication in the recommended disruption buffer. Evaluation of Ab binding to the analogs was carried using an ELISA-based assay wherein 2% gelatin and side chains were deprotected. The pins with their irreversibly bound peptide-carbohydrate mimicry. The Ab was purified from ascites by 40% ammonium sulfate precipitation before ion exchange chromatography.

The purified Ab 7C4 showed equivalent binding to the immunizing 12-mer and its mannopyranoside mimic (Fig. 1A). It was therefore pertinent to analyze whether these mimicking Ags could displace each other during binding to the Ab. Owing to the size and nature of the Ags, BSA-conjugated forms of 12-mer and mannopyranoside were made to compete with their immobilized forms for binding to mAb 7C4 (Fig. 1B). It was found that both the peptide and the carbohydrate Ag can compete with one another suggesting that the Ab is capable of recognizing the mimicking ligands equivalently and with comparable affinities.

Toward understanding the mechanistic details of the Ab binding to the Ags, the kinetics of the interactions with the immunogen and its mimic were measured. Measurement of the affinity of mAb 7C4 for the 12-mer and mannopyranoside was conducted on the surface plasmon resonance-based biosensor, Biacore2000, at ambient temperature (25°C). Various kinetic parameters including association ($k_a$) and dissociation ($k_d$) rate constants of binding of the Ab with the two Ags were calculated (Table I).

Real-time binding measurements show that mAb 7C4 has similar affinities for 12-mer and mannopyranoside with an equilibrium dissociation constant ($K_d$) of 94.5 and 135.8 nM, respectively. The $k_a$ and $k_d$ for binding to both the ligands are also comparable. The kinetic parameters quantitatively reinforce the observation that mAb 7C4 not only binds to the immunizing peptide but also shows cross-reactivity to mannopyranoside with equivalent affinity. The modes of binding of the two Ags appear to be similar as the association and dissociation rate constants are comparable at ambient temperature.

### Influence of temperature on kinetics of Ab-Ag interaction

Measurement of $k_a$ and $k_d$, as a function of temperature, enables evaluation of the thermodynamics and energetics of binding of the Ab to these chemically dissimilar yet mimicking Ags. Correlation of these physical parameters with structural features of the paratope would help to delineate the modes and mechanisms of interaction of Ab with 12-mer and mannopyranoside. Sensograms were generated for the binding of the Ab to the two Ags and the data

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>$k_a \times 10^4$ (M$^{-1}$s$^{-1}$)</th>
<th>$k_d \times 10^3$ (s$^{-1}$)</th>
<th>$K_d$ (nM)</th>
<th>$t_{1/2}$ (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>6.02 ± 0.07</td>
<td>1.30 ± 0.02</td>
<td>53.1</td>
<td>21.3</td>
</tr>
<tr>
<td>20</td>
<td>5.98 ± 0.11</td>
<td>3.84 ± 0.09</td>
<td>64.2</td>
<td>180.4</td>
</tr>
<tr>
<td>25</td>
<td>5.84 ± 0.11</td>
<td>5.52 ± 0.12</td>
<td>94.5</td>
<td>125.5</td>
</tr>
<tr>
<td>30</td>
<td>5.43 ± 0.08</td>
<td>6.86 ± 0.11</td>
<td>126.3</td>
<td>101.0</td>
</tr>
</tbody>
</table>

Table I. Kinetic parameters of the binding of mAb 7C4 to the peptide and carbohydrate Ags at different temperatures
were analyzed using BIAevaluation 3.2 to determine the values of the association and dissociation rate constants of binding at different temperatures (Table I).

The affinity of the Ab 7C4 for both 12-mer and mannopyranoside does not undergo any significant change on increasing the temperature from 15 to 30°C (Table I). The kinetics of binding to both the peptide and carbohydrate as a function of temperature were analyzed using BIAevaluation 3.2 to determine the values of the association (k_a) and dissociation (k_d) rate constants of the binding of mAb 7C4 to 12-mer and mannopyranoside with temperature. An Arrhenius plot for the calculation of activation energy (E_a) during each phase wherein the natural log of k_a or k_d has been plotted against the inverse of temperature. The comparison of the changes in enthalpy (ΔH) and entropy (ΔS) of binding to mannopyranoside and 12-mer peptide by mAb 7C4 at 30°C.

FIGURE 2. Thermodynamics of the binding of mAb 7C4 to the peptide immunogen and its mimicking carbohydrate Ag. A. Variation in the association (k_a) and dissociation (k_d) rate constants of the binding of mAb 7C4 to 12-mer and mannopyranoside with temperature. B. Arrhenius plots for the calculation of activation energy (E_a) during each phase wherein the natural log of k_a or k_d has been plotted against the inverse of temperature. C. The comparison of the changes in enthalpy (ΔH) and entropy (ΔS) of binding to mannopyranoside and 12-mer peptide by mAb 7C4 at 30°C.

were analyzed using BIAevaluation 3.2 to determine the values of the association and dissociation rate constants of binding at different temperatures (Table I).

The affinity of the Ab 7C4 for both 12-mer and mannopyranoside does not undergo any significant change on increasing the temperature from 15 to 30°C (Table I). The kinetics of binding to both the peptide and carbohydrate as a function of temperature appear to be similar during the association phase (Fig. 2A). In contrast, during the dissociation phase, while kinetics of mannopyranoside binding were insensitive to changes in temperature, there was a marginal increase in k_d of mAb 7C4 binding to 12-mer (Fig. 2A). This is also reflected in the strength of the Ab-Ag complex, the half-life of which decreases by ~2- and 5-fold while binding to mannopyranoside and 12-mer, respectively (Table I).

Thus, the binding of the anti-peptide mAb 7C4 to the immunogen and its mimicking Ag does not appear to be significantly influenced by variation in temperature. The decrease in the affinity at higher temperature is similar to that observed for other protein-ligand interactions that do not undergo major conformational changes during binding (17, 18, 12).

Energetics of Ag binding

To gain insights into the possible modes of binding, the energetics of interactions of 12-mer and mannopyranoside were analyzed in the context of the nature of the mAb 7C4-binding site. The change in Gibbs free energy (ΔG_eq), a measure of the favorability of a reaction, can be calculated from the K_eq such that the variation in the affinity of the Ab for the Ags can be directly correlated to the thermodynamics of binding to them. The change in ΔG_eq of mAb 7C4 binding to 12-mer and mannopyranoside, as a function of temperature, was ~2 kJ/mol (Table II). Thus, consistent with the lack of variation in the affinity (K_eq), there do not appear to be any significant temperature-dependent changes in the free energy of binding of the Ab to either of the mimics.

The effect of temperature variation on Ag binding was further analyzed by computing changes in enthalpy (ΔH) and entropy (ΔS) for the association and dissociation phases as well as at equilibrium. The activation energy (E_a) for either phase of binding was determined from the slope of Arrhenius plot which enabled calculation of changes in ΔH and ΔS at 30°C (Fig. 2B). It is interesting to note that while binding to the 12-mer is driven by favorable changes in enthalpy alone that to mannopyranoside is propelled by favorable changes in both enthalpy and entropy (Fig. 2C). The large negative, and therefore favorable change in enthalpy (ΔH_eq is ~83 kJ/mol) more than compensates for the unfavorable change in entropy (ΔS_eq equals ~43 kJ/mol) and drives the 12-mer binding. In contrast, binding of mAb 7C4 to the mimicking carbohydrate is characterized by favorable changes in both ΔH_eq (~30 kJ/mol) and ΔS_eq (~10 kJ/mol).

Comparable changes in enthalpy and entropy occur during the association phase of binding to either Ag. The favorable change in ΔH_eq of mAb 7C4 binding to the 12-mer is reinforced by unfavorable ΔS_eq. This results in large negative enthalpy at equilibrium that overrides the unfavorable ΔS_eq, thus driving the binding to the peptide immunogen (Fig. 2C). During binding to mannopyranoside, the net negative ΔH_eq is contributed by both favorable changes in ΔH_eq and unfavorable changes in ΔH_d (Fig. 2C). Additionally, large unfavorable ΔS_eq results in positive and thus favorable net change in entropy. Consequently, the net negative ΔG_eq at 30°C, during Ab-mannopyranoside complexation, is contributed by favorable changes in ΔS_eq and unfavorable changes in ΔH_d. Thus, binding of the mimicry recognizing Ab 7C4 to the sugar and peptide Ags differs mainly in the relative contributions of changes in enthalpy and entropy during the dissociation phase; while dissociation of mannopyranoside is disallowed by a negative ΔTΔS, unfavorable ΔH_d does not favor the dissociation of 12-mer. Favorable changes in entropy that accompany the binding of the carbohydrate ligand may indicate the critical role of hydrophobic interactions in complex formation (19).

Table II. Changes in Gibbs free energy (ΔG) at equilibrium as a function of temperature during binding of mAb 7C4 to 12-mer and mannopyranoside

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>ΔG_eq (kJ mol⁻¹)</th>
</tr>
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<tbody>
<tr>
<td>15</td>
<td>−42.8</td>
</tr>
<tr>
<td>20</td>
<td>−40.3</td>
</tr>
<tr>
<td>25</td>
<td>−40.1</td>
</tr>
<tr>
<td>30</td>
<td>−40.0</td>
</tr>
</tbody>
</table>

Table II. Changes in Gibbs free energy (ΔG) at equilibrium as a function of temperature during binding of mAb 7C4 to 12-mer and mannopyranoside
Ag binding by the anti-peptide and anti-carbohydrate Abs

To compare and contrast the modes of recognition by the anti-peptide and anti-carbohydrate Abs, both of which recognize the mimicry between manno- and 12-mer, analyses of the kinetics of peptide binding by the previously generated anti-mannopyranoside mAb 2D10 (12), were also conducted. Unlike the binding of mAb 7C4, that of mAb 2D10 to the 12-mer was drastically influenced by changes in temperature: while $k_d$ decreased by >8-fold, $k_a$ of mAb 2D10 binding to 12-mer increased 6-fold. This results in >50-fold loss in 12-mer affinity to mAb 2D10 at 30°C as compared with a 5-fold loss in the case of Ab 7C4-12-mer interaction (Fig. 3A). Consequently, the free energy of the interaction of the Ab 2D10 with 12-mer is less favorable by about $-10$ kJ/mol at 30°C, while the stability of the mAb 7C4-12-mer complex appears to be comparatively insensitive to temperature. Drastic variation in the kinetics of binding as a function of temperature suggests that large structural changes may occur during mAb 2D10-12-mer complexation. Comparatively, the mAb 7C4-binding site may not undergo major conformational alteration during binding of either ligand and structural variability in its paratope may be limited.

Comparison of the energetics of binding to 12-mer by the anti-peptide mAb 7C4 and anti-carbohydrate mAb 2D10 revealed differences in the extent of conformational flexibility in their Ag-binding sites. With regard to the enthalpy contributions for mAb 2D10 binding to 12-mer, highly favorable $\Delta H_a$ was aided by a large unfavorable $\Delta H_d$ (Fig. 3B). Although the trends were similar, significantly smaller changes in enthalpy were observed for mAb 7C4 binding to 12-mer especially during the association phase. This resulted in a larger favorable $\Delta H_{eq}$ for mAb 2D10 than for mAb 7C4 binding to the same Ag. However, in the case of mAb 2D10, the large favorable $\Delta H_{eq}$ is severely attenuated by unfavorable entropic changes, with the changes in $T\Delta S_d$ being highly unfavorable and $T\Delta S_a$ marginally favorable (Fig. 3C). In contrast, the entropy changes during both phases were relatively smaller and unfavorable during mAb 7C4-12-mer complexation. Despite binding to the 12-mer Ag by both the Abs being enthalpy driven, the high entropy penalty paid by the anti-mannopyranoside Ab suggests that major conformational changes accompany the binding of the peptide unlike relatively minor structural adjustments in the anti-12-mer Ab paratope.

Structure of the Ab paratope

The V regions of the L and H chains of the anti-peptide mAb 7C4 were sequenced to ascertain the nature of residues that form the Ag-binding site and to assign their germline origins. Five of the six CDRs could be assigned to the known classes of canonical structures: L1:9, L2:1, L3:6, H1:1, and H2:3 (20, 21). Analysis of the amino acid sequences showed that CDR H3, which is the specificity defining loop, had a larger proportion (9 of 12) of aromatic/hydrophobic residues vis-à-vis other CDRs (Fig. 4A). Ig-basic local alignment search tool sequence homology search (15) was conducted to assign germline origin and identify relatedness, if any, with other Abs analyzed in the context of mimicry. The germline genes were identified for the V and J elements of the L chain and the V, D, and J elements of the H chain. The L chain of mAb 7C4 belonged to the $\lambda$ family to which only 5% of all mouse L chains belong (22). It had 98.3% identity with the V$\lambda$1 and 100% identity to J$\lambda$1 germline sequence. The V region of the H chain of mAb 7C4 originated from the V$\gamma$GalSL1 V gene segment while the D region bears 100% homology to DFL16.2, and the J region has 92% homology to the J$\gamma$3 segment. Interestingly, the anti-peptide Ab 7C4 shares its germline origin with Abs generated in an anti-galactan response (23) although different from the antimannopyranoside mAbs 2D10 and 1H7 (12). It has been previously suggested that anti-sugar Abs probably share a common origin, especially for the H chain (24). In other words, a common pool of germline gene segments may be used for immune responses against not just carbohydrates but also their peptide mimics and other nonproteinaceous haptons.

The structure of the V region of the anti-12-mer Ab 7C4 was homology modeled using the structure of a catalytic Ab, 34E4 (PDB ID: 1Y0L), as template. The H and L chains of the Ab 34E4 showed 74 and 85% sequence identity with those of mAb 7C4. Model building involved transfer of coordinates of the backbone and the identical side chains of the Fv region from 1Y0L to the sequence of 7C4 and selection of the optimum set of coordinates for the CDR H3 loop by searching the PDB database. The resultant structure was energy minimized and stereochemically validated.

Fig. 4B depicts the Ag-binding site of the 7C4 Ab. As observed from the sequence analysis, there are a large proportion of aromatic residues in the Ab paratope contributed by CDR H and L3 loops. The Ag-combining site appears to have an elongated region that is lined by these aromatic residues (Fig. 4C). This clustering of aromatic residues, mainly from the specificity determining CDR H3, suggests that the binding of Ags, both peptide as well as carbohydrate, may be dominated by nonpolar interactions involving aromatic residues.

Epitope mapping on the peptide Ag

Having established that the anti-peptide mAb 7C4 binds the 12-mer and manno- to a comparable extent, it was pertinent to explore the epitope on the peptide that the Ab specifically recognized. This is relevant considering that the motif, Tyr-Pro-Tyr, is thought to be a specific mimic of manno- (13, 14). Additionally, the influence of aromatic cluster in the Ab paratope on the nature of interactions involved in peptide recognition was meant to be probed. We therefore identified the residues of the
12-mer directly involved in recognition by the mAb 7C4 by sequentially mutating each residue to glycine and evaluating Ab binding to the peptide analog.

The three hydrophobic residues, i.e., Phe3, Tyr6, and Tyr8, wherein Tyr6 and Tyr8 are part of the second Tyr-Pro-Tyr motif of the 12-mer, were found to be important for binding to the Ab in addition to the two serine residues, Ser10 and Ser12 (Fig. 4D). The average loss in binding to Phe3Gly, Tyr6Gly, or Tyr8Gly mutants was greater for mAb 7C4 as compared with other Abs that recognize the 12-mer-mannopyranoside mimicry (12), suggesting a more significant role for these hydrophobic residues in binding.

The recognition pattern indicated that alternate residues are important for binding, implying that the peptide may be bound such that a face of it interacted with the Ab, while intervening side
The immune system is remarkable in its ability to counter a limitless array of Ags that are varied in their chemical nature and structure. What is even more extraordinary is that the vast recognition repertoire is combined with exquisite specificity of the response against any invading Ag. However, molecular mimicry which results in the functional similarity of chemically independent Ags may be a manifestation of the breakdown in this specificity of immune recognition. We had previously addressed the functional mimicry between methyl-a-D-mannopyranoside and a Tyr-Pro-Tyr motif containing dodecapeptide in the context of polyclonal immune responses (4, 7). Molecular analyses of the physiological responses against the mimicking Ags necessitated generation of mAbs and comparison of their differential recognition specificities. The mAb 7C4, raised against the 12-mer, binds to the mimicking carbohydrate moiety in addition to the peptide immunogen. The two Ags could compete with one another and bind with comparable affinity to the Ab. Furthermore, similar binding kinetics at ambient temperature suggested possibility of a common mode of recognition of the peptide and the carbohydrate Ag by the Ab 7C4.

Thermodynamic and energetic analyses of binding provided insights into the mode of interaction of mAb 7C4 with 12-mer and mannopyranoside. Binding of the Ab to both the Ags appears to be considerably independent of changes in temperature. Although the association rate constants during binding to either 12-mer or mannopyranoside remain almost unchanged with respect to temperature, marginal increase in $k_a$ results in a minor loss in affinity of the 12-mer-Ab interaction. In terms of the temperature-dependent changes in $\Delta G_{eq}$ for binding to peptide or carbohydrate Ag, mAb 7C4 behaved differently from the mimicry-recognizing Ab 2D10 that was raised against mannopyranoside. Interestingly, the thermodynamic behavior of mAb 7C4 was similar to the mannopyranoside-specific Ab 1H7, which did not recognize carbohydrate-peptide mimicry (12).

The lack of significant temperature-dependent variation in the kinetics of binding indicates the absence of unrestricted flexibility in the Ag-combining site (17, 25). Deficiency in entropy penalty may also be indicative of the minor role of conformational alteration in the paratope during complexation with either Ag (26). In contrast, favorable entropy contribution during binding of mannopyranoside highlights the importance of hydrophobic/aromatic associations (27, 28, 19). Thus, it can be inferred that the binding of the Ab 7C4 to the two ligands does not involve large structural changes in the backbone of the CDR loops. However, minor variation in the side chain orientations is possible. Consequently, unlike the anti-mannopyranoside mAb 2D10, manifestation of mimicry in the anti-12-mer Ab 7C4 may not be attributable to conformational flexibility in the paratope that allows for the possible existence of multiple and/or additional Ag-binding subites in dynamic equilibrium.

It was indeed intriguing that neither paratope conformational flexibility nor epitope structural similarity facilitated recognition of the chemically distinct yet mimicking 12-mer and mannopyranoside by mAb 7C4. Exploring alternative explanations for the mimicry, nature of the residues involved in the epitope-paratope interaction was analyzed. It was observed that the L chain of Ab 7C4 belongs to the $\lambda$ subfamily to which only ~5% of all mouse L chains belong, which is however, common in anti-hapten Abs (29). In contrast, the H chain of mAb 7C4, which was raised against the dodecapeptide, shares its germline origin with anti-carbohydrate Abs. It is interesting to note that $V_\mu$ gene segment of mAb 7C4 was discovered from an immune response against $\beta$-(1,6)-galactan (23). Sharing of common germline origin by anti-peptide and anti-carbohydrate Abs may not be incidental but linked to molecular mimicry.

The structural model of the V region of the Ab 7C4 facilitated correlation of the topological features of the Ag-combining site with the thermodynamics of epitope-paratope interaction. The Ab paratope is characterized by predominance of aromatic residues which may mediate molecular mimicry in the absence of flexibility in the Ag-combining site. Critical involvement of aromatic residues has also been observed in carbohydrate-Ab interactions especially in the context of specificity of ligand recognition (30, 31). Epitope mapping analysis of the 12-mer peptide indicated maximum contributions from Phe$^3$, Tyr$^6$, and Tyr$^8$ for Ab binding. Thus, the clustering of aromatic residues in the paratope as well as in the epitope reinforced the hypothesis that plasticity associated with aromatic (hydrophobic and $\pi-\pi$) and van der Waals interactions dominate in recognition of the 12-mer and mannopyranoside, both of which possess ring-like structural components with associated aromaticity.

It is important to note that 12-mer and mannopyranoside are functionally equivalent to an impressive extent although they do not share a high level of structural similarity (4, 7). The two Ags bind to the Ab 7C4 with high affinity and specificity mediated by interactions involving aromatic residues, especially tyrosine. Such interactions are known to broaden the specificity repertoires of not just Abs but have also been implicated in receptor-ligand, enzyme-substrate, and lectin-carbohydrate multispecificity (32–37). Significance of aromatic residues in defining the specificity of epitope-paratope interaction has been extensively addressed, highlighting the importance of tyrosine residues. It has been suggested that this residue is capable of mediating favorable interactions with a diverse array of surfaces endowing the amino acid with a privileged role in Ag recognition (38). Numerous Ab CDR H3s have been identified with high tyrosine content and many of these Abs possess diversity in their recognition potential (39).

Our laboratory has explored diverse aspects of molecular mimicry in terms of immune responses or receptor binding (4–12). The correspondence between mannopyranoside and 12-mer provided an elegant model for correlating functional and structural aspects of molecular mimicry. The functional similarity between the two mimicking ligands in the humoral immune response, both at the monoclonal as well as at the polyclonal level, was overwhelmingly more significant than the topological relationship between these Ags. Despite insufficient structural correlation, polyclonal Abs elicited against the 12-mer could recognize the carbohydrate mimic and the peptide can act as a booster for enhancing the anti-mannopyranoside immune response and vice versa (4, 7). Thus, the two Ags exhibit functional similarity in the context of the humoral immune response. Similar binding to a common receptor by ligands with limited or no apparent structural similarity has also been illustrated in case of other promiscuous Abs (40, 41). Structural equivalence, involving not just similarity of shape and charge but also conservation of key interaction features in terms of a definable motif, such as a network of hydrogen bonds, would appear to be the most obvious rationale for functional mimicry (6, 10, 11). Contrary to expectations, molecular analyses of the anti-peptide and anti-carbohydrate Ab responses revealed diverse
Molecular mimicry was commonly considered to manifest by structural equivalence between them (Fig. 5). As seen in the present analysis with anti-peptide Ab 7C4, clustering of aromatic residues in the Ab paratope allowed binding of mimicking Ags (Fig. 5A). This could potentially allow chemically dissimilar Ags like carbohydrates and peptides to behave as functional equivalents. In addition to aromatic interactions, a combination of other noncovalent interactions could also bring about such plasticity—while maintaining structural integrity—of the binding site resulting in functional mimicry.

Another mechanism for manifestation of molecular mimicry is the invocation of conformational flexibility in the paratope (Fig. 5B). Existence of multiple conformational states of the Ag-combining site, in dynamic equilibrium, could potentially allow for the binding of various functional mimics. The mimicry-recognizing anti-carbohydrate Ab 2D10 exemplified this mode of molecular mimicry. In contrast, the paratope of the mannopyranoside specific mAb 1H7 perhaps is predesigned for the sugar immunogen and therefore does not bind to the peptide mimic (12). Alternatively, conformational flexibility in the Ag can also augment effective mimicry (9). Additional relevant factors that could modulate molecular recognition in favor of mimicry include imprinting by in nocuous molecules including ions and solvent molecules resulting in improvement in shape and/or electrostatic complementarity between the paratope and epitope (10, 42).

Molecular mimicry was commonly considered to manifest by topological equivalence between the Ags with the Ab receptors possessing well-defined structures. It was also anticipated to manifest without topological equivalence between the Ags if the paratope was capable of changing structure to complement the epitopes. Our present studies have revealed molecular mimicry in the absence of either structural similarity of the epitopes or conformational flexibility in the paratope. The adaptable nature of interactions between the Ab and the Ags arising from plasticity inherent to aromatic/hydrophobic interactions illustrated a distinctive mode of binding of mimicking Ags to a common conformation of the Ab paratope. Molecular mimicry being modulated by receptor-specific properties is relevant in the context of pathogens changing their antigenic determinants for evading immune surveillance; the immune system needs to recognize altered Ags despite their structural differences. Therefore, physiological responses against the invading pathogens may include immune receptors with broader recognition specificities to minimize the consequences of antigenic variation.

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